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- 34 Abstract
- 35 *INTRODUCTION*: Ischaemic preconditioning (IPC) may enhance
- 36 endurance performance. No previous study has directly compared
- 37 distinct IPC protocols for optimal benefit. The aim of this study was
- to determine whether a specific IPC protocol (i.e. number of cycles,
- 39 amount of muscle tissue, and local vs remote occlusion) elicits
- 40 greater performance outcome.
- 41 *METHODS:* Twelve cyclists performed five different IPC protocols
- 42 30-min prior to a blinded 375 kJ cycling time trial (TT) in a
- 43 laboratory. Responses to traditional IPC (4x5-min legs) were
- compared to: i. 8x5-min legs and SHAM ("dose-cycles"), ii. 4x5-
- 45 min unilateral legs ("dose-tissue"), and iii. 4x5-min arms
- 46 ("remote"). RPE and blood lactate were recorded at each 25% TT
- 47 completion. Power (watts), heart rate (bpm), and $\dot{V}O_2$ (ml.kg.min⁻¹)
- 48 were measured continuously throughout TT's. Magnitude based
- 49 inference statistics were employed to compare variable differences
- to the minimal practically important difference.
- 51 RESULTS: Traditional IPC was associated with a 17 (0, 34) secs
- faster TT time compared to SHAM. Applying more "dose-cycles"
- 53 (8x5-min) had no impact on performance. Traditional IPC was
- associated with "likely trivial" higher blood lactate and "possibly
- beneficial" lower $\dot{V}O_2$ responses vs. SHAM. Unilateral IPC was
- associated with 18 (-11, 48) secs slower performance compared to

- 57 bilateral ("dose-tissue"). TT times following remote and local IPC
- were not different [0 (-16, 16) secs].
- 59 CONCLUSION: The traditional 4x5-min (local or remote) IPC
- stimulus resulted in the fastest TT time compared to SHAM, there
- was no benefit of applying a greater number of cycles or employing
- 62 unilateral IPC.
- 63 **Key words:** Exercise, Occlusion, Ischaemia, Time Trial, Endurance

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Introduction

- 66 Ischaemic preconditioning (IPC) refers to the phenomenon whereby
- 67 3-4 brief periods of ischaemia, followed by tissue reperfusion,
- 68 confers subsequent tissue protection against ischaemic insult ¹. IPC
- 69 can be applied remotely by placing a blood pressure cuff around a
- 70 limb and inflating to supra-systolic pressure. Studies have generally
- 71 employed remote IPC in clinical populations relating to cardio-
- 72 protection, but there is accumulating evidence that remote IPC can
- 73 impact on other organs (e.g. skeletal muscle), and vascular beds to
- facilitate increased blood flow 2,3 . These finding have resulted in the
- 75 application of IPC to determine its efficacy as a potential pre-
- 76 exercise priming strategy.
- 77 The first study to investigate IPC in a human exercise model
- demonstrated a 3% improvement in maximal oxygen uptake ($\dot{V}O_2$)

following a 3x5-min bilateral leg cuff inflation (220 mmHg) protocol ⁴. A "traditional" IPC protocol consists of 3x5- or 4x5-min bouts of occlusion. More recently, studies have separately employed alternative IPC protocols (altering the number of IPC cycles, tissue occlusion area, and cuff location), with the aim of observing greater performance and clinical outcomes. There are now (pre)clinical studies providing evidence for a "dose"-dependency, where repeated daily IPC improves (cerebro)vascular function and clinical outcomes ^{5,6}. Nonetheless, a potential 'hyper-conditioning' effect from excessive cycles of IPC cannot be excluded ⁷. Corroborating the "dose"-hypothesis, recent work suggests that bilateral, but not unilateral cuff inflation leads to improved exercise performance ⁸. Finally, most studies to date have opted for cuff positioning directly on the exercising limb 9, but cuff placement on remote, nonexercising limbs has also been performed ¹⁰ to examine a systemic effect. In line with clinical observations in the protection of organs against ischaemic injury, local or remote application of IPC may induce comparable benefits ^{2,11}. Recently, a systematic review and meta-analysis reported a small beneficial effect of IPC on exercise performance, with the largest effect observed in aerobic-based tasks ¹². Despite the effect sizes being small, the potential benefits of IPC may translate to meaningful differences in competitive (time trial-based) events.

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102 Interestingly, no study has directly compared the capacity of distinct IPC protocols with the aim of electing greater performance 104 improvement. Therefore, the aim of this study was to examine whether the (i) number IPC cycles (i.e. "dose-cycles"), (ii) the amount of muscle mass occluded ("dose-tissue"), and (iii) the 106 107 application of IPC to either local or remote limbs ("remote") offers greater improvements to endurance cycling performance.

