

1 **Title: The effects of 10 days of separate heat and hypoxic exposure on heat acclimation and temperate**
2 **exercise performance**

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10 **Running head:** Hypoxia and heat acclimation

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23 **Abstract**

24 Adaptations to heat and hypoxia are typically studied in isolation, but are often encountered in combination.
25 Whether the adaptive response to multiple stressors affords the same response as when examined in isolation
26 is unclear. We examined: i) the influence of overnight moderate normobaric hypoxia on the time course and
27 magnitude of adaption to daily heat exposure; ii) whether heat acclimation (HA) was ergogenic and if this
28 was influenced by an additional hypoxic-stimulus. Eight males ($\dot{V}O_{2\max}=58.5[8.3]$ mL \cdot kg $^{-1}\cdot$ min $^{-1}$) undertook
29 two 11-day HA programmes (balanced-crossover design), once with overnight normobaric hypoxia (8[1] h
30 per night; 10 nights; $F_{I}O_2=0.156$; $S_pO_2=91[2]\%$ [HA_{Hyp}]) and once without (HA_{Con}). Days 1, 6, 11 were
31 exercise-heat stress tests (HST [40°C, 50% RH]); days 2-5 and 7-10 were isothermal-strain (target rectal
32 temperature [T_{re}] \sim 38.5°C), exercise-heat sessions. A graded exercise test and 30-minute cycle trial were
33 undertaken pre, post and 14-days after HA in temperate-normoxia (22°C, 55% RH; $F_{I}O_2=0.209$). HA was
34 evident on day 6 (*e.g.* reduced T_{re} , mean skin temperature [\bar{T}_{sk}], heart rate, sweat [Na^+], $P<0.05$) with
35 additional adaptations on day 11 (further reduced \bar{T}_{sk} , heart rate). HA increased plasma volume (+5.9[7.3]%)
36 and erythropoietin concentration (+1.8[2.4] mIU/mL); tHb_{mass} was unchanged. Peak power output (+12[20]
37 W), lactate threshold (+15[18] W) and work done (+12[20] kJ) increased following HA. The additional
38 hypoxic-stressor did not affect these adaptations. In conclusion, a separate moderate overnight normobaric
39 hypoxic-stimulus does not affect the time-course or magnitude of HA. Performance may be improved in
40 temperate-normoxia following HA, but this is unaffected by an additional hypoxic stressor.

41

42 **Key words (×3-5)**

43 Thermoregulation; Acclimatization; Altitude; Training; Combined-stress

44

45 **Introduction**

46 Historically, adaptation to environmental stressors has been examined in isolation, yet multiple
47 environmental stressors can be encountered in the natural world, either simultaneously or in close proximity,
48 for instance, heat or cold *and* hypoxia [78]. It cannot be assumed that the adaptive response to multiple
49 stressors affords the same response as when examined in isolation and it has recently been highlighted that
50 three broad types of interaction (additive, synergistic, antagonistic) can occur when combining independent
51 stressors [46]. Consequently, there is a need to better understand adaptations to multiple stressors [78].

52 Heat acclimation (HA) occurs when core (T_c) and skin temperature (T_{sk}) are frequently and repeatedly
53 elevated to a level challenging thermoeffector responses, commonly as a consequence of exercise-heat stress
54 [e.g. 45, 61]. At a systemic level, plasma volume (PV) expansion occurs within ~3 days [74], the resulting
55 hypervolemia increases stroke volume, maximal cardiac output [48], and arterial blood pressure [56], and
56 lowers heart rate for a given work-rate [45, 63]. PV expansion also increases the total specific heat capacity
57 of blood [7], aiding core-skin heat transfer and reducing cutaneous blood-flow requirements [62]. Sudomotor
58 changes (lower threshold and greater sweating sensitivity) are complete after ~10 days [62]. Together these
59 adaptations improve cardiovascular stability [74] and reduce thermal strain (lower T_{sk} and T_c , [21]. There is
60 also evidence of metabolic adaptation, characterized by reduced reliance on carbohydrate metabolism [83]
61 and lower exercise muscle and blood lactate accumulation [48]. At a cellular level, heat exposure activates
62 the heat shock response [42], increasing heat shock protein (HSP70 and HSP90) concentration; these
63 proteins are multi-functional, but are primarily cytoprotective [41, 64]. However, heat exposure may also
64 stimulate the hypoxia inducible factor-1 pathway [4, 44], which primarily controls oxygen-related genes.

65 Systemic adaptations to hypoxia develop within ~7 to 21 days of living at high-altitude (1,500 m-3,500 m) or
66 intermittent hypoxic exposure [19]. Stimulation of aortic-arch chemoreceptors and carotid bodies increases
67 sympatho-adrenal activity, elevating heart rate, cardiac output, and ventilation [81]. In the early stages of
68 acclimation PV decreases due to diuresis [40] and possibly extra- to intra-cellular fluid shifts [27]. The
69 resultant hypovolemia causes hemoconcentration, increasing oxygen carrying capacity per unit volume [82]
70 and reducing heart rate and cardiac output for a given oxygen demand. Together these effects improve tissue
71 oxygen delivery. With chronic hypoxia, erythropoiesis increases erythrocyte volume (EV) [22], although

72 reticulocytosis occurs more rapidly [20] and changes in EV may present after removal of the hypoxic
73 stimulus. Metabolically, adaptations to hypoxia may increase reliance on carbohydrate for ATP resynthesis
74 [59] whereas at the cellular-level, hypoxic stress primarily activates the HIF-1 pathway [73], which
75 stimulates a cascade of effects including erythropoiesis, but also induces the heat shock response [42].

76 Although recent studies have examined the cross-acclimation (attenuated physiological-strain) or cross-
77 tolerance (improved cellular protection) afforded by adaption to heat during subsequent hypoxic exposure
78 [24, 44], the effect of the addition of a hypoxic-stressor on the adaptive response to heat *i.e.* a combined-
79 stressor approach, has received little attention. Although mechanistically important, this question is also
80 practically relevant; athletes often sleep in hypoxic environments (*i.e.* hypoxic tents/nitrogen houses) to try
81 and gain an ergogenic benefit [8], whilst at the same time they may undergo HA prior to competition in a hot
82 environment. Likewise, high ambient temperatures may be encountered at popular high-altitude training
83 venues *e.g.* Colorado (up to 40°C and ~2,000 m). It has been hypothesised that the impact of individual
84 stressors on exercise capacity dictates the interaction; mild stressors producing an additive effect, with a
85 move towards antagonistic interactions as the individual stressors impact increases [46]. Thus, addition of a
86 modest hypoxic stimulus might be hypothesised to potentiate HA. Alternatively it has been suggested that
87 additive effects result from combining stressors with independent mechanisms, whilst interactive effects
88 arise from mechanistically similar stressors [47]. Although there are clearly independent mechanisms by
89 which heat and hypoxic stress elicit adaptation, there are also potential synergies in aspects of the cellular
90 (*e.g.* heat shock response and HIF-1), and systemic (*e.g.* reduced sub-maximal exercise heart rate, improved
91 tissue oxygen delivery) adaptive responses. However, antagonistic effects are also possible; PV is expanded
92 with HA [74], but reduced with hypoxia [68], whereas HA may reduce reliance on glycolysis [37], but this
93 may be increased with hypoxia [29].

94 An ancillary question which we sought to investigate was whether HA was ergogenic in temperate
95 conditions, and if this was influenced by the addition of hypoxia, *i.e.* a cross-stressor effect between
96 adaptation to heat-hypoxia and performance in temperate-normoxia. Although the ergogenic benefit of
97 hypoxia for endurance exercise is well established [8], the ergogenic potential of HA for prolonged exercise
98 has recently received increased attention (*e.g.* [15]). Although HA could be ergogenic via multiple
99 mechanisms [15] it is suggested that PV expansion is primary among these, due to its positive effect on

100 cardiac output and $\dot{V}O_{2\max}$ [48]. However, other studies have shown no ergogenic effect of HA induced PV
101 expansion [33, 36], possibly due to a hemodilution effect sufficient to offset any increase in cardiac output
102 [16]. Currently it is unclear if the addition of the erythropoietic stimulus of hypoxia is sufficient to offset the
103 hemodilution effect of HA, or whether hypoxia negates normal PV expansion with HA. Although Takeno *et*
104 *al.* [76] demonstrated increased PV, EV and $\dot{V}O_{2\text{peak}}$ with 10-daily exercise bouts (60 min·day⁻¹) in hot,
105 (30°C, 50% RH) hypobaric hypoxic (2,000 m), conditions, these data are limited by the small sample ($n=5$)
106 and similar adaptations were evident in a cool-normoxic control group, indicating a possible training-effect.
107 Likewise, Buchheit *et al.* [11] reported PV expansion in both normobaric hypoxic ($F_{I}O_2 \sim 0.150$; 14 ± 1
108 h·day⁻¹) and normoxic two-week HA programme (~27 h total heat exposure, ~32°C, 39% RH) although total
109 hemoglobin mass (tHb_{mass}) was increased in the hypoxic condition only. However, these hematological
110 changes were not related to the temperate-normoxic performance improvement following both regimens.
111 More recently, McCleave *et al.* [50] showed a 3.3% improvement in temperate-normoxic 3 km running trial
112 performance three weeks (but not immediately) after completing a 21-day intermittent HA programme.
113 However, the ergogenic effect was absent when normobaric hypoxia was added to the HA programme
114 ($F_{I}O_2=0.144$; 13 h·day⁻¹) and although tHb_{mass} did increase with the additional hypoxic stressor, PV
115 expansion was 'possibly less' and the hematological changes were not related to the performance effects.

