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Isolating the independent effects of hypoxia and hyperventilation-induced hypocapnia on cerebral haemodynamics and cognitive function

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DOI: 10.1113/EP087602

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Document Version Peer reviewed version

Citation for published version (Harvard):

Friend, A, Balanos, G & Lucas, S 2019, 'Isolating the independent effects of hypoxia and hyperventilationinduced hypocapnia on cerebral haemodynamics and cognitive function', *Experimental Physiology*, vol. 104, no. 10, pp. 1482-1493. https://doi.org/10.1113/EP087602

Link to publication on Research at Birmingham portal

Publisher Rights Statement: Checked for eligibility: 02/08/2019

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1	Isolating the independent effects of hypoxia and hyperventilation-induced hypocapnia
2	on cerebral haemodynamics and cognitive function.
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8	
9	Running title: Cognitive and cerebrovascular responses to hypoxia and hypocapnia
10	Keywords: hypoxia, hypocapnia, cerebral blood flow, cognition
11	Word count: 6692 (Excluding references and figure legends)
12	Reference Count: 52
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17 What is the central question of this study?

To determine the independent effects of hypoxia and hypocapnia on cerebral haemodynamicsand cognitive function.

20 What is the main finding and its importance?

Our data indicates that exposure to hyperventilation-induced hypocapnia causes cognitive impairment in both normoxia and hypoxia. In addition, supplementation of carbon dioxide during hypoxia alleviates the cognitive impairment and reverses hypocapnia-induced vasoconstriction of the cerebrovasculature. These data provide new evidence for the independent effect of hypocapnia on the cognitive impairment associated with hypoxia.

Abstract

Hypoxia, which is accompanied by hypocapnia at altitude, is associated with cognitive impairment. This study examined the independent effects of hypoxia and hypocapnia on cognitive function and assessed how changes in cerebral haemodynamics may underpin cognitive performance outcomes. Single reaction time (SRT), five-choice reaction time (CRT) and spatial working memory (SWM) tasks were completed in 20 participants at rest and after one hour of isocapnic hypoxia (IH, end-tidal oxygen partial pressure ($P_{ET}O_2$) = 45mmHg, end-tidal carbon dioxide partial pressure (PerCO₂) clamped at normal), and poikilocapnic hypoxia (PH, $P_{ETO_2} = 45 \text{mmHg}$, P_{ETCO_2} not clamped). A subgroup of 10 participants were also exposed to euoxic hypocapnia (EH, $PetO_2 = 100mmHg$, $PetCO_2$) clamped 8mmHg below normal). Middle cerebral artery velocity (MCAv) and prefrontal cerebral haemodynamics were measured with transcranial Doppler and near infrared spectroscopy, respectively. IH did not affect SRT and CRT performance from rest (566 \pm 50ms and 594 \pm 70ms), whereas PH (721 \pm 51ms and 765 \pm 48ms) and EH (718 \pm 55ms and 755 ± 34 ms) slowed response times (p<0.001 vs IH). Performance on the SWM task was not altered by condition. MCAv increased during IH compared to PH (p<0.05), which was unchanged from rest. EH caused a significant fall in MCAv and prefrontal cerebral oxygenation (p<0.05 vs baseline). MCAv was moderately correlated to cognitive performance ($R^2=0.266-0.289$), whereas prefrontal cerebral tissue perfusion and saturation were not (p>0.05). These findings reveal a role of hyperventilation-induced hypocapnia per se on the development of cognitive impairment during normoxic and hypoxic exposures.

26

27 Table of Abbreviation	IS
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28	CANTAB	Cambridge Neuropsychological Test Automated Battery
29	CaO_2	arterial oxygen content
30	CBF	cerebral blood flow
31	CMRO ₂	cerebral metabolic rate of oxygen
32	CRT	five-choice reaction time task
33	EH	euoxic hypocapnia
34	HCO ₃ ⁻	bicarbonate ion
35	IE	isocapnic euoxic
36	IH	isocapnic hypoxia
37	MAP	mean arterial pressure
38	MCAv	middle cerebral artery velocity
39	NIRS	near infrared spectroscopy
40	nTHI	total haemoglobin normalised to the initial value
41	PaCO ₂	partial pressure of arterial carbon dioxide
42	PaO ₂	partial pressure of arterial oxygen
43	PetCO ₂	end-tidal partial pressure of carbon dioxide
44	PetO ₂	end-tidal partial pressure of oxygen
45	РН	poikilocapnic hypoxia
46	SRT	single reaction time task
47	SWM	spatial working memory
48	TCD	transcranial Doppler
49	TOI	total oxygenation index

50 <u>Introduction</u>

51 Exposure to high altitude can cause a number of hypoxia-induced physiological complications such as acute mountain sickness, pulmonary and/or cerebral oedema, and 52 impairment of cognitive function (Hackett & Roach, 2001). Individuals become quickly 53 aware of physical symptoms such as dizziness, headaches and nausea at altitude (Hackett & 54 Roach, 2001), but they are less aware of the impairment to their cognitive function (Asmaro, 55 Mayall, & Ferguson, 2013). The degree to which cognitive function is impaired is related to 56 the severity of the hypoxic stimulus, particularly for tasks that require a higher order of 57 cognitive ability (Petrassi, Hodkinson, Walters, & Gaydos, 2012; Yan, 2014). This higher 58 59 order ability is essential for decision-making and attentional processes in individuals who venture to unfamiliar and dangerous environments, such as is typical of the high-altitude 60 environment. 61

The brain relies on two variables to maintain sufficient oxygen supply and its functional 62 63 capacity; namely, arterial oxygen content (CaO₂) and cerebral blood flow (CBF). During 64 exposure to hypoxia, partial pressure of arterial oxygen (PaO_2) will fall (and related CaO_2) and subsequently the cerebrovasculature dilates in order to increase CBF to maintain global 65 oxygen delivery to the brain (Kety & Schmidt, 1948; Willie, Tzeng, Fisher, & Ainslie, 2014). 66 Simultaneously, the peripheral chemoreceptors activate the hypoxic ventilatory response to 67 increase oxygen intake via the lungs. Consequently, this increased respiration gives rise to 68 hypocapnia, a known vasoconstrictor of the cerebrovasculature (Kety & Schmidt, 1946). 69 Therefore, the change in CBF is influenced by two conflicting stimuli, with the balance of 70 71 these changes in oxygen and carbon dioxide tensions key factors in the overall change in CBF during exposure to hypoxia (Lucas et al., 2011; Bruce et al., 2016). Given this, 72 hypocapnic-induced vasoconstriction could play a defining role in the cognitive impairment 73

experienced at altitude through compromising cerebral tissue perfusion via its effect on thecapacity of the vasculature to dilate in response to hypoxaemia.

