

The influence of short-term high-altitude acclimatization on cerebral and leg tissue oxygenation post orthostasis

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1	The influence of short-term high-altitude acclimatization on cerebral and leg tissue
2	oxygenation post orthostasis
3	
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17 Abstract

18

19	syncope. High altitude reduces brain and peripheral muscle tissue oxygenation. This study
20	determined the effect of short-term altitude acclimatization on cerebral and peripheral leg
21	tissue oxygenation index (TOI) post-orthostasis.
22	Method: Seven lowlanders completed a supine-to-stand maneuver at sea level (450 m) and
23	for 3 consecutive days at high altitude (3,776 m). Cardiorespiratory measurements and near-
24	infrared spectroscopy-derived oxygenation of the frontal lobe (cerebral TOI) and vastus
25	lateralis (leg TOI) were measured at supine and 5 min post-orthostasis.
26	Results: After orthostasis at sea level, cerebral TOI decreased (mean Δ % [95% CI]: -4.5%, [-
27	7.5, -1.5], <i>P</i> < 0.001) whilst leg TOI was unchanged (-4.6%, [-10.9, 1.7], <i>P</i> = 0.42). High
28	altitude had no effect on cerebral TOI following orthostasis (day 1 to 3: -2.3%, [-5.3, 0.7]; -
29	2.4%, [-5.4, 0.6]; -2.1%, [-5.1, 0.9], respectively, all $P > 0.05$) whereas leg TOI decreased
30	(day 1 to 3: -12.0%, [-18.3, -5.7]; -12.1%, [-18.4, -5.8]; -10.2%, [-16.5, -3.9], respectively, all
31	P < 0.001). This response did not differ with days spent at high altitude, despite evidence of
32	cardiorespiratory acclimatization (increased peripheral oxygen saturation [supine: $P = 0.01$;
33	stand: $P = 0.02$] and decreased end-tidal carbon dioxide [supine: $P = 0.003$; stand: $P = 0.01$]).

Purpose: Orthostasis at sea level decreases brain tissue oxygenation and increases risk of

34	Conclusion: Cerebral oxygenation is preferentially maintained over leg oxygenation post-
35	orthostasis at high altitude, suggesting different vascular regulation between cerebral and
36	peripheral circulations. Short-term acclimatization to high altitude did not alter cerebral and
37	leg oxygenation responses to orthostasis.
38	Keywords: altitude, blood pressure, heart rate, hypoperfusion, hypoxia, tissue oxygenation
39	
40	Declarations
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44	Conflicts of interest/Competing interests (include appropriate disclosures)
45	No conflict of interest, financial or otherwise, are declared by the authors.
46	Ethics approval and consent to participate
47	All study procedures were approved by the ethical committee of the Mount Fuji Research
48	Institute in Japan (ECMFRI-01-2014) and performed in accordance with the Declaration of
49	Helsinki 2013, with written informed consent obtained from all study participants.

50 Availability of data and material (data transparency)

51 All relevant data are within the paper. The data that support the findings of this study are

52 available from the corresponding author upon reasonable request.

53 **Code availability (software application or custom code)**

54 Not applicable.

55 Authors' contributions

- 56 The M.H., K.A., and K.O. conceived and designed the study. M.H., K.A., and K.O.
- 57 performed the experiments. M.H., K.O., A.T.F., G.M.K.R., and S.J.O. analyzed data and
- 58 interpreted results. M.H., G.M.K.R., and A.T.F. prepared tables and figures. M.H. drafted the
- 59 first manuscript. M.H., K.A., K.O., A.T.F., G.M.K.R., and S.J.O. critically revised the
- 60 manuscript, and all authors approved the final version of the manuscript.