Methods

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Participants

Twelve trained cyclists (mean±SD: age, 36±7 years; body mass, 78 ± 4 kg; height, 179 ± 6 cm; $\dot{V}O_{2max}$, 59 ± 4 ml.kg⁻¹.min⁻¹) were recruited. Participants were undertaking regular weekly training sessions (5±3 sessions) and mean weekly training volume was 8±4 hours. The mean training experience was 9±8 years. Following verbal and written explanation of procedures, all participants provided written informed consent. Physical Activity Readiness Questionnaires were administered to ensure no participant had any health implications that would prevent participation. All individuals refrained from exercise and alcohol consumption 24 hours, and consumption of caffeine at least 6 hours, respectively prior to all laboratory visits. The study was approved by the local Ethics Committee.

Research Design

The study was divided into three comparisons as illustrated in figure 1. All participants completed a maximal graded cycling test and at least two familiarization TT. Prior to commencement of the five experimental cycling TT's, an IPC protocol was administered. A traditional (4x5-min) IPC protocol was compared firstly to SHAM, and a larger (8x5-min) IPC protocol for the "dose-cycles" comparison. Whilst it was compared to a unilateral (4x5-min) IPC protocol for the "dose-tissue" comparison. Finally, to assess the importance of cuff placement, a 4x5-min bilateral IPC protocol was applied to the non-exercising upper limb for the "remote" comparison.

Experimental Protocol

In a randomized, counterbalanced, crossover study, participants reported to the laboratory at the same time of day on five separate occasions, at least 4 days apart, receiving a different pre-exercise IPC protocol during each visit. Following each IPC protocol, a 20-minute rest period, and a standardized warm up was performed before the completion of a 375 kilojoule (kJ) cycling time trial (TT). The TT was intended to simulate the demands of a 16.1 km TT. During each TT, heart rate and oxygen uptake ($\dot{V}O_2$) was measured continuously, whilst blood lactate and rate of perceived of exertion

(RPE) was recorded at every 25% completed of the TT kilojoule target.

Measurements

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Assessment of maximal oxygen uptake ($\dot{V}O_{2max}$). At least 7 days prior 149 150 to the first familiarisation trial, participants performed a continuous 151 incremental step test on an electromagnetically braked cycle 152 ergometer (SRM, Julich, Germany) to determine lactate threshold 153 and $\dot{V}O_{2max}$. The incremental protocol consisted of 3-minute cycling 154 stages, commencing at 95 watts (W) and increasing 35W until 155 volitional exhaustion occurred. Blood lactate concentration was 156 obtained via finger prick capillary sampling using a safety lancet 157 (BD Microtainer® Contact-Activated Lancet) after administration 158 of a disposable sterile isoprophyl alcohol swab (China MEHECO 159 Co., Ltd.). Blood was collected into a sodium-heparinized blood gas 160 capillary tube (Marienfeld Superior, Germany) and immediately 161 analysed in duplicate (ABL90 FLEX, Radiometer Medical ApS, 162 Denmark) during the last 30 seconds of each incremental stage. 163 Throughout the incremental cycling test, breath-by-breath expired 164 gases were monitored for oxygen consumption, ventilation and 165 respiratory exchange ratio (RER) (MasterScreenTM CPX, 166 Carefusion, Germany) and the highest 30-second average was taken 167 3 consecutive 10-second bins from subsequently to determine $\dot{V}O_{2max}$. Heart rate (HR) was also monitored continuously 168

169 (Polar H1, Kempele, Finland). W_{max} was calculated from the last 170 completed workload, plus the fraction of time spent in the final non-171 completed stage multiplied by the work rate increment ¹³.

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Familiarisation. At least 2 familiarisation trials were undertaken prior to the first experimental TT to ensure performance was reliable. Data from familiarisation sessions revealed a mean coefficient of variation (CoV) of 1.06% which was deemed to be acceptable for the purpose of this TT study.

IPC protocols. For the IPC and SHAM trials, 13.5 cm wide cuffs were used. Participants lay in the supine position and cuff inflation pressure was set at a standardized pressure (220mmHg) in all IPC conditions with the aim of preventing arterial inflow 14 and 20mmHg in SHAM (i.e. cuffs were placed but only inflated to 20mmHg) with the use of an automatic rapid cuff inflator (Hokanson, Washington, USA). Subsequently, cuffs were deflated for 5 minutes, allowing reperfusion. This process was repeated four times in all protocols except for the "dose-cycles" protocol where 8 cycles were used (Figure 1). For IPC on the leg, the cuff was placed (unilaterally or bilaterally) on the most proximal portions of the upper thigh (distal to the inguinal fold). For remote IPC, cuffs were placed on the most proximal portions of the upper arms. Each participant gave a "perceived discomfort" rating at four time points (every 25%) throughout the IPC or SHAM protocols. The discomfort rating was