To investigate the physiological effects of hypocapnia participants are often instructed to 76 voluntarily hyperventilate. Studies using this method have demonstrated that hypocapnia 77 compromises brain function through its effect on the cerebrovasculature and produces similar 78 impairment to that experienced at altitude, as evidenced by reports of light-headedness and 79 80 dizziness (Bresseleers, Van Diest, De Peuter, Verhamme, & Van den Bergh, 2010), and impairment of complex cognitive tasks such as Stroop Test performance (Van Diest, Stegen, 81 Van de Woestijne, Schippers, & Van den Bergh, 2000). The ambient gas compositions 82 83 experienced at altitude are as consequence of a reduction in atmospheric pressure (hypobaric hypoxia), but can be mimicked in the laboratory setting through a reduction in partial 84 pressure of oxygen (normobaric hypoxia). Despite some evidence suggesting different 85 86 physiological responses between hypobaric hypoxia and normobaric hypoxia (Savourey, Launay, Besnard, Guinet, & Travers, 2003), the ability to tightly control gas composition in 87 the laboratory setting enables the comparison of poikilocapnic hypoxia (PH), as it occurs 88 naturally from hypoxia-induced hyperventilation, to that of isocapnic hypoxia (IH), where the 89 effects of hypoxia *per se* can be separated from hypocapnia by clamping partial pressure of 90 arterial carbon dioxide (PaCO₂) at its normal value. Using such an approach, Van Dorp et al. 91 (2007) compared the effects of PH with that of IH on a combination of vigilance and multi-92 attribute cognitive tasks and found that carbon dioxide supplementation during hypoxia (IH) 93 alleviated the impairment in cognitive function such that performance was similar to that 94 under normoxic conditions. The authors concluded that the hypocaphic element of PH may 95 be directly related to the compromised cognitive function. 96

97 However, the independent contribution of hypocapnia to cognitive function and its link to98 CBF during hypoxia remains unclear. To our knowledge, no study has attempted to separate

99 the roles of hypoxia *and* hypocapnia on cognitive function, as well as the associated changes 100 in cerebral haemodynamics and task performance. Therefore, the present study was designed 101 to examine the isolated effects of hypocapnia and hypoxia on simple and complex cognitive 102 tasks, as well as to explore how changes in global and prefrontal cerebral haemodynamics 103 might relate to changes in cognitive performance.

104 <u>Methods</u>

105 <u>Ethical Approval</u>

Ethical approval for this study was provided by the Safety and Ethics Subcommittee of the School of Sport, Exercise and Rehabilitation Sciences at the University of Birmingham (reference: MW 07/10/14) and was conducted in accordance with the standards of the *Declaration of Helsinki*, except for registration in a database, with written informed consent obtained from participants before they took part in the study.

111 <u>Participants</u>

Twenty young healthy males (aged 22.4 ± 6.3 years) participated in this study. All participants completed a general health questionnaire and were invited to participate if they were healthy, non-smokers, and had no history of cardiorespiratory disease. Participants were asked to refrain from consuming alcohol and from undertaking strenuous exercise within 24 hours of each experimental session. Participants were also asked not to consume caffeinated drinks within six hours, and food within two hours prior to reporting to the laboratory.

118 <u>Study Design and Procedures</u>

All participants visited the laboratory on three occasions, once for a familiarisation session and then for two experimental sessions performed in a random order and separated by at least hours. A subgroup of 10 participants completed a third experimental session. All participants completed an IH session (end-tidal partial pressure of oxygen (PerO₂) = 45mmHg and end-tidal partial pressure of carbon dioxide (PerCO₂) clamped at each participant's normal value) and a PH session (PerO₂ = 45mmHg and PerCO₂ not controlled), while the subgroup completed an additional euoxic hypocapnia (EH) session (PerO₂ = 100 mmHg and PerCO₂ clamped at 8 mmHg below each participant's normal value) (see Figure 2). Participants were blinded to IH and PH conditions only, as participants were coached to maintain a ventilation rate during EH.

129 Familiarisation

Participants visited the laboratory to familiarise themselves with the equipment and procedures that were used in the study. During this session, participants completed one repeat of the reaction time tasks and three repeats of the spatial working memory (SWM) task of the Cambridge Neuropsychological Test Automated Battery (CANTAB) programme to minimise any learning effect on performance outcomes during the experimental conditions.

135 Isocapnic Hypoxia (IH)

Participants were comfortably seated while being instrumented to measure cerebral 136 haemodynamics, peripheral arterial oxygen saturation, mean arterial blood pressure and heart 137 rate. The pulse oximeter probe and blood pressure finger cuff were attached to fingers on 138 their non-dominant hand, allowing their dominant hand to be used for the cognitive function 139 tests. Once instrumentation was complete and the signals were optimised, participants 140 breathed through a mouthpiece whilst wearing a nose clip. Control of end-tidal gases was 141 142 achieved by means of a dynamic end-tidal forcing system described in detail elsewhere (Robbins, Swanson, & Howson, 1982). Participants completed the first battery of cognitive 143 function tests under isocapnic euoxic (IE) conditions ($PetO_2 = 100 \text{ mmHg}$ and $PetCO_2$ 144 145 clamped at participant's normal value). This was followed by a 60-minute intervention period

146	during which participants were exposed to IH, followed by a repeat of the cognitive function
147	tests whilst remaining under IH conditions. Once the cognitive function tests were completed
148	participants were returned to breathing room air and equipment was removed.

149 *Poikilocapnic Hypoxia (PH)*

This protocol was identical to the one described for IH except that PETCO₂ was not controlled
 during the 60-minute intervention and during the repeat of the cognitive function tests.