116 Accordingly, the aims of the present study were two-fold. First, to examine the addition of a daily hypoxic
117 stimulus on the time course and magnitude of adaption to heat and second, to investigate whether HA was
118 ergogenic under temperate-normoxic conditions, and if this was influenced by the addition of a daily hypoxic
119 stimulus. Our null hypotheses were that the addition of a moderate daily hypoxic stimulus would not affect
120 the time course or magnitude of HA, and would not influence any effect of HA on temperate-normoxic
121 exercise performance.

122

123 **Materials & Methods**

124 **Participants**

125 Sample size was calculated *a priori* using G*Power software; effect size data were derived from the change
126 in exercise T_{re} ($\eta^2=0.16$) observed following an identical HA programme (without hypoxia) in our laboratory

127 [55]. For two-way (Condition \times Time) repeated measures analysis of variance with sufficient power ($\beta \geq 0.80$)
128 at an α level of 0.05 a minimum of eight participants was required. Similar sample-size estimates were
129 obtained with effect-size data derived from other key outcome variables, including mean body temperature
130 (\bar{T}_b) and heart rate. To account for attrition 12 male participants were recruited; four did not complete the
131 study due to injury (unrelated to study, $n=1$), illness ($n=1$) and logistics ($n=2$). Eight performance level three
132 [17] males (Age: 25[6] years; $\dot{V}O_{2\max}$: 58.6[8.9] mL \cdot min $^{-1}\cdot$ kg $^{-1}$; peak power output: 348[53] W) completed
133 this study. Participants were all trained endurance athletes (cyclists/triathletes/runners). The study was
134 approved by the University's Ethics Committee and conformed to the Declaration of Helsinki, and all
135 participants provided written informed consent.

136 **Experimental design**

137 A within-participant, balanced cross-over design was employed, with participants undertaking both control
138 (heat acclimation [HA_{Con}]) and experimental (heat acclimation with hypoxic exposure [HA_{Hyp}]) HA
139 programmes. Each HA programme lasted 11-days and consisted of three bouts of exercise at a fixed external
140 work rate (heat stress test [HST]), undertaken on day 1 (HST_{pre}), day 6 (HST_{mid}) and day 11 (HST_{post}),
141 interspersed with eight isothermal heat strain exercise-heat exposures (ISO). A temperate graded exercise
142 test (GXT) and 30 minute work done trial (T30) were performed before (GXT_{pre}; T30_{pre}) and after (GXT_{post};
143 T30_{post}) each HA programme; an additional retention T30 was undertaken 14-days after completing HA
144 (T30_{ret}) (

145

146 Figure 1). HA programmes were identical apart from the addition of daily (overnight) normobaric hypoxic
147 exposure in HA_{Hyp} . A minimum three-month wash-out period was prescribed between HA programmes [14]
148 and all testing was completed outside of the UK summertime (average weather conditions: 8.7°C, 77% RH).

149

INSERT FIGURE 1 HERE

150 **Experimental procedures**

151 *Graded Exercise Test*

152 GXTs were performed in a temperate environment (22°C, 50% RH) (pre- and post-HA_{Con} and HA_{Hyp}) on a
153 Lode Excalibur cycle ergometer (Lode B.V. Groningen, the Netherlands). Participants exercised for 20
154 minutes at 85 or 110 W, dependent upon the estimated fitness of the participant (fixed within-participant for
155 pre-post tests and between-conditions). Thereafter, work-rate was incremented by 25 W every three minutes
156 until blood lactate concentration [Lac] was ≥ 4 mmol·L⁻¹, following which, the participant was given a five
157 minute break before beginning cycling again at 100 W for five minutes. Work-rate was then increased 25
158 W·min⁻¹ until volitional exhaustion. [Lac] was determined from fingertip capillary blood obtained at the end
159 of each exercise stage (Biosen C-line, EKF Diagnostic, Cardiff, UK). Convective cooling was provided at a
160 rate of 3.5 m·s⁻¹.

161 ***30 Minute maximal cycling trial***

162 T30s were conducted to obtain an index of endurance performance. All trials were performed on a Lode
163 Excalibur cycle ergometer (Lode B.V. Groningen, the Netherlands) in a temperate environment (22°C, 50%
164 RH). After a standardized warm up participants commenced a 30 minute ‘all-out’ performance trial;
165 ‘performance’ was defined as the total work done (kJ). A fan provided some convective cooling (3.5 m·s⁻¹)
166 to reduce the likelihood of having to end the test early due to reaching withdrawal criteria for T_{re} of 40°C.

167 ***Heat Stress Test (HST)***

168 HSTs were completed pre-, mid- and post-HA in both conditions as described previously [54, 55]. Briefly,
169 participants cycled in a hot environment (target ambient conditions: 40°C; 50% RH) on a calibrated
170 COMPUTRAINER™ cycle ergometer (RacerMate Inc., Seattle, WA, USA) for 60 minutes at 35% of peak
171 power output (PPO) reached in the pre-HA GXT. 1.25 L of 3.6% carbohydrate solution (Science in Sport Go
172 Electrolyte drink, Nelson, UK) (drink temperature 20°C) was ingested to replace fluid losses, divided into
173 five equal boluses (0.25 L) and consumed immediately prior to commencing exercise and every 15 minutes
174 thereafter. Convective cooling was provided at a rate of 3.5 m·s⁻¹; this prevented participants from reaching
175 the T_{re} withdrawal criteria, whilst maintaining an acceptably high mean skin temperature (\bar{T}_{sk}) and allowing
176 thermoeffector responses to be assessed.

177 ***Isothermal heat strain sessions (ISO)***

178 Participants exercised in a hot environment (target ambient conditions: 40°C; 50% RH) on a calibrated
179 COMPUTRAINER™ cycle ergometer (RacerMate Inc., Seattle, WA, USA), initially selecting a work rate
180 eliciting a rating of perceived exertion (RPE [9]) of 15. This was maintained until T_{re} reached 38.5°C, at
181 which point external power output was adjusted as appropriate to maintain this target temperature ($\pm 0.2^\circ\text{C}$)
182 and a small amount of convective cooling ($3 \text{ m}\cdot\text{s}^{-1}$) was used to facilitate the exercise component and provide
183 some perceptual benefit, whilst maintaining a high T_{sk} . Participants completed eight 90 minute ISO sessions
184 in both the HA_{Con} and the HA_{Hyp} condition and were provided with fluid replacement ($7 \times 0.25 \text{ L}$, 3.6%
185 carbohydrate, boluses every 15 minutes during ISO sessions).

186 *Hypoxic exposure*

187 During the HA programme participants in the HA_{Hyp} condition were exposed to nightly moderate normobaric
188 hypoxia (10 nights, 8-10 h exposure per night, $F_{I\text{O}_2} = 0.156$) comparable to a simulated altitude of ~2,400 m,
189 using ‘portable altitude tents’ (Hypoxico, New York City, New York, USA). This hypoxic stimulus exceeds
190 the threshold required for erythropoiesis in humans [53], is consistent with the hypoxic stimulus used in
191 previous studies [11, 76] and is similar to the altitude of many popular training camp locations e.g. Flagstaff
192 AZ., USA (2,106 m); Sierra Nevada, Spain (2,320 m); Iten, Kenya (2,400 m). Although the hypoxic and
193 heat stimuli were not delivered simultaneously, as might occur with residing at a high altitude training camp,
194 some individuals (athletes) may live or sleep in a hypoxic environment and undertake their training in a
195 normoxic (hot) environment Participants were familiarized with sleeping in the tents (without a reduced PO_2)
196 for several nights prior to commencing HA_{Hyp} to become accustomed to any changes in ambient noise and
197 minimize sleep disturbances. Participants wore a physiological monitoring system (EQUIVITAL™ ,
198 Cambridge, UK) which recorded heart rate (EQO₂ LifeMonitor, EQUIVITAL™, Cambridge, UK) and
199 oxygen saturation (Nonin iPod SpO₂, EQUIVITAL™, Cambridge, UK) (sampling every 15 seconds, and for
200 two minutes every 10 minutes, respectively) throughout each of the 10-nights.

201 *General procedures*

202 Participants wore the same clothes on each day, abstained from alcohol throughout the experimental periods
203 or caffeine for 12 hours prior to exercise, consumed a similar diet before each test and drank 0.5 L of water
204 two hours before every attendance. Participants were instructed to maintain their normal high-intensity

205 training (except 24 h before HSTs, GXTs, T30s) and replace an equivalent duration of low/moderate training
206 with that completed in the laboratory to maintain usual training volume. Additionally, participants recorded
207 the number of hours spent in the tent and the evening and morning F_{iO_2} (independent reading taken with a
208 calibrated VN202 mkII oxygen analyser, Vandagraph Ltd, Keightly, UK) within the tent each night.