established using a Numerical Rating Scale (NRS) ranging from 0 193 (no discomfort) to 10 (maximum discomfort) and are included for descriptive purpose (Table 4) ¹⁵. 194 195 375 kJ TT. After 20 minutes of rest following cessation of 196 IPC/SHAM, a capillary blood lactate sample was obtained from the 197 finger and analysed for resting lactate levels (ABL90 FLEX, 198 Radiometer Medical ApS, Denmark). Participants then completed a 199 standardized warm up on an electromagnetically braked cycle 200 ergometer (SRM, Julich, Germany). The warm up lasted 201 approximately 10 minutes (5-min at 100W, 2-min at 150W, [15-secs 202 at W_{max} , 30-secs at 150W, repeat x3], 45-secs at 150W). Once the 203 flywheel had completely stopped turning, the SRM clock was reset 204 to zero and a 375 kJ TT was performed (exactly 35 minutes after 205 completion of IPC in all trials). Participants were instructed to 206 produce a maximum effort throughout TT's, but were blinded to 207 power output, elapsed time and HR. Breath-by-breath expired gases 208 and HR were measured continuously, while RPE and blood lactate 209 measurements were acquired at 25%, 50%, 75% and 100% time 210 points (all described previously). Participants were notified once 211 they had completed each quarter of the TT and when they had 30 kJ 212 of work remaining. No encouragement or feedback was given

Statistical Analysis

throughout any trial.

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The primary outcome variable was TT time and was analyzed using a repeated measures general linear modelling for "dose-cycles" (3 levels: SHAM, 4x5-min, 8x5-min) and paired t-tests for "dosetissue" (2 levels: unilateral, bilateral) and 'remote' (2 levels: local, remote). For TT measures, $\dot{V}O_2$, power, lactate, HR, and RPE were analyzed using repeated measures general linear modelling. The least significant method was employed for pairwise comparisons ¹⁶. Using a magnitude based inferences framework, the mean effect of each TT comparison for each variable was presented with the uncertainty of the estimates presented as 90% confidence intervals (appropriate SI units used for a given variable). The mean difference between each comparison were evaluated for their practical significance by pre-specifying the smallest worthwhile change (SWC) ¹⁷. For TT time and power output, the SWC was calculated using 0.3 x coefficient of variation from the familiarization trials, equating to 4.5 seconds and 1 watt, respectively ¹⁸. The noise to signal ratio was determined by calculating the typical error (SD of between-trial differences divided by $\sqrt{2}$). The typical error for time and power was 18 seconds and 4 watts, respectively. For blood lactate and $\dot{V}O_2$ the SWC was calculated using the standardized mean difference of 0.2 between subject standard deviations (SD) as they were not measured during the familiarisation trials ¹⁹. The SHAM values were used for this purpose. The mean difference

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between each comparison, together with its uncertainty, the probability (percent chances) that the true population effect was beneficial (>SWC), harmful (>SWC with opposite sign), or trivial (within \pm SWC) was calculated ¹⁸. Using mechanistic inferences, qualitative probabilistic terms for benefit were assigned to each effect using the following scale; <0.5%, most unlikely or almost certainly not; 0.5 to 5%, very unlikely; 5 to 25%, unlikely or probably not; 25 to 75%, possibly; 75 to 95%, likely or probably; 95 to 99.5%, very likely; >99.5%, most likely or almost certainly ¹⁸. An unclear effect is possibly beneficial (>25%) with an unacceptable risk of harm (>0.5%) and an odds ratio for benefit:harm of <66 interpreted from current recommendations; all other effects are clear. Data that were lower than the typical error (noise > signal) for TT performance were interpreted as "unclear" and reported with the confidence limits within the text and in figure 2.

Results

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Dose-cycles

TT time: TT time was 17 secs (90% CI: 0, 34 secs; P=0.097) faster following the traditional IPC protocol compared to SHAM. The mean change is lower than the noise so is interpreted as "unclear" with the following confidence limits 89% chance beneficial, 9% chance trivial and 2% chance harmful (Figure 2b). Increasing the "dose" by applying more cycles (8x5-min) did not result in a faster