152 Euoxic Hypocapnia (EH, n=10)

This protocol was identical to the one described for IH except that participants were exposed 153 to EH during the 60-minute intervention and during the repeat of the cognitive function tests. 154 Hypocapnia was achieved through voluntary hyperventilation. For this, participants were 155 coached to hyperventilate enough to reduce their PETCO₂ to approximately 10 mmHg below 156 their normal value, allowing the dynamic end-tidal forcing system to then adjust PETCO₂ to 8 157 mmHg below accurately. Figure 1 shows a schematic of the protocol during each 158 experimental visit, as well as examples of each of the CANTAB tests completed under each 159 condition. 160

161 Equipment and Measures

162 Cognitive Function Assessment

Cognitive function was measured via a touch screen CANTAB cognition computer test (Cambridge Cognition Ltd., United Kingdom). The CANTAB is a valid neuropsychological testing instrument of cognitive function (Smith, Need, Cirulli, Chiba-Falek, & Attix, 2013), and is regularly used to assess cognitive function in both healthy and neurodegenerative cohorts (e.g. mild cognitive impairment (Saunders & Summers, 2010) and Alzheimer's 168

disease (Matos Goncalves, Pinho, & Simoes, 2018)). Reaction time tasks and the SWM tests were performed representing simple and complex cognitive tasks respectively. 169

Reaction Time Tasks 170

Reaction time was measured through two tasks; single reaction time task (SRT) and five-171 choice reaction time task (CRT). Both tasks required participants to hold down a pressure pad 172 placed in front of the computer and to tap a circle on the monitor as quickly as possible after 173 a yellow spot was displayed within it. The time taken for the yellow spot to appear was 174 randomised between trials. This task was completed with a single response circle for the 175 SRT, whilst the spot had the option to flash in any one of five response circles in the CRT 176 (see Figure 1a). Participants were given practice attempts of both tasks prior to the test period 177 in which their performance was recorded. Performance time was recorded as the sum of 178 reaction time (time taken between the yellow spot appearing and releasing the pad) and 179 movement time (time taken between releasing the pad and tapping the circle). Additionally, 180 error count (releasing the pad too early or missing the correct circle) was measured for both 181 182 reaction time tasks.

Spatial Working Memory Task 183

SWM was measured through a visuospatial task. The participant was presented with a 184 selection of coloured boxes on the screen and the aim was to find all of the tokens hidden 185 inside these boxes. Participants were required to use working memory and a process of 186 elimination to find all of the tokens as only one token was hidden at a time and would never 187 188 be found in the same box again. Three sets of practice trials (three boxes within each set) were completed before performance was recorded across three stages of increasing difficulty, 189 with each stage consisting of four sets of trials with four, six and eight boxes, respectively, 190 191 for each level of increasing difficulty. The total number of errors were recorded as the 192 measure of performance. Errors were recorded when participants returned to a box where a 193 token had already been found, or when a box that had been previously selected was selected 194 again in a subsequent search.

195 Cerebral blood flow velocity and prefrontal cerebral haemodynamics

Bilateral measures of blood flow velocity from the left and right middle cerebral artery 196 (MCAv) were measured using a 2 MHz pulsed Transcranial Doppler (TCD) ultrasound 197 system (Doppler Box, DWL, Compumedics Ltd, Germany) using standardised procedures 198 (Willie et al., 2011). Probes were placed over the left and right temporal windows and 199 secured in place via an adjustable head piece. Photographs of the probe position and angle 200 were used to replicate the placement between sessions, and signal depth and gain settings 201 were also replicated. Left and right side MCAv measures were averaged, reported as a pooled 202 mean, and expressed as a change from resting baseline. 203

In the subgroup of 10 participants that completed all three protocols prefrontal cerebral 204 haemodynamics was also monitored non-invasively on the left and right side of the forehead 205 using near infrared spectroscopy (NIRS; NIRO-200NX, Hamamatsu Photonics KK; 206 Hamamatsu, Japan). The NIRS probes were housed in light-shielding cases and attached to 207 the forehead skin with tape in the same position for each session. Probes were placed as 208 lateral and superior as possible to avoid the frontal sinus and to allow the TCD head piece to 209 fit between the probes and the superior orbital ridge (i.e. probe centre points were located 210 approximately 4 cm from the midline and approximately 3 cm above the orbital ridge). The 211 212 NIRO-200NX device measures changes in chromophore concentrations of oxyhaemoglobin and deoxyhaemoglobin via the modified Beer-Lambert law and provides depth-resolved 213 214 measures of tissue oxygen saturation [total oxygenation index (TOI)] and tissue haemoglobin content (i.e., relative value of the total haemoglobin normalised to the initial value, nTHI) 215

using the spatially resolved spectroscopy (SRS) method. The SRS-derived NIRS parameters
limit contamination from superficial tissue via depth-resolved algorithmic methods, providing
an index of targeted local tissue saturation (TOI) and perfusion (nTHI) (Davies et al., 2015).
Given the inter-individual variability of baseline measures using this imaging technology
(Davies et al., 2017) and in accordance with recommendations of others (Subudhi, Miramon,
Granger, & Roach, 2009), these NIRS data are expressed as the magnitude of the change
from the resting baseline value.

Cerebrovascular haemodynamics, cardiovascular and respiratory variables were all acquired
continuously at 200 Hz using an analogue-to-digital converter (Powerlab/16SP ML795;
ADInstruments, New Zealand) interfaced and displayed in real time using LabChart software
(Chart v7.5, ADInstruments) on a computer.

227 Data Analysis

Mean SRT and CRT performance time and error count, and SWM task mean error count were collected from each CANTAB trial. A 60 s mean for MCAv, TOI and nTHI data were collected from the two baseline measures that preceded each CANTAB battery under IE or experimental conditions. During CANTAB battery periods, MCAv, TOI and nTHI data were averaged from the final 20 s of each reaction time task (SRT and CRT) and the final 30 s of the SWM task. One participant's TOI data was lost due to corruption of the containing file.