209 To monitor daily hydration status, urine osmolality was assessed prior to exercise (Osmometer 3320,
210 Advanced Instruments Inc., Norwood, MA, USA). Nude body mass (dry) was measured pre- and post- each
211 test session (Industrial Electronic Weight Indicator, Model I10, Ohaus Corporation, Parsippany, NJ, USA);
212 body mass changes were used to determine whole-body sweat rate, adjusted for fluid ingested. Ambient
213 conditions were measured by a WBGT logger (Squirrel 1000, Grant Instruments, Cambridge, UK), T_{re} by a
214 thermistor (Grant Instruments, Cambridge, UK) self-inserted 15 cm beyond the anal sphincter and cardiac
215 frequency (f_c) by short-range telemetry (Polar RS800, Polar Electro, Kempele, Finland). During HSTs and
216 GXTs skin temperature (T_{sk}) was measured using thermistors on the chest, biceps, thigh and calf (Grant
217 Instruments, Cambridge, UK) and local sweat rate at the upper right back (Q-Sweat, WR Medical
218 Electronics, Maplewood, MN, USA) and forearm skin blood flow (MoorLAB, Moor Instruments, Devon,
219 UK) were recorded. During HSTs expired gases (Douglas bag method), RPE [9], thermal sensation [84] and
220 thermal comfort [85] were measured at 15 min intervals. A sample of sweat was collected using a custom
221 patch constructed from TEGADERM™ (TEGADERM™ Dressings, 3M, St. Paul, Minnesota, USA) and
222 PARAFILM® (Bemis NA, Neenah, WI, USA) for determining sodium concentration [Na^+] by flame
223 photometry (Flame Photometer 410, Sherwood Scientific Ltd, Cambridge, UK). During GXTs oxygen
224 uptake was measured breath-by-breath throughout (Quark B2, Cosmed, Rome, Italy).

225 *Hematological procedures*

226 Immediately before and after ISO1 and prior to HSTs a 10 mL venous blood sample was obtained (K2 EDTA
227 blood collection tubes, Beckton Dickson & Company, Plymouth, UK) from the antecubital vein following 15
228 min of seated rest. Whole blood samples were centrifuged (1500 g for 15 min at 4°C, HERAEUS™
229 MULTIFUGE™ 3 S-R, Thermo Electron Corporation, Karlsruhe, Germany) and 20 μ L of the resultant
230 plasma was assessed for osmolality (Osmometer 3320, Advanced Instruments Inc., Norwood, MA, USA)
231 and the remainder aliquoted and stored at -80°C for subsequent biochemical analyses using enzyme linked

232 immunosorbent assays (ELISA). Resting tHb_{mass} , (CV=4.2%), blood volume (BV) (CV=3.4%) and PV
233 (CV=4.4%) were determined using the optimised carbon monoxide rebreathing technique [68] with a 1.0
234 $mL \cdot kg^{-1}$ body mass CO bolus [79], the day before and after the HA programmes, and 14-days after
235 completion of HA. Fingertip capillary samples were taken in triplicate during the CO rebreathing technique
236 to assess the percentage of carboxyhemoglobin (ABL80 CO-OX Flex Hemoximeter, RADIOMETER™,
237 Copenhagen, Denmark) in the blood. Venous blood samples were also collected to determine hemoglobin
238 concentration [Hb] (201⁺ HEMOCUE®, Ängelholm, Sweden) and hematocrit (Hct) (Hawksley, Lancing,
239 UK) in triplicate. Together, these were used to determine tHb_{mass} , PV and BV, before and after the HA
240 programmes, due to potential for a change in red cells which is not accounted for in the Dill & Costill [18]
241 method.

242 **Data analyses**

243 \bar{T}_{sk} was calculated according to Ramanathan [59] and \bar{T}_b as the weighted mean of T_{re} (0.9) and \bar{T}_{sk} (0.1)
244 according to Jay *et al.* [30]. For GXT data the lactate threshold was defined as the power output at [Lac] of 4
245 $mmol \cdot L^{-1}$, gross mechanical efficiency was calculated at 185 W (highest work rate below lactate threshold
246 achieved by all participants), and $\dot{V}O_{2max}$ was defined as the highest 15 s $\dot{V}O_2$. Physiological strain index
247 (PSI) was determined according to Moran *et al.* [52] and metabolic heat production (MHP) was calculated
248 according to ISO 8996 Malchaire [49].

249 Extracellular HIF-1 α and erythropoietin (EPO) concentration, in EDTA plasma, were measured using
250 colorimetric sandwich ELISAs (Thermo Fisher Scientific, Waltham, MA, USA, and; Abcam, Cambridge,
251 UK, respectively) and read at 450 nm (450 and 550 nm for EPO) on a plate reader (SPECTRAMAX® i3x,
252 Molecular Devices, Wokingham, UK) with SOFTMAX® Pro (version 6.5.1, Molecular Devices,
253 Wokingham, UK). Results were calculated using the standard curve and the average absorbencies from
254 samples in duplicate. The HIF-1 α assay's detection range was 81.92-20,000 pg/mL and limit of detection
255 was <30 pg/mL. The intra-assay precision was determined from duplicates of standards/controls within the
256 same plate (3.2%) and inter-assay precision determined from standards/controls assessed across plates
257 (8.7%). The EPO assays' detection range was 1.6-100 mIU/mL and had a sensitivity of 0.17 mIU/mL, with

258 an intra-assay precision of 8.0% and an inter-assay precision of 8.6%. Pre-post programme changes in both
259 conditions were assessed on the same plate for each individual.

260 *Statistical analyses*

261 Statistical analyses were undertaken using SPSS (IBM Version 22, IBM, New York, NY, USA).
262 Significance was set *a-priori* at $P \leq 0.05$; data are presented mean(SD) unless otherwise stated. Following
263 Shapiro-Wilk tests for normality, two-way repeated measures ANOVA were used to analyze the main
264 effects, *i.e.* responses over Time (HST: pre/mid/post; GXT and T30: pre/post/ret; ISO: 1-8) and Condition
265 (HA_{Con} vs. HA_{Hyp}), as well as the interaction effect (*i.e.* Time \times Condition). Effect sizes are presented using
266 eta squared (η^2 , calculated as the sum of squares for an effect/total sum of squares) for ANOVAs (η^2
267 ≤ 0.02 =small; 0.02-0.13=medium; 0.13-0.26=large effect size). The Huynh-Feldt statistic was employed to
268 account for violations of sphericity; Bonferroni adjusted Students *t*-tests were used *post-hoc* for analysis of
269 main and interaction effects. *Post-hoc* analysis of significant time effects for ISO sessions were made
270 relative to ISO1 only, with alpha adjusted accordingly. A one-way ANOVA was used to assess changes in
271 the daily degree of hypoxic strain, as indicated by overnight oxy-hemoglobin saturation during the HA_{Hyp}
272 condition. Non-parametric tests (Friedman's test for change over time and Wilcoxon signed ranks tests for
273 condition effects at each time point) were used to assess ordinal (RPE) data. Correlations were assessed
274 using Pearson's *r* for parametric data and Spearman's rank comparisons for non-parametric data.

275

276 **Results**

277 **Daily heat and hypoxic exposure**

278 Ambient conditions during ISOs did not differ between conditions (39.6[0.3] $^{\circ}$ C, 53.3[4.1]% RH, $P > 0.05$).
279 Participants sustained a mean power of 105(16) W (not different between conditions, $F_{(1,7)} = 0.071$, $P = 0.797$,
280 $\eta^2 < 0.01$) with a 5 minute peak power of 189(40) W (not different between conditions, $F_{(1,7)} = 0.379$, $P = 0.558$,
281 $\eta^2 < 0.01$). A T_{re} of 38.5 $^{\circ}$ C was achieved in 31(11) mins (not different between conditions $F_{(1,7)} = 0.698$,
282 $P = 0.431$, $\eta^2 = 0.02$) and the average T_{re} for the final 60 minute of each ISO was 38.52(0.17) $^{\circ}$ C. Power output
283 increased over the eight ISO sessions ($F_{(4.4,30.6)} = 2.823$, $P = 0.038$, $\eta^2 = 0.08$) but this did not differ between

284 conditions ($F_{(1,7)}=0.071$, $P=0.797$, $\eta^2=0.02$). Whole-body sweat rate was increased over time
285 ($F_{(4,0,28,2)}=18.038$, $P<0.001$, $\eta^2=0.12$) and also differed between conditions ($F_{(1,7)}=15.278$, $P=0.006$, $\eta^2=0.01$)
286 although the location of differences could not be located *post-hoc*. Pre-exercise urine osmolality was higher
287 in the HA_{Hyp} condition compared to the HA_{Con} condition ($F_{(1,7)}=11.142$, $P=0.012$, $\eta^2=0.05$) with significant
288 differences between conditions evident on ISO6 only ($P=0.024$); urine osmolality did not change over the
289 course of HA ($F_{(7,49)}=0.223$, $P=0.978$, $\eta^2=0.01$). An interaction effect was evident for pre-exercise mass
290 ($F_{(7,49)}=3.316$, $P=0.006$, $\eta^2<0.01$) which increased over time in the HA_{Con} condition and decreased in the
291 HA_{Hyp} condition, although *post-hoc* comparisons could not locate these differences (Table 1). The overnight
292 hypoxia ($F_1O_2 = 0.156(0.008)$) during HA_{Hyp} was sustained for 8(1) hrs on 10 consecutive nights and elicited
293 an average S_pO_2 of 91(2)% (Table 2).