- TT time compared to traditional IPC (4x5-min) [13 secs (-19, 44
- secs); *P*=0.49, (beneficial 67%, trivial 15%, harmful 18%)] Figure
- 263 2]. The effect between IPC with 8x5-min cycles and SHAM on
- exercise performance was interpreted as "unclear" (beneficial 50%,
- 265 trivial 19%, harmful 31%).
- 266 $\dot{V}O_2$: $\dot{V}O_2$ was 0.99 ml.kg.min⁻¹ (-1.7, -0.3 ml.kg.min⁻¹) lower
- 267 following traditional IPC compared to SHAM, interpreted as
- 268 "possibly beneficial" (beneficial 59%, trivial 41%, harmful 0%;
- 269 P=0.03). A "likely trivial" difference was evident between
- traditional IPC and the 8x5-min protocol [0.51 ml.kg.min⁻¹(-1.2, 0.2)
- 271 ml.kg.min⁻¹); (beneficial 17%, trivial 83%, harmful 0%) P=0.25].
- 272 Lactate: Blood lactate increased throughout TT performance, with
- 273 highest values observed during the 4th quarter (Table 1). Traditional
- 274 IPC was associated with a higher mean TT blood lactate compared
- 275 to SHAM [0.73 mmol.L⁻¹ (0.1, 1.5 mmol.L⁻¹); P=0.06, "possibly
- 276 trivial" (beneficial 42%, trivial 58%, harmful 0%)] and to the 8x5-
- 277 min protocol [0.9 mmol.L⁻¹ (0.4, 1.9 mmol.L⁻¹); P=0.006, "possibly
- beneficial" (beneficial 73%, trivial 27%, harmful 0%)].
- 279 Power / HR / RPE: HR and RPE increased significantly across time
- 280 (P<0.05), whilst power was highest during the 1st quarter. No
- further differences were evident for power, HR, or RPE between
- traditional, SHAM and 8x5-min (all P>0.05; Table 1).

Dose-tissue

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- 284 TT Time: Traditional bilateral IPC resulted in an 18 secs (-11, 48
- secs, P=0.29; Figure 2) faster TT performance than unilateral IPC.
- Nevertheless, this change was interpreted as "unclear" (beneficial
- 287 78%, trivial 12%, harmful 10%).
- 288 $\dot{V}O_2$: The lower resultant $\dot{V}O_2$ following traditional IPC compared
- to unilateral IPC [0.8 ml.kg.min⁻¹; (-2, 0.4 ml.kg.min⁻¹); (beneficial
- 290 45%, trivial 54%, harmful 1%) P=0.26)] was interpreted as
- 291 "possibly trivial". The time-dependent effect (Table 2), was not
- 292 different between the 2 trials.
- 293 Lactate: Blood lactate increased throughout TT performance, with
- 294 highest values during 4th quarter (Table 2). The mean blood lactate
- 295 difference of 0.05 mmol.L⁻¹ (-1.3, 1.4 mmol.L⁻¹); (beneficial 11%,
- 296 trivial 81%, harmful 9%; P=0.95) between protocols was
- interpreted as "unclear".
- 298 Power / HR / RPE: HR and RPE increased significantly across time,
- 299 whilst power was highest during the 1st quarter (Table 2) .No further
- differences were evident for power, HR, or RPE (Table 2).

Remote

- TT time: The comparison of traditional IPC and remote IPC resulted
- in a negligible difference in mean TT time [0 secs (-16, 16 secs;

- 304 P=1.0, Figure 2a)]; interpreted as an "unclear" (beneficial 50%,
- 305 trivial 0, harmful 50%).
- 306 $\dot{V}O_2$: $\dot{V}O_2$ was 1.1 ml.kg.min⁻¹ (-1.9, -0.2 ml.kg.min⁻¹; (beneficial
- 307 71%, trivial 29%, harmful 0%) P=0.04) lower following the
- 308 traditional protocol compared to remote IPC, interpreted as a
- 309 "possibly beneficial" reduction.
- Lactate: Blood lactate increased throughout both TT performances,
- with highest values observed during 4th quarter (Table 3). A mean
- blood lactate difference of 0.2 mmol.L⁻¹ occurred (-1.2, 1.6 mmol.L⁻¹
- 313 1 ; P=0.8) between both protocols, interpreted as an "unclear"
- difference (beneficial 18%, trivial 74%, harmful 8%).
- Power / HR / RPE: HR and RPE increased significantly across time,
- 316 whilst power was highest during the 1st quarter. No further
- differences were evident for power, HR, or RPE between traditional
- and remote IPC (Table 3).