A repeated-measures analysis of variance (IBM SPSS Statistics v23) was used to assess the relations between condition (IH, PH, EH), time (IE, Experimental) and task phase (Baseline, SRT, CRT, SWM) for each physiological variable. A repeated measures analysis of variance was also used to assess the relations between condition (IH, PH, EH) and time (IE, Experimental) for each CANTAB performance variable. Pairwise comparisons (Bonferroni adjusted) were applied to evaluate main effects and interactions. The relationship between changes in selected physiological variables (MCAv, TOI, nTHI) and change in reaction time task performance (SRT and CRT) were determined using Pearson's correlations. Data are presented as mean \pm SD and statistical significance was accepted at p < 0.05.

243 <u>Results</u>

There were marked differences between IE baseline and experimental measures of $PetO_2$, PetCO₂, MCAv, TOI and nTHI (see Table 1). This general pattern was consistent during cognitive testing (see Figures 2 and 3), with no significant differences between the measured time points within each condition (all p > 0.05). Nevertheless, we have presented the haemodynamics for each specific time point in Figure 3, but for brevity we have summarised our findings using pooled data across the cognitive tasks and report differences between condition (IH, PH, EH) and time (IE, Experimental) for each dependent variable.

251 End Tidal Gas Control

Baseline and experimental end-tidal values are shown in Table 1, and a representative 252 example of the differences shown in Figure 2. By design, end-tidal gases were similar during 253 IE conditions, and were successfully manipulated and held consistent during cognitive testing 254 under experimental conditions. Specifically, PETCO₂ remained clamped at IE values during 255 IH (41.1 \pm 2.0 mmHg), whereas PETCO₂ declined during the PH (37.4 \pm 2.7 mmHg; p < 256 0.001 vs IE and p < 0.001 vs IH). For the subgroup completing the EH condition, PetCO₂ 257 was lowered to 32.6 ± 2.3 mmHg (p < 0.001 vs IE), significantly lower than IH (40.9 ± 1.8 258 mmHg; p < 0.001) and PH (36.6 ± 3.0 mmHg; p < 0.01). The reductions in PetO₂ during IH 259 260 $(44.2 \pm 1.7 \text{ mmHg})$ and PH $(43.2 \pm 2.4 \text{ mmHg})$ interventions were similar (both p < 0.05 vs IE). For the subgroup, PetO₂ during the EH condition remained clamped at IE levels (98.6 \pm 261 6.0 mmHg), which was significantly greater than IH (44.2 \pm 2.2 mmHg; p < 0.001) and PH 262 $(43.5 \pm 3.2 \text{ mmHg}; p < 0.001).$ 263

264 Haemodynamic Measurements (Isocapnic Euoxic vs Experimental conditions)

Baseline absolute measures of heart rate, mean arterial pressure (MAP), MCAv, nTHI and 265 TOI in IE conditions were consistent between all sessions and are shown in Table 1. There 266 was no difference in heart rate between IE and experimental conditions, whereas there was a 267 main effect of time for MAP (p < 0.05) representing elevated values during the experimental 268 conditions compared to IE baseline. Compared to IE, MCAv increased during IH (up $6.7 \pm$ 269 7.2 cm·s⁻¹; p < 0.001 vs IE) whereas it remained similar during PH (p = 0.63 vs IE) and thus 270 lower than IH (p < 0.001). In the subgroup, similar results for IH (up 6.6 ± 8.5 cm s⁻¹; p < 271 0.05 vs IE) and PH (p = 0.16 vs IE, and p < 0.05 vs IH) conditions were seen, while MCAv 272 decreased by $9.2 \pm 6.4 \text{ cm} \cdot \text{s}^{-1}$ from IE (p < 0.001) during the EH condition (p < 0.01 vs IH, 273 and p = 0.18 vs PH). Measures of prefrontal cerebral haemodynamics collected via NIRS in 274 the subgroup completing all three conditions demonstrated that prefrontal perfusion (as 275 indexed by nTHI) increased from IE for IH (up 0.05 ± 0.05 au; p < 0.05 vs IE) and PH (up 276 0.05 ± 0.08 au; p = 0.071 vs IE), while nTHI decreased in EH (down 0.05 ± 0.04 au; p < 0.05 277 vs IE, and p < 0.05 IH vs PH). All conditions recorded a significant decline in prefrontal 278 tissue saturation (indexed by TOI) compared to IE (p < 0.001), with a greater decrease 279 recorded in IH (down $8.8 \pm 3.2\%$) and PH (down $9.4 \pm 3.3\%$) conditions relative to EH 280 (down $4.2 \pm 2.0\%$; p < 0.05 vs IH and PH). Figure 3 shows these cerebral haemodynamic 281 changes for each experimental condition relative to the proceeding IE baseline. 282

283 Cognitive Task Performance

Simple and Complex Reaction Time: Performance scores for both reaction time tasks are shown in Table 2. Baseline IE measures were consistent between all conditions (p > 0.05). During IH, performance times for SRT (566 ± 50 ms) and CRT (594 ± 70 ms) tasks were unaffected with respect to IE (p > 0.05), whereas PH caused a significant slowing of both SRT (by 149 \pm 81 ms; p < 0.001 vs IH) and CRT (by 152 \pm 82 ms; p < 0.001 vs IH) performance. For the subgroup, EH produced similar performance decrements as was observed during PH (p > 0.05) for both SRT (slower by 174 \pm 42 ms; p < 0.001 vs IH) and CRT (slower by 167 \pm 70 ms; p < 0.001 vs IH) performance. There was no effect of condition on SRT and CRT error count.

Spatial Working Memory Task: Error count for the SWM task is shown in Table 2. There was
no significant change in error count during the experimental conditions compared to IE
conditions for any protocols.

296 <u>Relation between cerebral haemodynamics and cognitive task performance</u>

Finally, as shown in figure 4A and B, changes in MCAv were moderately correlated ($R^2 = ~0.28$) with both SRT and CRT, such that increases in MCAv were associated with maintained reaction time task performance. These correlations were not apparent in the NIRS-derived prefrontal cortex measures of tissue saturation and perfusion (as indexed by TOI and nTHI, respectively), with no significant correlations observed (all p > 0.05, see Figures 4C-F).