294 INSERT TABLE 1 HERE

295 INSERT TABLE 2 HERE

296 Heat acclimation

297 Ambient conditions did not differ between the HSTs (39.4(0.5)°C, 50.5(1.6)% RH, $P>0.05$) and metabolic
298 heat production (8.1(0.8) W·kg⁻¹) did not differ throughout HSTs (main effect of time: $F_{(2,14)}=0.465$,
299 $P=0.637$, $\eta^2=0.01$) or between conditions ($F_{(1,7)}=3.426$, $P=0.107$, $\eta^2=0.06$).

300 Both HA protocols successfully induced HA, with a number of thermophysiological adaptations evident at
301 HST_{mid} and some further adaptations developing by HST_{post} (Figure 2 and Supplemental Table 1). However,
302 the addition of nightly hypoxic exposure to the regimen did not affect HA; no significant interaction effects
303 were observed for parameters measured in the HST (Figure 2 and Supplemental Table 1). Although end
304 exercise f_c recorded in each HST was significantly greater in the HA_{Hyp} condition than then HA_{Con} condition
305 (main effect for condition: $F_{(1,7)}=13.656$, $P=0.008$, $\eta^2=0.06$), Bonferroni corrected *post-hoc t*-tests comparing
306 conditions at each time point could not locate specific differences. No other condition effects were evident.

307 INSERT FIGURE 2 HERE

308 Two participants were unable to complete the retention period hematological tests, therefore data in the 3 × 2
309 (Time × Condition) ANOVA are for $n=6$. tHb_{mass} was unchanged over time ($F_{(2,10)}=2.275$, $P=0.153$, $\eta^2=0.03$)

310 and condition ($F_{(1,5)}=0.852$, $P=0.398$, $\eta^2=0.01$) and there were no interaction effects ($F_{(2,10)}=0.263$, $P=0.774$,
311 $\eta^2=0.01$) (**Error! Reference source not found.3**). On the other hand, PV ($F_{(2,10)}=8.974$, $P=0.006$, $\eta^2=0.10$)
312 and BV ($F_{(2,10)}=8.678$, $P=0.007$, $\eta^2=0.10$) changed over time; *post-hoc* comparisons identified a significant
313 decrease from post to retention time points (PV: -8.9[5.2]% ($P=0.015$); BV: -6.2[4.4%] ($P=0.027$)), but the
314 pre-HA and retention PV and BV values were not different. PV and BV were also unchanged between
315 conditions and there were no interaction effects (Table 3). To account for the reduced participant number and
316 increased potential for type II error, we undertook a further analysis (*i.e.* a 2×2 repeated measures
317 ANOVA), for the time points where $n=8$ (*i.e.* HA_{pre} vs. HA_{post}); with this further analysis both PV
318 (+5.9(7.3)%, $F_{(1,7)}=10.981$, $P=0.013$, $\eta^2=0.07$) and BV (+3.5(5.9)%, $F_{(1,7)}=10.083$, $P=0.016$, $\eta^2=0.05$) were
319 expanded pre to post-HA, but there were no condition or interaction effects.

320 The concentration of plasma EPO (pre-exercise in HST) was increased over time with HA ($F_{(1,7)}=6.646$,
321 $P=0.037$, $\eta^2=0.06$), *post-hoc* analysis indicated that the increase was significant from HST_{pre} (8.3(3.6)
322 mIU/mL) to HST_{post} (10.1(3.9) mIU/mL). There was no difference between conditions ($F_{(1,7)}=0.273$,
323 $P=0.618$, $\eta^2<0.01$) or interaction effect ($F_{(1,7)}=0.005$, $P=0.948$, $\eta^2<0.01$) (Supplemental Table 1). EPO
324 concentration did not differ following a single bout of overnight hypoxia compared to normoxic exposure
325 ($t_{(7)}=0.041$, $P=0.968$, $d=0.02$). HIF-1 α was largely undetectable in the plasma at these time points.

326 INSERT TABLE 3 HERE

327 **Temperate exercise performance following HA**

328 *Graded exercise test*

329 Data from the GXTs are shown in Figure 3. No interaction (Time \times Condition) effects were reported for the
330 parameters measured ($\dot{V}O_{2max}$, PPO, LT, GME, maximal heart rate) in the temperate GXT completed
331 immediately before and after each HA programme, although a condition effect was detected for PPO
332 ($F_{(1,7)}=9.632$, $P=0.017$, $\eta^2=0.05$), *post-hoc* analysis indicated that this was partly due to a higher baseline
333 PPO in the HA_{Hyp} condition (359(48) W) than the HA_{Con} condition (342(48) W) ($P=0.048$) as well as
334 following HA (HA_{Hyp}: 373(38) W; HA_{Con}: 353(30) W; $P=0.021$). PPO and lactate threshold ($F_{(1,7)}=11.700$,
335 $P=0.011$, $\eta^2=0.02$) were improved over time (+12(20) W and +15(18) W, respectively) and f_{Cmax} was reduced
336 (-5(5) $b \cdot \min^{-1}$, $F_{(1,7)}=37.840$, $P=0.001$, $\eta^2=0.17$) following the medium-term HA, but GME remained

337 unchanged with time ($F_{(1,7)}=1.189$, $P=0.312$, $\eta^2=0.03$) or condition ($F_{(1,7)}=0.394$, $P=0.550$, $\eta^2=0.02$). Results
338 for $\dot{V}O_{2\max}$ showed different effects depending on whether oxygen uptake was in relative or absolute terms;
339 relative $\dot{V}O_{2\max}$ was unchanged with time ($F_{(1,7)}=0.913$, $P=0.371$, $\eta^2=0.01$) or condition ($F_{(1,7)}=4.641$,
340 $P=0.068$, $\eta^2=0.02$). On the other hand, a main effect for condition was reported for absolute $\dot{V}O_{2\max}$
341 ($F_{(1,7)}=6.735$, $P=0.036$, $\eta^2=0.04$); *post-hoc* tests indicated a trend ($P=0.094$) for a higher $\dot{V}O_{2\max}$ at baseline in
342 the HA_{Hyp} (4.36(0.62) L·min⁻¹) condition than the HA_{Con} condition (4.13(0.48) L·min⁻¹), but there was not a
343 main effect over time ($F_{(1,7)}=0.808$, $P=0.399$, $\eta^2=0.01$).

344

INSERT FIGURE 3 HERE

345 **30 minute work done trial (T30)**

346 Environmental conditions for the T30 were matched between conditions and over time: 22.1(0.2)°C,
347 52.5(3.0)% RH). Data from the T30 are shown in Figure 4. Two participants in the HA_{Hyp} condition did not
348 complete the retention trial therefore $n=6$ in the 3 (Time) \times 2 (Condition) repeated measures ANOVA. Work
349 done was not different between conditions ($F_{(1,5)}=3.341$, $P=0.127$, $\eta^2=0.02$) and there was no interaction
350 effect ($F_{(2,10)}=0.505$, $P=0.618$, $\eta^2<0.01$) but it was changed over time ($F_{(2,10)}=5.283$, $P=0.028$, $\eta^2<0.01$).
351 Although *post-hoc* comparisons could not locate these differences. We undertook a further analysis (*i.e.* a 2
352 \times 2 repeated measures ANOVA), for the time points where $n=8$ in both conditions (*i.e.* T30_{pre} and T30_{post}),
353 which indicated that work done was improved by +12(20) kJ ($F_{(1,7)}=5.939$, $P=0.045$, $\eta^2=0.01$) immediately
354 following HA. There were no significant differences between conditions ($F_{(1,7)}=4.102$, $P=0.082$, $\eta^2=0.03$)
355 and there was no interaction effect ($F_{(1,7)}=0.036$, $P=0.854$, $\eta^2<0.01$). The improvement in work done was not
356 correlated with the increased LT ($r_{(16)}=0.088$, $P=0.746$) or PPO ($r_{(16)}=0.476$, $P=0.062$).

357

INSERT FIGURE 4 HERE

358

359 **Discussion**

360 This study was the first to examine the effect of adding a moderate overnight hypoxic stimulus on the time
361 course and magnitude of adaption to heat, with an ancillary aim of investigating the ergogenic potential of
362 combined adaptation to heat and hypoxia on exercise performance in a temperate, normoxic environment.

363 The main finding of the present study was that the addition of 80(8) hours normobaric hypoxia did not alter
364 the rate or magnitude of the development of HA, as indicated by key thermophysiological and hematological
365 indices; regardless of the intervention condition some HA was acquired with short-term heat exposure
366 (totaling seven hours over five-days), with a more pronounced heat-acclimated phenotype evident following
367 medium-term heat exposure (totaling 14 hours over 10-days). Furthermore, although there was evidence
368 supporting an ergogenic effect of HA under temperate-normoxic conditions (improved lactate threshold,
369 PPO and work done), this was not affected by the addition of normobaric hypoxia, which did not notably
370 affect the hematological adaptations to HA.