319 Discussion

- 320 The aim of this study was to determine the impact of different IPC
- 321 protocols on cycling endurance performance. Specifically we
- explored, for the first time, whether the "dose" of IPC, reflected by
- 323 either the number of cycles, or the amount of muscle tissue
- occluded, affects endurance cycling TT performance. We provide
- evidence that the traditional (4x5-min) occlusion/reperfusion cycles

326 resulted in the fastest TT times. Our data may support application of 327 a traditional IPC "dose" of cycles, since increasing the "dose" by applying more cycles and reducing the "dose" by applying unilateral 328 329 IPC, resulted in no further benefit to endurance performance. 330 Furthermore, our study provides evidence that the same magnitude of change in TT time (17 seconds) occurs when exposed to either 331 332 local or remote application of IPC. 333 Ischaemic preconditioning, applied using the traditional (4x5-min) 334 inflation/reperfusion cycles ^{9,20–24}, mediated an effect that was an 335 unclear performance improvement in a 375 kJ cycling TT based on 336 a the signal to noise ratio. The improvement of 17 seconds following 337 traditional IPC vs SHAM is marginally below the calculated error 338 and the confidence intervals do not cross zero therefore we are 339 confident that a directional change is present in favor of a 340 worthwhile performance improvement. Furthermore, 341 observation of a 1.4% performance change is largely in line with 342 previous reports examining the impact of traditional IPC on 343 endurance-type exercise tasks ¹², but it is important to emphasise 344 that we included a trained population (natural coefficient of 345 variation of 1.1%); something not commonly observed to date in 346 time-trial based performance tasks, with the exception of competitive swimmers ^{20,25,26}. The research evidence suggests IPC 347 348 can improve exercise capacity in recreationally trained participants

349 ⁴, but one recent study demonstrated that in highly trained athletes, 350 IPC provided little benefit in improving exercise capacity ²⁷. 351 Whether a higher aerobic capacity blunts the ergogenic effect of IPC 352 on exercise performance using sports specific tasks remains to be 353 determined. 354 Importantly, the difference in TT time following a larger "dose", 355 through applying more (8x5-min) cycles in one session, was not 356 deemed substantial enough, when compared to SHAM, to be of 357 benefit. In addition, a smaller "dose" by applying unilateral IPC had 358 little beneficial impact on performance. These results suggest for the 359 first time, that IPC-mediated performance improvements are 360 unlikely amplified by doubling the "traditional" number of IPC 361 cycles. Nevertheless, it is unclear whether an area threshold is 362 present for the "dose-tissue". Whilst no negative impact on TT time 363 was suggested from the magnitude based inference, the lack of 364 additional benefit on exercise performance after the 8x5-min 365 protocol provides support for the 'hyperconditioning' hypothesis, in 366 that too many cycles may negate the beneficial effects of IPC ⁷. 367 A recent animal model corroborated these findings and 368 demonstrated four to six cycles yielded cardioprotection, with no further benefit after using eight cycles ²⁸. Additionally, it was found 369 370 that when using four cycles, both unilateral and bilateral hind-limb

occlusion offered similar cardioprotection ²⁸. The current study

findings suggest a bilateral "dose", but not unilateral "dose", may result in greater endurance performance; an outcome in line with one previous human study showing bilateral, but not unilateral IPC improved anaerobic sprint cycling performance 8. Whilst our data is specific to aerobic exercise performance, it may be possible that an "area threshold" i.e. a required amount tissue occlusion, is required to stimulate IPC-induced performance improvements, regardless of intensity ^{8,29}. Remote IPC can elicit cardio protective effects, comparable to local IPC, possibly as a result of a humoral trigger signal or circulating factor ²⁰. To date, the comparison between remote and local IPC has not been directly examined in an human performance setting, although both protocols have been previously reported to enhance performance when compared to SHAM ^{8,9}. In our study, we provide the first direct evidence that local and remote application of IPC resulted in the same TT performance (288 watts, respectively). Whether a systemic pathway contribution towards improved exercise performance occurs, such as a humoral trigger signal or circulating factor similar to that shown with cardioprotection ²⁰ remains to be seen. Interestingly, clinical application of IPC locally or remotely is associated with a comparable protective effect against ischaemia-reperfusion injury in animals and humans ¹¹.