303 <u>Discussion</u>

The present study was designed to investigate the independent roles of hypoxia and 304 hypocapnia on simple and complex cognitive ability, and how changes in global and 305 prefrontal cerebral haemodynamics were associated with altered cognitive performance. We 306 found that acute exposure to PH impaired both SRT and CRT performance, but it had no 307 apparent effect on SWM task performance. Hypocapnia alone (i.e. EH) produced similar 308 309 decrements to those seen during PH, whilst the supplementation of carbon dioxide to maintain PetCO₂ relieved the hypoxia-induced cognitive impairment. The associated changes 310 in cerebral haemodynamics indicate that differences in CBF between the experimental 311 312 conditions may mediate this effect, with the changes in global flow (as indexed by MCAv) moderately correlated to cognitive task performance. Interestingly, despite differences in global flow and the associated link to performance, prefrontal cerebral tissue perfusion and saturation were not different between hypoxic trials and not linked to cognitive performance. Overall, these findings reveal a significant role of hypocapnia *per se* on the development of cognitive impairment during normoxic *and* hypoxic exposures.

318 Cognitive Function during Hypoxia and Hypocapnia

The observed detriment to cognitive function during PH reported in the current study is 319 consistent with previous work showing impairment in CRT during exposure to high altitude 320 (Dykiert et al., 2010). Further, our findings of the recovered cognitive performance during 321 carbon dioxide supplementation in hypoxia has also been previously demonstrated (Van Dorp 322 et al., 2007). However, to our knowledge no such cognitive impairment has been found when 323 tasks are completed under hypocapnia when controlling for hypoxia, nor demonstrated how 324 cerebral haemodynamic changes may mediate this effect (discussed below). Interestingly, 325 326 Bloch-Salisbury and colleagues reported significant changes to electroencephalographic 327 signals under hypocapnia during a series of rapid-response cognitive tasks (Bloch-Salisbury, Lansing, & Shea, 2000); however, these changes did not reflect impairment to response time 328 or error scores despite a similar hypocapnic stimulus (PerCO₂ of ~30mmHg) to that induced 329 in the current study. The present data exhibited a speed-accuracy trade-off for SRT and CRT 330 performance during PH and EH conditions, with significantly slower performance times 331 recorded with no change to error count. An unexpected finding of the present study was that 332 the performance of the SWM task was unaffected by all conditions. It is widely accepted that 333 as altitude increases, complex cognitive abilities, such as working memory become 334 progressively impaired (reviewed in Yan, 2014). Studies using test batteries to examine 335 executive function performance during hypoxia have found impairments in the Paced 336 337 Auditory Serial Addition Test (PASAT) (Malle et al., 2013) and Stroop Test (Turner, Barker-

Collo, Connell, & Gant, 2015) task performance. Despite differences in mean average error 338 count, it is likely that we did not have the power (effect size = 0.279, observed power = 339 0.498) to detect any significant differences in SWM task performance as a consequence a 340 lack of sensitivity of the SWM CANTAB task. Further, Lowe and Rabbitt (1998) described 341 that for executive function to be measured effectively tasks must remain novel to the 342 participant due to the rapid improvements in performance once an optimal strategy is 343 discovered. Specifically, the familiarisation session conducted to minimise the learning 344 effects may have provided a ceiling effect for SWM task performance The CANTAB SWM 345 346 task used here is designed to test memory retention, strategy, and visuospatial abilities as a representation of executive function. The version of the SWM task used in this present study 347 produces 15 identical arrangements of coloured boxes for each repeat, which may diminish 348 its ability to reliably measure executive function. Patients with mild cognitive impairment 349 and Alzheimer's disease completing the CANTAB SWM task in a 6 month test-retest 350 assessment are shown to exhibit a practice effect by optimising their strategy search patterns, 351 which was maintained at the 12-month re-test assessment (Cacciamani et al., 2018). 352 Subsequently in the present study, the acute test-retest period that was used (within ~ 1 hr) 353 would likely have been compromised by this learning effect. In addition, the measurements 354 of error collected by the SWM task may not provide adequate information to determine 355 whether there is impairment to performance. Based on our reaction time task performance 356 357 decrements, it was the speed of the response that was impaired as opposed to the accuracy. As such, including a time pressure during a cognitive task may be a more effective way to 358 demonstrate the hypoxic impairment effect given its effect on a recall task (Earles, Kersten, 359 Berlin Mas, & Miccio, 2004). Indeed, this is consistent with observations of hypoxia-related 360 impairment of PASAT test performance (Malle et al., 2013), a task which includes a time 361 362 pressure.

363 Cerebral Haemodynamics and Cognitive Function

Exposure to hypoxia is well known to cause a cerebral vasodilatory response but is 364 compromised by the reflex hypoxia-induced hyperventilation response lowering PaCO₂ and 365 causing cerebral vasoconstriction (Ainslie & Ogoh, 2010). In the present study, there was no 366 change in MCAv observed during PH, reflecting the contrasting cerebrovascular activity that 367 hypoxia and hypocapnia stimulate (Mardimae et al., 2012). Consistent with previous 368 observations, the supplementation of carbon dioxide to maintain PerCO₂ constant during the 369 hypoxic exposure (i.e. IH) allowed the cerebrovasculature to dilate and thus to increase 370 oxygen delivery to the brain via elevated flow (Van Dorp et al., 2007). Indeed, higher blood 371 372 flow velocity was associated with maintained reaction time task performance (Fig 4A and 4B). Interestingly, while increases in global cerebral haemodynamics were observed during 373 IH compared to PH (and EH), the NIRS-based measures of regional tissue perfusion as 374 indexed by haemoglobin content (i.e. nTHI) at the prefrontal cortex was not different 375 between the hypoxia conditions, which increased similarly in both hypoxic conditions. A 376 potential explanation is that this may reflect a global increase in CBF during IH, compared to 377 a regional shift of blood towards active areas of the brain during PH, particularly at the 378 prefrontal cortex. Binks and colleagues reported a global increase in CBF to all areas of the 379 380 brain during IH, but not necessarily each to the same magnitude (Binks, Cunningham, Adams, & Banzett, 2008). Additionally, Lawley et al. reported an active heterogeneous CBF 381 response following two hours of PH, with increased perfusion observed in the anterior 382 portions of the brain and reductions to the posterior regions (Lawley, Macdonald, Oliver, & 383 Mullins, 2017). It is known that different portions of the brain are activated depending on the 384 task completed, with working memory processes stimulating the prefrontal cortex (van 385 Asselen et al., 2006), whereas reaction time tasks activate both the premotor and primary 386 sensorimotor areas (Kwon, Kwon, & Park, 2013). This regional activation may explain why 387

no significant haemodynamic differences were seen between the impaired reaction time tasks
and the unimpaired SWM task as only prefrontal cortex measurements were recorded.
Further investigation using whole-head functional imaging would enable a clearer
understanding of the regional differences in CBF during cognitive tasks under hypoxia and
hypocapnia.