371 Importantly, for our experimental model, thermal-strain, cardiovascular-strain and external work-rate were
372 matched between the HA_{Con} and HA_{Hyp} conditions, whereas oxy-hemoglobin saturation was significantly
373 reduced overnight in HA_{Hyp}. Moreover, the degree of thermal strain experienced by the participants was
374 sufficient to exceed the adaptation threshold [77]; reduced T_{re} , \bar{T}_{sk} , \bar{T}_b , f_c and sweat $[Na^+]$ and augmented
375 sweat rate were evident within five days of HA, with a more developed heat acclimated phenotype
376 (expansion of PV and BV, further reduced \bar{T}_{sk} and f_c) evident after 10-days of HA. Whilst a pronounced
377 adaptive response was evident within five days, the observation that a longer term HA regimen is superior to
378 a shorter regimen is in keeping with a recent meta-analysis [80], whereas the finding that the time-course and
379 magnitude of the adaptive response to heat was unaffected by the addition of 80(8) hours of moderate
380 normobaric hypoxia is novel, although there are some relevant comparison data. For instance, Buchheit *et al.*
381 [11] demonstrated similar reductions in f_c and sweat $[Na^+]$ following a 14-day warm-weather training camp,
382 which was unaffected by the addition of a hypoxic stressor (170 h, $F_{I}O_{2\sim} \sim 0.15$), but no measures of body
383 temperature were reported. However, Takeno *et al.* [76] reported reduced esophageal temperature and
384 exercising f_c following 10 ($1 \text{ h} \cdot \text{day}^{-1}$) exercise-heat (30°C , 50% RH) and hypobaric hypoxic (2,000 m
385 altitude) sessions, but surprisingly \bar{T}_{sk} and sweat loss were unchanged and similar adaptation were evident in
386 a cool-normoxic group, indicating that some of this adaptation may have been a training effect [1].

387 A key focus of the present study was the hematological responses to the combined thermal and hypoxic-
388 stressors. Typically, HA is associated with an increase in PV and BV [74], whereas PV and BV are reduced
389 following hypoxic exposure [27, 40]. Our data demonstrated that both PV (+5.9(7.3)%) and BV
390 (+3.5(5.9)%) were increased with HA, irrespective of the additional hypoxic-stressor. This finding is

391 consistent with Takeno *et al.* [76] who demonstrated ~6% PV and ~5% BV increase following 10-days (1
392 h·day⁻¹) exercise-heat (30°C, 50% RH) and hypobaric hypoxic (2,000 m altitude) and Buchheit *et al.* [11]
393 who reported 6% PV and 4% BV changes following a 14-day warm-weather training camp including ~14(1)
394 h·day⁻¹ normobaric hypoxia (F₁O₂≈0.15). Together, these data suggest that the exercise-heat stimulus
395 predominates over the effect of hypoxia on PV and BV, at least for these magnitudes of hypoxic exposure.
396 However, a recent study demonstrated that PV expansion was ‘possibly less’ when a hypoxic stressor
397 (F₁O₂=0.144; 14 h·day⁻¹) was added to a 21 day HA programme, suggesting that a larger hypoxic stimulus
398 could blunt PV expansion [50]. Two-weeks after HA the PV and BV had returned to baseline, in line with
399 the typical decay following HA [58]. tHb_{mass} was unchanged following HA, with or without hypoxic
400 exposure; although some hematological changes can present in a delayed manner following exposure to a
401 hypoxic-stressor [6], there were also no changes in tHb_{mass} evident 14-days after cessation of either
402 intervention. Whilst data supporting the positive effect of adaptation to heat alone on tHb_{mass} are limited [72],
403 tHb_{mass} is typically increased with hypoxic exposure [10], whilst Buchheit *et al.* [11] reported a 3% increase
404 in tHb_{mass} following 14-days and McCleave *et al.* [50] reported a 4% increase following 21-days of combined
405 exercise-heat and hypoxia intervention. However, the erythropoietic effect is proportional to the magnitude
406 of hypoxic stimulus [23, 13] and participants in Buchheit *et al.* [11] and McCleave *et al.* [50] received a
407 greater hypoxic dose than participants in the present study. Moreover, Brugniaux *et al.* [10] have shown that
408 tHb_{mass} increases ~4% with ~100 h hypoxic exposure (~2,500-3,000 m); given the hypoxic dose in the
409 present study, the anticipated increase in tHb_{mass} would have approximated the CV for the CO rebreathing
410 method, possibly limiting detection.

411 Cross-stressor research has identified commonalities between heat and hypoxic stress in the HSP and HIF-1α
412 pathways, with some evidence for cross-tolerance between environments [24, 44], but the effect on these
413 pathways of concurrent exposure to these stressors is unexplored. Unfortunately, we were unable to detect
414 HIF-1α, with either HA programme, possibly due to the extracellular samples collected and the short half-
415 life of HIF-1α in normoxia [31]. However, the plasma concentration of EPO, a downstream effect following
416 the translocation of HIF-1α and subsequent gene expression in hypoxia [73], was increased following
417 medium-term HA, but this was unaffected by the addition of hypoxia to the programme. Indeed the extent of
418 the increase as a consequence of heat exposure (+28%) was similar to that reported following exposure to

419 hypoxic stress alone (+42%, five nights, 8-11 h per night, simulated altitude of 2650 m [2]). Our own
420 (unpublished) data indicate that EPO concentration is unchanged by exercise of the same duration and
421 similar intensity to our HA programme when undertaken in cool conditions (11°C), suggesting that the
422 increase was due heat-stress, or the interaction of exercise and heat-stress, rather than a training-effect, or
423 hypoxia. The lack of an additive effect of hypoxia on plasma EPO concentration during HA is not easily
424 explained. It has been suggested that combining mild stressors produces an additive effect, with a move
425 towards antagonistic interactions as the individual stressors impact increases [46], alternatively if EPO
426 production was maximally stimulated as a consequence of the heat stimulus, then the addition of a hypoxic
427 stressor would be of little consequence. Nevertheless, given the increase in EPO it is perhaps surprising that
428 there was no increase in tHb_{mass}. It may be that a greater, or more sustained, change in EPO concentration is
429 required to increase tHb_{mass} and erythrocyte volume [71]. Although reticulocytosis has been demonstrated
430 with exposure to altitude increasing serum EPO by 31-73% [38, 26, 75], other studies reporting similar
431 increases in EPO did not detect increased red blood cell production or tHb_{mass} [2, 3].

432 There was evidence for an ergogenic effect of HA on performance in a temperate-normoxic environment as
433 shown by an increase in work done in a 30 minute cycling trial (+4%) and GXT PPO (+4%), although it
434 should be noted that the performance benefit in a time trial would be somewhat less given that power is
435 related to cycling velocity with an exponent of between 2.6 and 3 [5]. However, this effect was not
436 influenced by the addition of a hypoxic-stressor and the ergogenic benefits were no longer evident two-
437 weeks after completing the HA programmes. An ergogenic effect of adaptation to heat on temperate-
438 normoxic performance has been demonstrated previously by some (*e.g.* [12, 48, 54]), but not all studies [33,
439 36], and the ergogenic efficacy of HA is controversial [15, 51, 57]. Similarly, a meta-analysis by Bonetti &
440 Hopkins [8] observed a clear ergogenic effect of adaptation to hypoxia on normoxic performance. A relatively
441 small number of studies have previously examined the ergogenic potential of adaptation to heat and hypoxia
442 in combination, but the data are equivocal. For instance, Buchheit *et al.* [11] reported an improvement in
443 temperate-normoxic performance (44% Yo-YoIR2) following HA, which was unaffected by an additional
444 hypoxic exposure. In contrast, McCleave *et al.* [50] showed a 3.3% improvement in temperate-normoxic 3
445 km running trial performance three weeks (but not immediately) after completing a 21-day intermittent HA

446 programme, but the ergogenic effect was absent when hypoxia was added to the HA programme (3,000 m,
447 13 h·day⁻¹).