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394 TT performance after the traditional IPC "dose" was accompanied by a lower $\dot{V}O_2$ when compared to SHAM. Our data also reveal a 395 396 lower TT $\dot{V}O_2$ for the same given workload (288w average) 397 following local, compared to remote IPC. Whilst local IPC 398 application can increase pig skeletal muscle metabolic efficiency 399 under ischaemic conditions ², it remains unknown whether 400 previously observed local IPC-induced metabolic adaptations ^{9,30} 401 may have contributed to these findings. Nevertheless, the current 402 data are suggestive that traditional IPC, applied locally, enhances 403 the ability to sustain the same workload for a relatively lower 404 oxygen cost compared to both SHAM and remote IPC, but this does 405 not necessarily relate to clear improvements in power output. 406 We recorded lactate measurements at each 25% stage of TT 407 performance and found the traditional "dose" of IPC increased 408 blood lactate during exercise when compared to both SHAM and the 409 8x5-min condition. This finding is somewhat intriguing given that 410 we have previously reported a lower onset of blood lactate 411 accumulation (OBLA) during submaximal exercise following 4x5-412 min (traditional) bilateral IPC compared to SHAM, hypothesizing 413 greater lactate removal and transportation for uptake ⁹. A logical 414 explanation for this apparent contrasting result is that workload in 415 the current cycling TT task markedly exceeds that at OBLA. The 416 increased blood lactate response in the current study following 4x5min local bilateral IPC, combined with lower VO2, could be suggestive of alterations in substrate utilisation, with a proposed heightened anaerobic energy contribution. This was recently inferred by Cruz et al. ³¹, who demonstrated 4x5-min cycles of IPC improved 60-second sprint cycling performance and lead to an increased skeletal muscle activation during exercise, whilst during recovery produced higher amplitude of blood lactate kinetics and increased excess post-exercise oxygen consumption (EPOC), when compared to SHAM exercise. This, in combination with our data, suggests the potential ergogenic mechanisms relating to IPCinduced metabolic alteration, is likely task and/or intensity specific. The capability of IPC to enhance aerobic exercise capacity ^{4,29,30}, yet have smaller ergogenic effects on fixed-end-point performance ¹² is a relationship also observed following the use of nitrate based dietary interventions ³² and might provide some insight into potential mechanisms. A systematic review and meta-analysis ¹² recently reported IPC can enhance incremental exercise performance, time to exhaustion task performance, and fixed-end-point task performance by 2.4%, 5.8% and 0.5%, respectively. Additionally, Ferreira et al. ²⁵ stated the estimated performance improvement of IPC was 1.5% based on some previous study findings 9,20,29. The current observed performance changes (1.4%) are broadly in line with the above

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studies, yet the cycling mode we employed was a fixed-end-point task. We further delimited the impact of pacing strategy with rigorous familiarization trials (mean co-efficient of variation in TT time between trials was $1.1\% \pm 0.8\%$), and selecting only trained cyclists as participants.

Practical Applications:

IPC is a well-tolerated intervention for the competing individual (table 4). The magnitude of improvement after a bilateral 4x5-min protocol, independent of whether cuffs are placed locally (upper thighs) or remotely (upper arms), lead to improvements in finish time. This conclusion is based on the calculated typical error of our laboratory based test. Given the performance changes in laboratory based tests are different to the field and in competition (e.g. power-velocity relationship on the road is cubic and not linear) this needs to be taken into account when applying these findings to road competition.

Conclusion

Our results suggest the "traditional" protocol of IPC involving 4x5-min occlusion is associated with the fastest TT time compared to SHAM, in a laboratory 375 kJ TT task, aimed to simulate demands of a 16.1 km road TT race. Moreover, by applying different IPC protocols in a within-subject cross-over design, our data suggests no

- 462 benefit when increasing the "dose" by doubling the number of 463 cycles or reducing the "dose" via implementing unilateral IPC. 464 Finally, TT performance after IPC appears to be independent of the 465 localization of the cuffs, as IPC applied to the upper limbs resulted 466 in the same TT time. 467 Acknowledgements: The authors would like to thank the 468 committed participants for giving up their time to undertake 469 physically demanding protocols, in order to obtain the data for this 470 study. 471 **Conflict of Interest:** None to declare. Results of the present study 472 do not constitute endorsement by any party and all results are 473 presented clearly, honestly, and without fabrication or falsification. **References:** 474 475 Murry CE, Jennings RB, Reimer KA. Preconditioning with 476 ischemia: a delay of lethal cell injury in ischemic 477 478 myocardium. Circulation. 1986;74(5):1124-1136. 479 Pang CY, Yang RZ, Zhong A, Xu N, Boyd B, Forrest CR. 480 Acute ischaemic preconditioning protects against skeletal muscle infarction in the pig. Cardiovasc Res. 1995;29(6):782-481 788. 482 483 3. Kraemer R, Lorenzen J, Kabbani M, et al. Acute effects of 484 remote ischemic preconditioning on cutaneous microcirculation-a controlled prospective cohort study. BMC 485 486 Surg. 2011;11(1):32.
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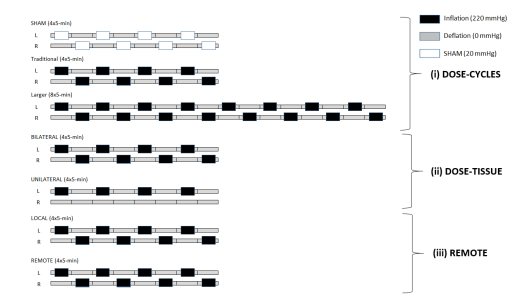
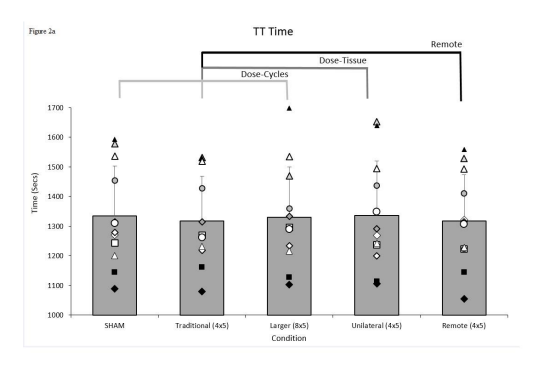


Figure 1 – Schematic of different of IPC protocols (i) comparison of dose-cycles (ii) comparison of dose-tissue and (ii) comparison remote. (N.B. traditional dose of IPC was performed once in the experimental design but is shown 3 times on schematic to highlight the comparisons).