Despite possible differences in the maintenance of local blood flow, there was an equivalent 393 394 fall in cerebral oxygenation (TOI) observed in both IH and PH, indicating that insufficient delivery of oxygen to the tissue is not the defining factor behind the cognitive function 395 difference. This is demonstrated with no meaningful correlations found between TOI and 396 397 reaction time task performance (Fig 4C-D). Hypocapnia causes haemoglobin to have an increased affinity for oxygen and reduce oxygen unloading at the tissues (Collins, Rudenski, 398 Gibson, Howard, & O'Driscoll, 2015). This may be a defining factor between the two 399 400 hypoxic conditions in the development of cognitive impairment, with the supplementation of carbon dioxide reversing the leftward shift of the oxygen-haemoglobin dissociation curve, 401 402 allowing adequate offloading of oxygen into the tissue. This is highlighted during EH given that there was less of a fall in TOI but still a cognitive impairment. With hypoxia-induced 403 hypocapnia comes respiratory alkalosis and acid-base adjustment via renal compensation 404 through excretion of bicarbonate ion (HCO_3) , although this is typically reported with longer 405 exposures than the 60 minutes we used here. Further, it remains undefined whether PaCO₂ or 406 pH acts as the primary stimulant responsible for cerebral vasoconstriction (Willie, Tzeng, 407 Fisher, & Ainslie, 2014). Nonetheless, hypocapnia-induced vasoconstriction has been shown 408 to impact the neurovascular coupling response, such that it overwhelms the neuronal 409 activated vasodilation response to visual stimulation, and compromises oxygen supply to the 410 411 brain (Szabo et al., 2011). The combination of a compromised oxygen supply and reduced oxygen unloading causes hypocapnia-induced brain ischaemia (Laffey & Kavanagh, 2002) 412

and could well be an underlying factor in the development of the cognitive impairment during 413 hypoxic exposure. In addition to altering the neurovascular coupling response, the cerebral 414 metabolic rate of oxygen (CMRO₂) does not change during isocapnic hypoxia (Ainslie et al., 415 2014), with MRI-based evidence indicating that increased neural excitability (and subsequent 416 CMRO₂) during hypoxia are as a consequence of hypoxic ventilatory response-induced 417 hypocapnia (Smith et al., 2012; Vestergaard et al., 2015). This increase in CMRO₂ has been 418 shown to be mitigated during hypoxia with the administration of acetazolamide (Wang, 419 Smith, Buxton, Swenson, & Dubowitz, 2015), which indicates an important role of 420 421 hypocapnia and alkalosis in cerebral metabolism during acute hypoxia.

422 In the present study, the use of an acute exposure to normobaric hypoxia enables the controlled manipulation of oxygen and carbon dioxide to investigate their impact on 423 cognitive function. During extended or chronic exposures to hypobaric hypoxia (i.e. the 424 425 natural high-altitude environment), a complex integrative response to hypoxia will also include haematological and extended nephrological compensation in addition to regulation by 426 427 arterial blood gases. Consequently, the effect of respiratory alkalosis on CBF, metabolism and cognitive function is likely to be influenced by the degree of hypoxic ventilatory 428 response and renal compensation during acclimatization. Similarly, haemoglobin increases 429 occur within weeks of high altitude exposure and improve CaO₂ and global oxygen delivery 430 (Subudhi et al., 2014). Therefore, cognitive impairment to tasks involving sustained attention 431 (i.e. tasks involving reaction time) often occur during the initial exposure to high altitude 432 (4,350m and 5,050m), but are reversed within the days following acclimatization (Davranche 433 et al., 2016; Pun et al., 2018). 434

435 <u>Methodological Considerations</u>

An important consideration to acknowledge is that during EH PETCO2 was not matched to the 436 changes in P_{ET}CO₂ induced during PH (i.e. PETCO₂ significantly different between PH and 437 EH conditions). Our aim was to elicit a hypocapnic state that resembled the one that results 438 from the natural hyperventilation caused by hypoxia, but in reality we overestimated this 439 response when selecting the target P_{ET}CO₂ in EH. This could have been avoided if all 440 participants had undertaken the EH condition after the PH condition, but of course this would 441 442 then introduce a problematic order effect. Nevertheless, studies report that there is a linear graded response of cerebral saturation with carbon dioxide tensions (Mutch et al., 2013), and 443 444 so mechanisms by which hypocapnia induces cognitive impairment may also work in a graded fashion. 445

Active hyperventilation is attention consuming when compared to passive hyperventilation 446 (Gallego, Perruchet, & Camus, 1991), and subsequently may confound any interpretation of 447 hypocapnia on cognitive functioning. To overcome this, previous studies have assessed 448 cognitive function during the minutes of recovery from hyperventilation-induced hypocapnia 449 (Van Diest et al., 2000). However, our battery of cognitive tasks took approximately 15 450 minutes to complete, which was too long to use such an approach. Indeed, Malatino and 451 colleagues demonstrated that MCAv returns to near baseline values within five minutes 452 following hyperventilation-induced hypocapnia, and this was from a greater level of 453 hypocapnia (P_{ET}CO₂=20 mm Hg) than induced in the current study (Malatino et al., 1992). 454 Nonetheless, completing a normocapnic normoxic hyperventilation trial would determine the 455 effect of active hyperventilation on the cognitive function. Transcranial Doppler measures 456 blood flow velocity as an index of vessel blood flow based on the assumption that the 457 diameter of the MCA remains constant. This assumption has recently been questioned 458 (Ainslie & Hoiland, 2014) and evidence for altered MCA diameter in conditions where blood 459 gas content is affected has been demonstrated (Coverdale, Gati, Opalevych, Perrotta, & 460

Shoemaker, 2014; Verbree et al., 2014; Wilson et al., 2011). Nevertheless, if the diameter of
the MCA was increased (in IH) or decreased (EH) as a consequence of the manipulated blood
partial pressures, our TCD-based findings would only underestimate the true effect observed
here.