448 The reasons for these discrepant findings between studies are uncertain, and where an ergogenic effect has
449 been demonstrated the physiological mechanisms are often unclear. Accordingly, in an attempt to provide
450 insight into any ergogenic effect we also assessed some of the key physiological determinants of
451 performance under temperate-normoxic conditions. Neither $\dot{V}O_{2max}$ nor GME were increased following
452 either programme. Indeed, the evidence supporting an effect of HA on GME is limited, and where an effect
453 has been demonstrated performance was not measured [67]. However, a positive effect of hypoxia on cycling
454 efficiency and running economy has been demonstrated in some studies [25, 66] and is relatively well
455 established [65]. However, the hypoxic dose is typically larger than that included in the present study [34,
456 35] and previous studies demonstrating an effect have not included an additional heat-stressor. A small
457 number of previous studies have shown an effect of HA, with [76], or without [48, 69], an additional
458 hypoxic-stressor on $\dot{V}O_{2max}$. Takeno *et al.* [76] reported an increased $\dot{V}O_{2peak}$, following their combined heat
459 and hypoxic-stressor intervention, but this was not improved to a greater extent than either stressor alone or a
460 cooler control programme, indicating a potential training effect. Similarly, Lorenzo *et al.* [48] reported an
461 increase in $\dot{V}O_{2max}$ following a 10 day HA programme, which they attributed to an increase in PV and a
462 consequent increase in stroke volume and cardiac output [28]. Although PV was expanded to a similar extent
463 in the present study, if the hemodilution effect approximates any increase in cardiac output, then O₂ delivery
464 will be unchanged; this is commonly observed with acute PV expansion in trained individuals [16] and
465 would account for the lack of change in $\dot{V}O_{2max}$ in the present study. However, a significant increase in
466 power at LT (8.6[11.0]%) was evident; whilst the LT does not directly influence performance *per se*, it is
467 well correlated and is typically used as a surrogate of sustainable percentage of $\dot{V}O_{2max}$ [32]. Indeed, Lorenzo
468 *et al.* [48] and Neal *et al.* [54] have demonstrated an increased power at lactate threshold following HA, with
469 possible mechanisms including reduced carbohydrate metabolism [83], increased strength [39] or simply
470 dilution from PV expansion. However, the increased LT was not related to the individual performance
471 improvements in either total work done or GXT PPO, which was also the case in Neal *et al.* [54], whereas
472 Lorenzo *et al.* [48] did not report correlations. Taken together the results of our study and previous studies
473 (*e.g.* [11, 48]) are not able to clearly identify the mechanisms underpinning the ergogenic effect of adaption

474 to heat (with, or without hypoxia). While it is not possible for us to discount the possibility of either a
475 placebo or training effect, we are able to conclude that the addition of a moderate hypoxic-stressor to a HA
476 programme is of no greater benefit, or harm, than HA alone on temperate-normoxic exercise performance.

477 The present study was not without limitation. Although we employed a cross-over study design, which is
478 more powerful than a parallel-groups study design, a small sample-size will increase the potential for type II
479 error. Nevertheless, our *a-priori* power calculations indicated that our sample-size would have been
480 sufficient to detect change in our key outcome variables; we detected a number of statistically significant
481 time-effects, whereas the mean between-groups differences in many of our key outcome measures (*e.g.* T_{re} ,
482 \bar{T}_b , whole body sweat rate) were typically small at each time point and within the normal daily physiological
483 variation (see Supplemental Table 1). Finally, it was not possible to exclude a role of training on the adaptive
484 responses observed in HA_{Con} and HA_{Hyp}. However, our participants were well-trained and maintained their
485 usual training volume by replacing an equivalent duration of low/moderate training with that completed in
486 the laboratory, whereas any training effects will have been similar between groups due to the balanced cross-
487 over study design.

488 In conclusion, a moderate hypoxic stressor does not affect the time-course or magnitude of
489 thermophysiological or hematological adaptations to heat. Temperate-normoxic endurance performance is
490 improved following longer-term HA, but this is unaffected by the addition of a hypoxic stimulus.

491

492 **Perspectives and Significance**

493 Adaptations to heat and hypoxia are typically studied in isolation, yet they can be encountered in
494 combination, both in the natural environment, as well as artificially when athletes expose themselves to a
495 hypoxic-stressor in order to gain favorable hematological adaptations, whilst at the same time preparing to
496 compete in a hot environment. Whether the adaptive response to these combined stressors affords the same
497 response as when examined in isolation is unclear and there are potential additive and antagonistic
498 mechanisms by which heat and hypoxic-stress may interact. The present study, using a trained cohort and
499 employing a balanced cross-over design with washout, has shown, for the first time, that the addition of a
500 moderate overnight hypoxic stimulus (equivalent to an altitude of ~2,400 m) to a 10 day HA regimen does

501 not affect the time-course or magnitude of thermophysiological adaptation to heat. Temperate-normoxic
502 endurance performance is improved following HA, but this is unaffected by a concurrent hypoxic stimulus.
503 Although these findings are mechanistically important, this observation is also practically relevant; athletes
504 preparing for competition in a hot environment should not be concerned about concurrent exposure to a
505 moderate-hypoxic stressor such as that which would occur if sleeping in a hypoxic tent. Future research
506 should seek to characterize the adaptive responses to simultaneous (rather than separate) hypoxia and heat,
507 and over longer time periods, as might occur during a prolonged high-altitude sojourn.

508 **Acknowledgements**

509 We would like to acknowledge the assistance during data collection provided by Megan Davies, Jennifer
510 Wright, Liam Colley, Danny White, Geoff Long and Amanda Ward as well as the guidance provided by Dr
511 Victoria Downie.

512

513 **Grants**

514 RN was funded by a joint English Institute of Sport and University of Portsmouth research bursary.

515

516 **Disclosures**

517 No conflicts of interest, financial or otherwise, are declared by the authors.

518 Supplementary material: Supplemental Table.

519

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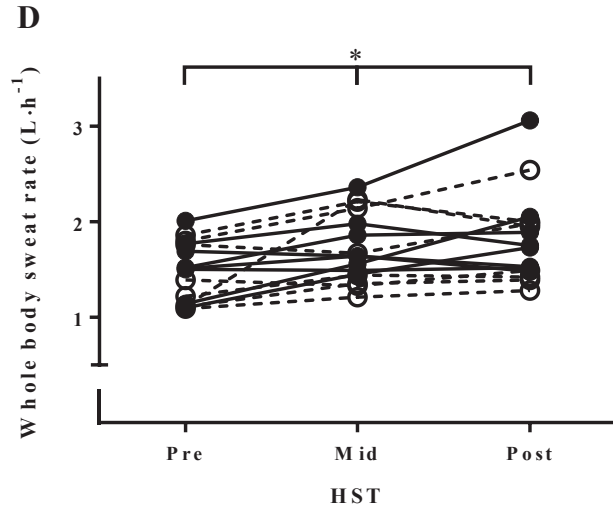
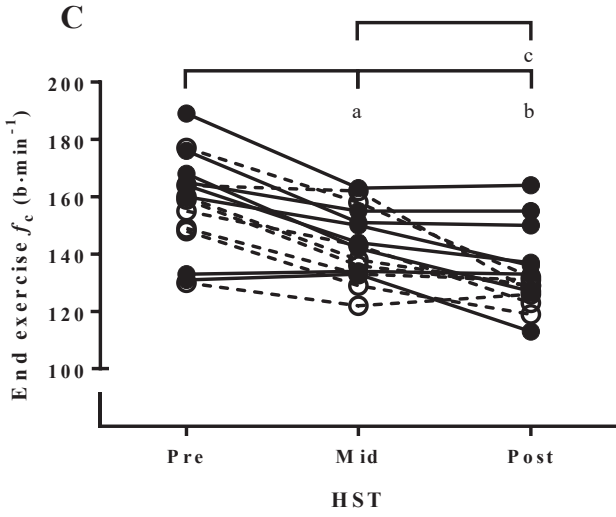
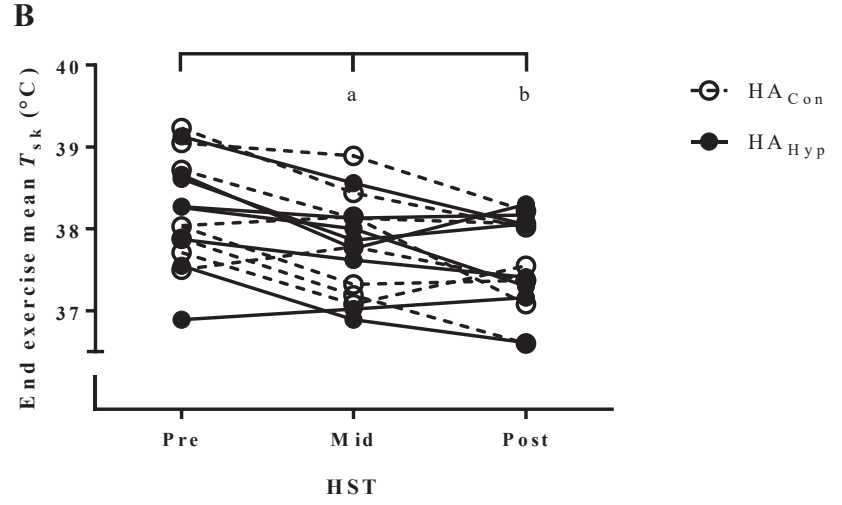
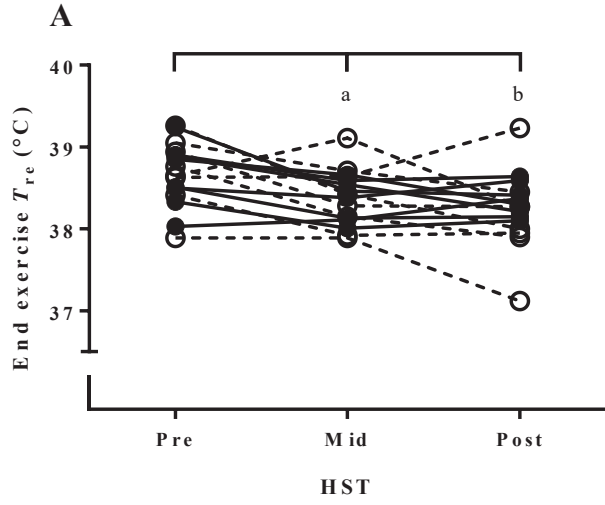
711 **Figure 1** Protocol diagram. Participants completed the heat acclimation protocol with pre/post-tests, twice,
712 in a within-subject balanced crossover design including a three to seven month washout period and two
713 conditions: HA_{Con} : Heat Acclimation Control; HA_{Hyp} : Heat Acclimation with Hypoxia. GXT =Graded
714 Exercise Test (22°C, 50% RH); $T30$ =30 minute work done trial (22°C, 50% RH); tHb_m =resting measurement
715 of total hemoglobin mass; HST =Heat Stress Test (40°C, 50% RH); ISO =Isothermal model of heat
716 acclimation (ambient conditions: 40°C, 50% RH; target T_{re} : 38.5°C); \uparrow indicates nightly hypoxic exposure in
717 the HA_{Hyp} condition ($F_{I}O_2$: 0.156).