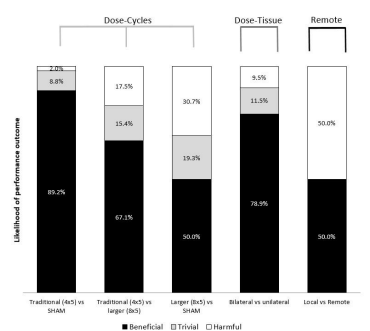


Figure 2a – Overall TT times (with individual times plotted) for IPC (i) comparison of dose-cycles (ii) comparison of dose-tissue (iii) comparison of remote.

Figure 2b – A between-condition representation of the likelihood of "beneficial", "trivial", or "harmful" performance outcome to endurance cycling TT performance.

Tables:
Table 1: The effect of "dose-cycles" on power,

Table 1: The effect of "dose-cycles" on power, heart rate, rate of perceived exertion and $\dot{V}O_2$ following 25%, 50%, 75% and 100% time points during time trial performance.

	Intervention					P values	
	Average	0-25%	25-50%	50-75%	75-100%		
Power (watts)							
4x5	288 ± 33	310 ± 38	284 ± 34	275 ± 34	285 ± 32	Condition	0.57
8x5	286 ± 35	307 ± 37	284 ± 35	273 ± 37	281 ± 36	Time	< 0.005
SHAM	285 ± 35	305 ± 38	282 ± 39	273 ± 35	284 ± 35	Condition x time	0.99
Lactate (mmol.L ⁻¹)							
4x5	11.8 ± 2.8	10.8 ± 3.4	11.8 ± 3.2	12.4 ± 2.9	13.4 ± 2.8*	Condition	0.02
8x5	11.2 ± 3.1	10.6 ± 3.7	11.2 ± 3.2	11.3 ± 3.3	11.6 ± 3.2*	Time	< 0.005
SHAM	11.4 ± 4.3	10.1 ± 4	10.7 ± 5.1	11.5 ± 4.5	13.2 ± 4.4	Condition x time	0.69
HR (BPM)							
4x5	168 ± 11	158 ± 12	168 ± 11	170 ± 10	173 ± 10	Condition	0.45
8x5	167 ± 13	158 ± 15	166 ± 13	170 ± 13	173 ± 13	Time	< 0.005
SHAM	166 ± 14	154 ± 15	165 ± 14	168 ± 14	171 ± 14	Condition x time	0.96
RPE (Borg scale 6-21)							
4x5	17.7 ± 1.1	16.1 ± 1.6	17.2 ± 1.5	17.7 ± 1.3	19.3 ± 0.9	Condition	0.83
8x5	17.7 ± 1.1	16.6 ± 1.1	17.3 ± 1.3	17.9 ± 1.2	18.8 ± 1.1	Time	< 0.005
SHAM	17.6 ± 1	16.2 ± 1.3	17.1 ± 1.2	17.8 ± 1.4	19 ± 0.9	Condition x time	0.64
V O2 (ml.kg.min ⁻¹)							
4x5	52.6 ± 4.4	49.8 ± 3.3	54.6 ± 4.8	53.2 ± 5.3	52.8 ± 4.7	Condition	0.08
8x5	52.8 ± 4.3	50.3 ± 3.6	54.8 ± 4.7	53.6 ± 5.3	52.8 ± 4.7	Time	< 0.005
SHAM	53.3 ± 4.4	50.4 ± 3.7	55.6 ± 4.7	54.1 ± 4.8	53.3 ± 4.9	Condition x time	0.1

Table 2: The effect of "dose-tissue" on power, heart rate, rate of perceived exertion and $\dot{V}O_2$ following 25%, 50%, 75% and 100% time points during time trial performance.