As mentioned above, we only measured prefrontal cerebral haemodynamic changes with our 465 NIRS and so the regional perfusion shifts proposed would need to be confirmed via whole-466 head NIRS imaging (or with functional magnetic resonance imaging). Further, NIRS is 467 limited to the cortex surface and currently available technology and analysis approaches does 468 not differentiate between skin and skull blood flow, and cerebrospinal fluid. However, 469 470 despite its spatial limitations NIRS is clearly able to measure changes in haemodynamic responses, and which are more likely to result from neural activation than haemoglobin 471 content shift within the blood vessels of the skin under this experimental paradigm (Davies et 472 473 al., 2016; Davies et al., 2017). Finally, these apparatuses only reflect global CBF and regional haemoglobin content, representing vascular flow and oxygenation changes to our measured 474 475 areas of interest. Neither of these imaging devices provided any measure of cerebral metabolic rate of oxygen, which may better reflect the mechanisms of cognitive (dys)function 476 during hypoxia and hypocapnia exposure and warrants future study. 477

478 <u>Conclusion</u>

Hyperventilation-induced hypocapnia impairs performance of simple and five-choice reaction time tasks during normoxia and hypoxia, but not working memory cognitive performance. Furthermore, supplementation of carbon dioxide during hypoxia preserved cognitive function and facilitates an appropriate vascular response. The associated changes in global cerebral haemodynamics between the experimental conditions may mediate this effect, with the changes in MCAv moderately correlated to cognitive task performance. Taken together, these

- findings reveal the significant role of hypocapnia *per se* on the development of cognitive
- 486 impairment during normoxic *and* hypoxic exposures.

Author Contributions: GB and SL conceived and designed the study. All authors 487 contributed to the acquisition, analysis, or interpretation of data for the work. AF and SL 488 drafted the manuscript, with GB reviewing and providing critical feedback important for 489 intellectual content. All authors have approved the final version of the paper and agree to be 490 accountable for all aspects of the work in ensuring that questions related to the accuracy or 491 integrity of any part of the work are appropriately investigated and resolved. All persons 492 designated as authors qualify for authorship, and all those who qualify for authorship are 493 listed. 494

Acknowledgements: We would like to thank the participants for their time and effort in this
study. We also thank Alex Peart, Shaun Webster, Matthew Holloway, Jamie Phillips,
Timothy Riviere, Sebastian Cox, Ciaran O'Connell, Daniel Slater, Josh Kelly and Patrick
Gravett-Curl for their contribution to data collection.

549 **Funding:** No funding received.

550 **Conflict of interest:** There authors declare that they have no conflicts of interest.