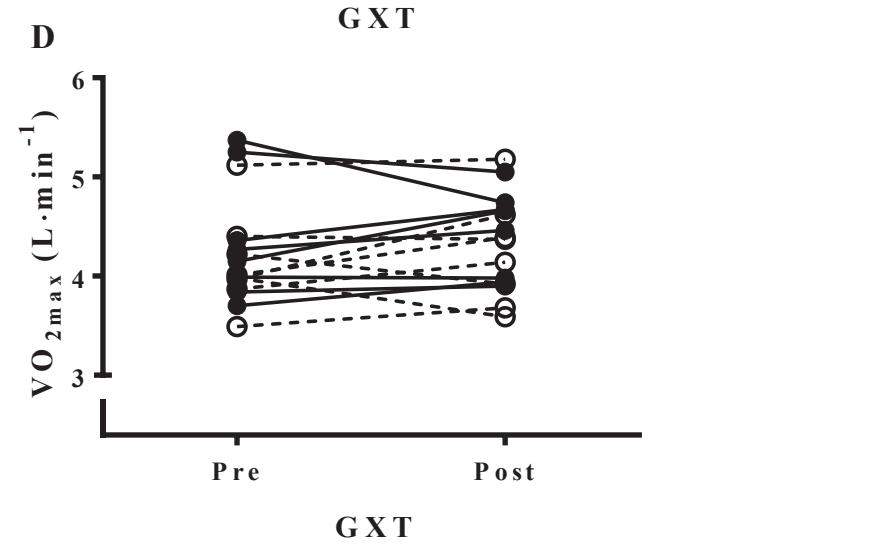
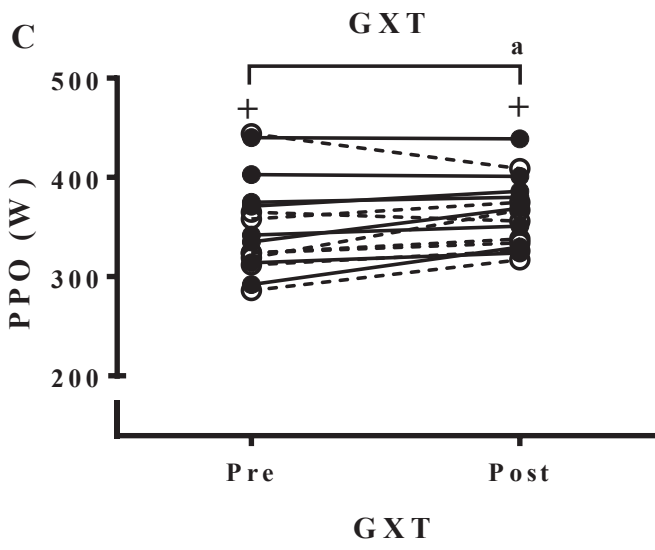
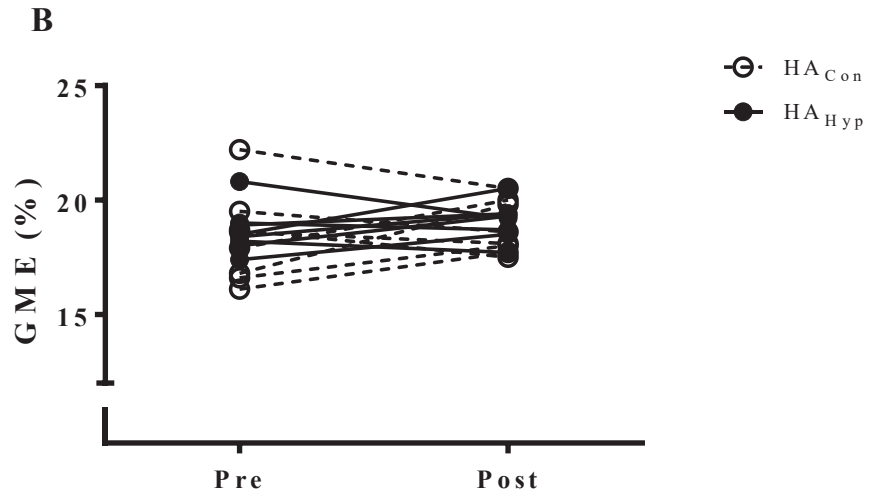
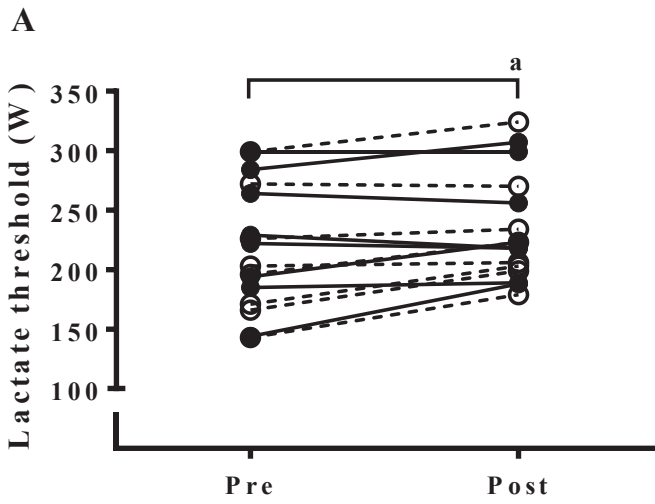
718 **Figure 2** Individual responses ($n=8$) to exercise in the heat stress test (HST) (40°C, 50% RH) before (Pre)
719 and following short- (Mid) and longer-term (Post) heat acclimation, with (HA_{Hyp} , filled circles) and without
720 (HA_{Con} , open circles) overnight normobaric hypoxia, for: A : end exercise rectal temperature; B : end exercise
721 mean skin temperature; C : end exercise cardiac frequency; D : whole-body sweat rate. * refers to a significant
722 overall time effect; ^a refers to a change from Pre-Mid, ^b from Pre-Post and ^c from Mid-Post ($P\leq 0.05$).

723 **Figure 3** Individual data ($n=8$) from the graded exercise test (GXT) in a temperate environment (22°C, 50%
724 RH) before (Pre) and after (Post) heat acclimation with (HA_{Hyp} , filled circles) and without (HA_{Con} , open
725 circles) overnight normobaric hypoxia. A : lactate threshold; B : gross mechanical efficiency (GME); C : peak
726 power output (PPO); D : maximal oxygen uptake ($\dot{V}O_{2max}$). ^a denotes a pre-post HA change over time; ⁺
727 denotes a condition effect, $P\leq 0.05$).

728 **Figure 4** Individual data from the 30 minute work done trial (T30) in a temperate environment (22°C, 50%
729 RH), before (Pre), immediately after (Post) and +14-days after (Retention) heat acclimation with (HA_{Hyp} ,
730 filled circles) and without (HA_{Con} , open circles) overnight normobaric hypoxia. *denotes a change over time
731 (over all three time points, $n=6$); ^a denotes a significant change over time (pre-post, $n=8$) ($P\leq 0.05$).

Day	-2	-1	0	1	2	3	4	5	6	7	8	9	10	11	12	13	14	15-23	24	25
Test	GXT Pre	T30 Pre	tHb _m Pre	HST Pre	ISO1	ISO2	ISO3	ISO4	HST Mid	ISO5	ISO6	ISO7	ISO8	HST Post	tHb _m Post	GXT Post	T30 Post	OFF	tHb _m Ret	T30 Ret
				↑	↑	↑	↑	↑	↑	↑	↑	↑	↑	↑						