•			P values				
	Average	0-25%	25-50%	50-75%	75-100%		
Power (Watts)							
BILATERAL	288 ± 33	310 ± 38	284 ± 34	275 ± 34	285 ± 32	Condition	0.43
UNI	285 ± 38	305 ± 45	282 ± 42	275 ± 36	282 ± 36	Time	< 0.005
						Condition x time	0.75
Lactate (mmol.L ⁻¹)							
BILATERAL	11.8 ± 2.8	10.7 ± 3.4	11.5 ± 3.2	12 ± 2.7	13.1 ± 2.9	Condition	0.83
UNI	11.7 ± 3.5	10.9 ± 3.9	11.6 ± 4.1	11.8 ± 3.6	12.9 ± 3.7	Time	0.001
						Condition x time	0.1
HR (BPM)							
BILATERAL	168 ± 11	158 ± 12	168 ± 11	170 ± 10	173 ± 10	Condition	0.21
UNI	168 ± 13	158 ± 15	169 ± 13	171 ± 14	173 ± 13	Time	< 0.005
						Condition x time	0.38
RPE (Borg scale 6-21)							
BILATERAL	17.7 ± 1.1	16.1 ± 1.6	17.2 ± 1.5	17.7 ± 1.3	19.3 ± 0.9	Condition	0.44
UNI	17.5 ± 1	16.3 ± 1.2	17.3 ± 1	17.7 ± 1.2	18.9 ± 1	Time	< 0.005
						Condition x time	0.77
V O2 (ml.kg.min ⁻¹)							
BILATERAL	52.6 ± 4.2	49.8 ± 3.4	54.6 ± 4.5	53.2 ± 5	52.8 ± 4.6	Condition	0.26
UNI	52.5 ± 5.6	49 ± 4.5	54 ± 6.2	53.8 ± 6.1	53.3 ± 6	Time	< 0.005
						Condition x time	0.06

Table 3: The effect of "remote" IPC on power, heart rate, rate of perceived exertion and $\dot{V}O_2$ at 25%, 50%, 75% and 100% time points during time trial performance.

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-			P values				
	Average	0-25%	25-50%	50-75%	75-100%		
Power (Watts)							
LOCAL	288 ± 33	310 ± 38	284 ± 34	275 ± 34	285 ± 32	Condition	0.8
REMOTE	288 ± 35	308 ± 39	286 ± 33	277 ± 35	286 ± 40	Time	< 0.005
						Condition x time	0.94
Lactate (mmol.L ⁻¹)							
LOCAL	11.8 ± 3	10.7 ± 3.4	11.5 ± 3.2	12 ± 2.7	13.1 ± 3	Condition	0.24
REMOTE	11.4 ± 5	9.8 ± 3.9	11.2 ± 4	11.4 ± 4	13.4 ± 6.1	Time	< 0.005
						Condition x time	0.93
HR (BPM)							
LOCAL	168 ± 11	158 ± 12	168 ± 11	170 ± 10	173 ± 10	Condition	0.56
REMOTE	167 ± 14	158 ± 15	168 ± 14	171 ± 13	173 ± 13	Time	< 0.005
					Condition x time	0.41	
RPE (Borg scale 6-21)							
LOCAL	17.7 ± 1.1	16.1 ± 1.6	17.2 ± 1.5	17.7 ± 1.3	19.3 ± 0.9	Condition	0.72
REMOTE	17.6 ± 1.1	16.5 ± 1.2	17.3 ± 1.4	17.6 ± 1.2	19 ± 1	Time	< 0.005
						Condition x time	0.57
V O2 (ml.kg.min ⁻¹)							
LOCAL	52.6 ± 3.8	49.8 ± 3.1	54.6 ± 4.1	53.2 ± 4.6 *	52.8 ± 4.4	Condition	0.04*
REMOTE	53.4 ± 4.3	$.4 \pm 4.3$ 50.4 ± 3.3	55.1 ± 4.6	54.5 ± 5*	53.7 ± 5	Time	< 0.005
						Condition x time	0.36

Table 4: Perceived discomfort of IPC and SHAM interventions.

		Pe	Mean discomfort rating				
_		Average	0-10 min	10-20 min	20-30 min	30-40 min	
	Traditional 4x5 IPC (legs)	3.7 ± 1.2	4.5 ± 1.5	3.5 ± 1.1	3.5 ± 1.1	3.4 ± 1.1	Light to moderate
	Larger 8x5 cycles	3.6 ± 1.7	3.9 ± 1.8	3.5 ± 1.6	3.3 ± 1.8	3.5 ± 1.8	Light to moderate
	Unilateral 4x5 IPC	3.1 ± 1.5	3.5 ± 1.9	3.1 ± 1.5	2.8 ± 1.3	2.8 ± 1.5	Light to moderate
	Remote 4x5 IPC (arms)	3.7 ± 2.1	4.1 ± 2	3.6 ± 2	3.7 ± 2	3.4 ± 2.3	Light to moderate
685	SHAM	0 ± 0	0 ± 0	0 ± 0	0 ± 0	0 ± 0	No discomfort
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687							
688							