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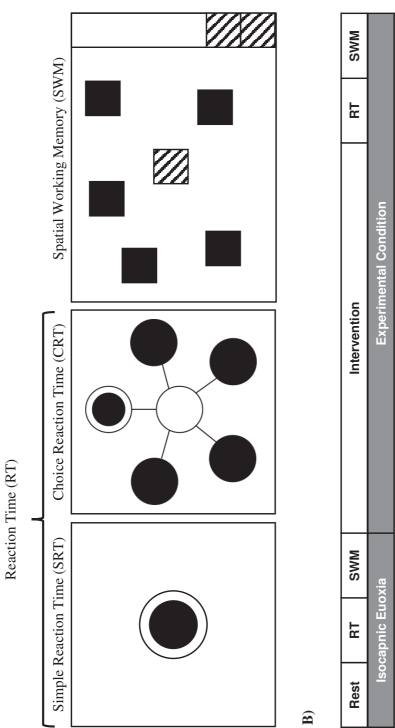
			HR (bpm)		MAP (mmHg)	Ag)	(c V	MCAv (cm·s ⁻¹)		TOI (%)		nTHI (au)	Pr (mi	PETO ₂ (mmHg)	PETCO ₂ (mmHg)	2
	Isocapnic Euoxic (n=20)	~														
	Isocapnic Hypoxia	67.3	± 10.8		82.4 ±	= 15.7	65.4	± 11.8					98.9	± 3.9	41.1 ± 2	2.1
	Poikilocapnic Hypoxia	68.1	± 11.3		87.6 ±	= 14.7	64.9	± 12.2					98.8	± 4.2	41.5 ± 2	2.0
	Subgroup (n=10)															
	Isocapnic Hypoxia	60.9	± 7.2		77.4 ±	= 10.9	61.6	± 12.2	74.9	± 4.9	0.98	± 0.08	9.66	± 4.0	40.8 ± 1	1.7
	Poikilocapnic Hypoxia	63.4	± 8.2		82.3 ±	= 11.8	60.9	\pm 11.0	73.7	± 3.7	1.00	± 0.06	100.8	± 3.8	41.2 ± 2	4
	Euoxic Hypocapnia	63.4	± 8.7		77.4 ±	= 12.7	61.9	± 9.9	73.3	± 3.8	1.00	± 0.05	97.0	± 3.3	41.6 ± 1	ς;
	Experimental (n=20)															
	Isocapnic Hypoxia	68.9	± 10.3		90.8 ±	= 14.0	72.2	$\pm 11.8^{a^{**}}$					43.6	$\pm 1.7^{**}$	41.4 ± 2	2.2 ^α
	Poikilocapnic Hypoxia	71.0	± 10.6		86.4 ±	= 10.3	67.3	\pm 11.0					42.2	$\pm 2.9^{**}$	39.0 ± 3	3.2^{**}
	Subgroup (n=10)															
	Isocapnic Hypoxia	63.9	± 6.3		87.5 ±	= 14.4	68.5	$\pm 10.8^{\alpha\beta*}$	66.3	$\pm 4.6^{\beta^{**}}$	1.02	$\pm 0.09^{\beta*}$	43.4	$\pm 1.8^{\beta^{**}}$	41.3 ± 2	$2.1^{\alpha\beta}$
	Poikilocapnic Hypoxia	65.1	± 8.6		84.2 ±	± 9.9	62.2	± 9.1	64.7	$\pm 4.8^{\beta^{**}}$	1.03	$\pm 0.08^{\beta}$	42.5	$\pm 3.6^{\beta**}$	37.9 ± 3	$3.5^{\beta^{**}}$
	Euoxic Hypocapnia	61.0	± 9.4		85.6 ±	= 11.3	51.7	$\pm 7.6^{**}$	68.8	$\pm 4.1^{**}$	0.95	$\pm 0.05^{**}$	97.0	± 3.6	33.3 ± 1	1.5^{**}
731	Table 1. Absolute resting values for cerebral haemodynamics and end-tidal respiratory gases during isocapnic euoxic baseline and experimental	sting v	alues fo	or cere	sbral h	laemody	/namic	s and end-	tidal re	spiratory	gases c	luring isoca	pnic eu	oxic basel	line and exp	erimental
732	conditions. Experimental conditions were isocapnic hypoxia (IH), poikilocapnic hypoxia (PH), and euoxic hypocapnia (EH). Data are presented	ental cc	ndition	IS Wer	e isoc	apnic hy	/poxia	(III), poik	ilocapr	nic hypox	ia (PH)	, and euoxic	s hypoc	apnia (EH	l). Data are	presented
733	for the group which completed IH and PH conditions $(n = 20)$, and for the subgroup which completed the additional EH condition $(n = 10)$.	compl	eted IH	and	PH co	ndition	s (n =	20), and f	or the	subgroup	which	completed	the add	itional EF	H condition	(n = 10)
734	Significance notation represents differences between data pooled across four measured time points during each IE and experimental period. * p <	ו repres	ents di	fferen	ses be	tween d	ata po(oled across	four n	neasured	ime po	ints during	each IE	and exper	rimental per	iod. * p <
735	0.05 compared to IE. ** $p < 0.001$ compared to IE. $\alpha_{\rm J}$	> d ** .	¢ 0.001	comp	ared to	o IE. α I	0.0.0	5 compare	d to PF	I. $\beta p < 0$.05 con	$p < 0.05$ compared to PH. β $p < 0.05$ compared to EH. HR, Heart rate; MAP, Mean arterial	H. H.R., J	Heart rate	; MAP, Me	an arteria
736	pressure; MCAv, Middle cerebral artery velocity; TOI, Total oxygenation index; nTHI, Total haemoglobin index normalised to initial value;	iddle c	erebral	artery	velo	city; TC	JI, Tot	al oxygené	ttion ir	ıdex; nTF	II, Toté	ıl haemoglc	bin ind	ex norma	lised to ini	tial value
737	PETO ₂ , End-tidal partial pressure of oxygen; PETCO ₂ , End-tidal partial pressure of carbon dioxide Values are Mean ± SD	tial pre:	ssure of	oxyg	en; P _E	TCO ₂ , F	Ind-tid	al partial p	ressure	s of carbo	n dioxia	le Values a	re Mean	ı ± SD.		
																30

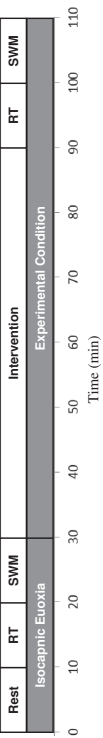
		SRT	CRT	T	SWM
	Time (ms)	Error Count	Time (ms)	Error Count	Error Count
Isocapnic Euoxic (n=20)					
Isocapnic Hypoxia	573 ± 55	0.3 ± 0.5	590 ± 60	0.8 ± 0.7	7.9 ± 12.8
Poikilocapnic Hypoxia	552 ± 52	0.4 ± 0.5	583 ± 74	0.9 ± 1.3	9.5 ± 12.9
Subgroup (n=10)					
Isocapnic Hypoxia	550 ± 47	0.3 ± 0.5	579 ± 58	0.8 ± 0.6	8.1 ± 15.9
Poikilocapnic Hypoxia	532 ± 46	0.6 ± 0.5	562 ± 72	1.4 ± 1.5	7.3 ± 15.1
Euoxic Hypocapnia	544 ± 31	0.5 ± 0.7	588 ± 69	0.6 ± 0.8	5.6 ± 11.8
Experimental (n=20)					
Isocapnic Hypoxia	575 ± 54	0.5 ± 0.6	600 ± 75	0.8 ± 0.8	8.1 ± 17.5
Poikilocapnic Hypoxia	$700 \pm 85^{\delta^{**}}$	0.3 ± 0.4	$735 \pm 86^{\delta **}$	0.6 ± 0.8	11.5 ± 13.3
Subgroup (n=10)					
Isocapnic Hypoxia	566 ± 51	0.3 ± 0.5	594 ± 70	1.0 ± 0.8	6.4 ± 14.0
Poikilocapnic Hypoxia	$721 \pm 51^{\delta^{**}}$	0.3 ± 0.5	$765 \pm 47^{8**}$	0.6 ± 1.0	9.8 ± 17.2
Euoxic Hypocapnia	$718 \pm 55^{\delta **}$	0.6 ± 0.8	$755 \pm 34^{\delta^{**}}$	0.2 ± 0.6	13.0 ± 15.5

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poikilocapnic hypoxia (PH) and euoxic hypocapnia (EH). Data have been presented for the group (n=20) which completed the IH and PH Table 2. Performance time and error count for simple reaction time (SRT) and five-choice reaction time (CRT) tasks, and error count for spatial working memory (SWM) task during isocapnic euoxic and experimental conditions. Experimental conditions were isocapnic hypoxia (IH), conditions, and for the subgroup (n=10) which completed the additional EH condition. ** p < 0.001 compared to IE. $\delta p < 0.001$ compared to IH. Values are Mean \pm SD 740 739 743 741 742

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