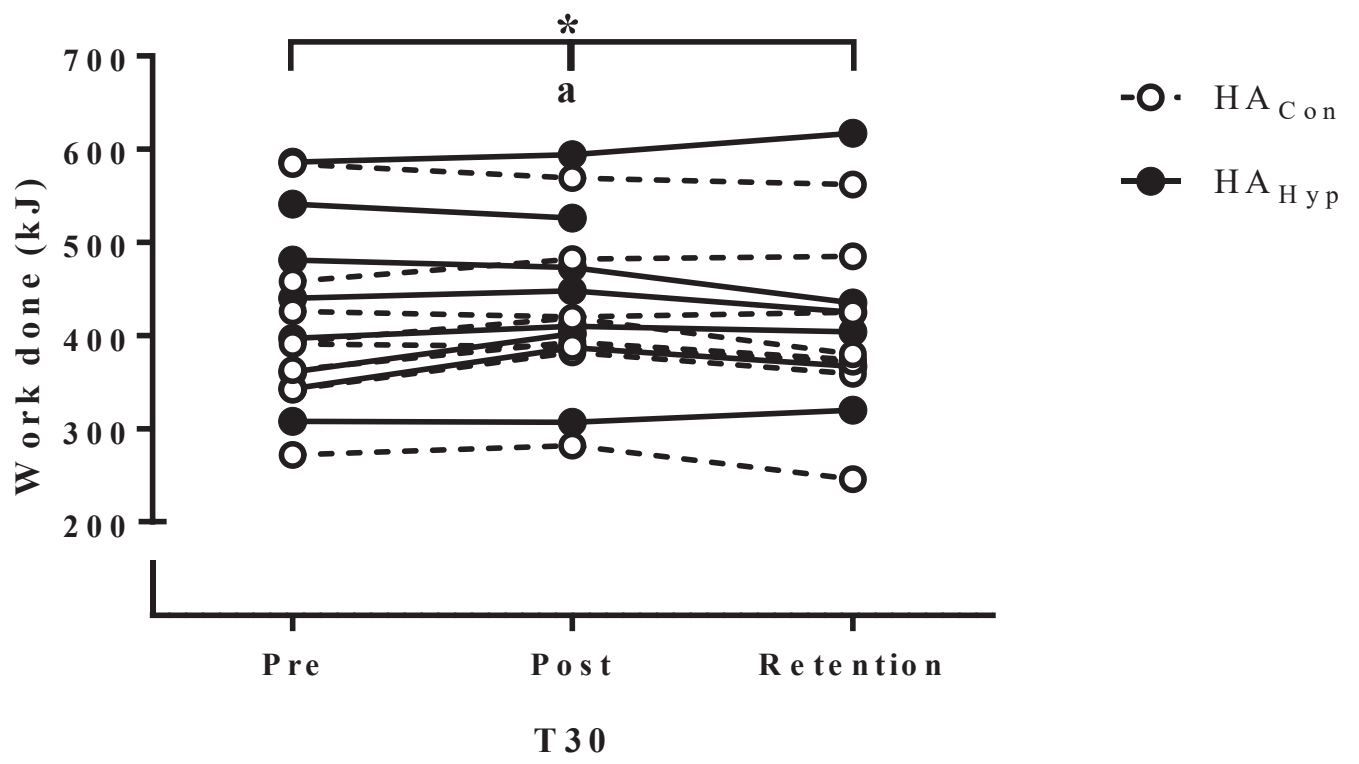


Table 1 Mean(SD) daily exercise responses ($n=8$) during medium-term heat acclimation with and without overnight hypoxia (HA_{Hyp} and HA_{Con} , respectively). In the case of a main effect for time, ^a refers to a (*post-hoc*) change between ISO1 and ISO8 ($P\leq 0.05$). In the case of a condition effect ^b denotes a significant difference between conditions at ISO6.

	ISO1		ISO2		ISO3		ISO4		ISO5		ISO6		ISO7		ISO8		Time	P value	
	HA_{Con}	HA_{Hyp}	HA_{Con}	HA_{Hyp}	HA_{Con}	HA_{Hyp}	HA_{Con}	HA_{Hyp}	HA_{Con}	HA_{Hyp}	HA_{Con}	HA_{Hyp}	HA_{Con}	HA_{Hyp}	HA_{Con}	HA_{Hyp}		Condition	Interaction
Time to target T_{re} (min)	27 (6)	30 (15)	35 (24)	33 (14)	30 (12)	34 (12)	28 (6)	32 (10)	28 (7)	33 (11)	31 (9)	27 (7)	31 (10)	32 (12)	31 (8)	30 (8)	0.556	0.431	0.806
Average T_{re} ($^{\circ}C$)	38.65 (0.14)	38.60 (0.23)	38.49 (0.36)	38.49 (0.21)	38.52 (0.15)	38.48 (0.16)	38.55 (0.10)	38.46 (0.10)	38.55 (0.09)	38.47 (0.16)	38.53 (0.14)	38.51 (0.12)	38.55 (0.13)	38.47 (0.12)	38.48 (0.09)	38.45 (0.24)	0.204	0.057	0.802
Average f_c ($b\cdot min^{-1}$)	148 (9)	142 (16)	143 (12)	142 (12)	144 (10)	140 (9)	142 (10)	142 (8)	142 (10)	142 (10)	140 (10)	140 (13)	139 (10)	140 (11)	140 (13)	139 (12)	0.166	0.194	0.419
Average power (W)	97 (18)	97 (29)	99 (20)	101 (13)	108 (18)	108 (14)	111 (22)	114 (15)	100 (17)	110 (11)	106 (12)	106 (15)	107 (11)	103 (13)	111 (16)	107 (11)	0.073	0.797	0.541
5 min peak power (W)	193 (45)	204 (65)	185 (42)	185 (47)	181 (36)	192 (48)	186 (39)	188 (38)	183 (35)	185 (41)	194 (34)	175 (33)	188 (22)	193 (54)	195 (36)	203 (35)	0.375	0.558	0.748
Pre-exercise mass (kg)	73.78 (6.51)	74.99 (7.73)	73.89 (6.69)	74.87 (7.79)	73.89 (6.65)	74.85 (7.88)	73.82 (6.51)	74.83 (7.94)	74.16 (6.82)	74.58 (8.07)	74.32 (6.82)	74.68 (7.76)	74.30 (7.00)	74.42 (7.54)	74.35 (6.80)	74.46 (7.88)	0.996	0.446	0.006
Whole body sweat rate ($L\cdot hr^{-1}$)	1.24 (0.27)	1.28 (0.43)	1.24 (0.34)	1.38 (0.26)	1.33 (0.32)	1.34 (0.29)	1.34 (0.33)	1.40 (0.26)	1.44 (0.36)	1.51 (0.34)	1.49 (0.37)	1.58 (0.41)	1.50 (0.40)	1.62 (0.39)	1.56 (0.38)	1.66 (0.34)	<0.00 ^{1a}	0.006	0.681
Urine osmolality ($mOsmo\cdot kg^{-1}$)	509 (353)	652 (335)	577 (387)	652 (335)	544 (204)	560 (274)	534 (324)	501 (288)	502 (329)	607 (249)	383 (245)	597 (289)	385 (265)	677 (323)	379 (203)	632 (290)	0.978	0.019 ^b	0.312

ISO: Isothermal strain session; HA_{Con} : Heat Acclimation Control condition; HA_{Hyp} : Heat Acclimation with Hypoxia condition; T_{re} : rectal temperature; f_c : cardiac frequency.

Table 2 Mean(SD) daily overnight responses ($n=8$) to moderate normobaric hypoxic exposure (15.6[0.9]%). Independent one-way ANOVA were performed and $P \leq 0.05$.

	HA _{Hyp} 1	HA _{Hyp} 2	HA _{Hyp} 3	HA _{Hyp} 4	HA _{Hyp} 5	HA _{Hyp} 6	HA _{Hyp} 7	HA _{Hyp} 8	HA _{Hyp} 9	HA _{Hyp} 10	<i>P</i> value
Overnight oxyhemoglobin saturation (%)	91 (1)	90 (2)	90 (2)	91 (2)	91 (2)	91 (1)	92 (2)	90 (4)	91 (1)	91 (2)	0.395
Overnight f_c ($b \cdot \text{min}^{-1}$)	65 (17)	57 (10)	61 (9)	57 (9)	54 (6)	55 (7)	54 (5)	54 (5)	57 (10)	52 (8)	0.263
Hours hypoxic exposure (h)	8.0 (1.0)	7.8 (1.2)	7.4 (1.2)	7.9 (1.5)	7.8 (1.0)	8.3 (1.5)	8.4 (1.4)	8.0 (0.5)	8.2 (0.6)	8.2 (0.8)	0.871

HA_{Hyp}: Heat Acclimation with Hypoxia condition; f_c : cardiac frequency

Table 3 Mean(SD) blood volumes ($n=6$) calculated using the optimised CO rebreathing technique pre-, post- and retention-HA for both HA_{Con} and HA_{Hyp} conditions. *Post-hoc* pairwise comparisons were performed following a significant main effect for time, ^a represents a significant change from post – retention-HA ($P \leq 0.05$).

	Pre		Post		Retention		Time	P value	
	HA _{Con}	HA _{Hyp}	HA _{Con}	HA _{Hyp}	HA _{Con}	HA _{Hyp}		Condition	Interaction
tHb _{mass} (g·kg ⁻¹)	11.7 (0.6)	11.9 (0.8)	11.6 (0.7)	12.1 (1.0)	11.4 (0.8)	11.7 (0.8)	0.153	0.398	0.774
Plasma volume (mL·kg ⁻¹)	44.9 (4.3)	44.9 (4.5)	48.0 (6.5)	47.8 (6.5)	43.8 (6.6)	43.4 (6.0)	0.006 ^a	0.889	0.955
Blood volume (mL·kg ⁻¹)	80.6 (5.6)	81.3 (5.5)	83.4 (8.2)	85.0 (7.9)	78.8 (8.0)	79.1 (7.3)	0.007 ^a	0.731	0.887

HA_{Con}: heat acclimation control condition; HA_{Hyp}: heat acclimation with hypoxia condition; tHb_{mass}: total hemoglobin mass.