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- 67 ANOVA: Analysis of variance
- 68 BP: blood pressure
- 69 DBP: diastolic blood pressure
- 70 HR: Heart rate
- 71 MAP: Mean arterial pressure
- 72 NIRS: near-infrared spectroscopy
- 73 P_{ET}CO₂: partial pressure of end tidal carbon dioxide
- 74 SBP: Systolic blood pressure
- 75 SD: standard deviation
- 76 SpO₂: peripheral arterial oxygen saturation
- 77 TOI: tissue oxygenation index

78 Introduction

79 In the upright stand position the cerebral arteries are positioned above the heart resulting in a 80 hydrostatic arterial pressure gradient between the heart and the brain, causing a reduction in 81 cerebral perfusion pressure (Rosner and Coley 1986). Consequently, the brain is prone to 82 incidents of hypoperfusion that may lead to orthostatic intolerance (Van Lieshout et al. 2003). 83 Orthostatic tolerance is reduced during exposure to severe (10% O₂) normobaric hypoxia (Rowell and Seals 1990) and to hypobaric hypoxia at moderate (ca. 2,800 m) (Nicholas et al. 84 85 1992), and high (ca. 6,000 m) altitudes (Westendorp et al. 1997). As impaired orthostatic tolerance is associated with syncope or collapse (Van Lieshout et al. 2003), this is of concern 86 to mountaineers and others working and visiting high altitude, for whom falls may be life 87 threatening (Firth et al. 2008). Short-term acclimatization at high altitude could be a strategy 88 to reduce orthostatic intolerance. However, evidence of high-altitude acclimatization effect on 89 90 orthostatic intolerance is limited. 91 To evaluate tolerance to orthostasis, measurements of cerebral tissue oxygenation 92 index (TOI) by near-infrared spectroscopy (NIRS) have been commonly used (Mehagnoul-

Schipper et al. 2000; Mehagnoul-Schipper et al. 2003). Cerebral TOI is determined by arterial
oxygen content, oxygen consumption, and total blood volume at the target tissue (i.e., directly
below the probes). Moreover, previous research has suggested that cerebral TOI is a sensitive

96	and robust indicator of orthostatic intolerance (Kuriyama et al. 2000). Although it is well
97	established that a common cause of syncope is initial (< 1 min) orthostatic hypotension
98	(Wieling et al. 2007), the potential risk of syncope during the later phase (> 1 min) should
99	also be considered. Previous studies have demonstrated that blood pressure (BP) and middle
100	cerebral artery blood flow velocity fall acutely from supine to stand within 10 s, but recovers
101	within 30 s (Thomas et al. 2010; Thomas et al. 2009; van Lieshout et al. 2001). However,
102	cerebral oxygenation remains lower after 5 min of stand (van Lieshout et al. 2001), suggesting
103	a possibility of syncope incidents even after initial orthostatic hypotension phase. Moreover, it
104	has been suggested that orthostatic intolerance is fundamentally due to a critical fall in
105	cerebral perfusion, rather than systemic BP per se (Van Lieshout et al. 2003). Thus,
106	evaluations of cerebral TOI and BP after initial orthostatic hypotension phase could provide
107	further understanding to orthostasis.
108	Orthostatic tolerance is positively related to the strength of sympathetic-driven
109	peripheral vasoconstriction (Brown and Hainsworth 2000; Claydon and Hainsworth 2004).
110	Enhanced muscle sympathetic nerve activity is accompanied by reductions in peripheral
111	oxygenation during lower body negative pressure during normoxia (Hansen et al. 2000;
112	Hansen et al. 1996; Vongpatanasin et al. 2011) and hypoxia (Hansen et al. 2000). Therefore,
113	the reduced peripheral oxygenation reported to orthostasis may be part of an adaptive
114	response to maintain cerebral oxygenation and prevent orthostatic intolerance. This study

115	aimed to investigate the cerebral and peripheral tissue oxygenation response to orthostasis at
116	sea level and high altitude. We hypothesized that with acute exposure to high altitude (day 1),
117	following the initial hypotension phase (5 min post-orthostasis) cerebral tissue oxygenation
118	would be maintained at the expense of peripheral tissue oxygenation. Secondly, we
119	hypothesized that following short-term high-altitude acclimatization (3 days) there would be
120	an attenuated reduction in peripheral oxygenation post-orthostasis compared to acute
121	exposure (day 1).
122	
123	Methods
124	Participants
125	This study was approved by the ethical committee of the Mount Fuji Research Institute in
126	Japan (ECMFRI-01-2014) and performed in accordance with the <i>Declaration of Helsinki</i>
127	vapan (2011) til of 2011) and performed in accordance with the Decian anon of Heisinna
	2013, with written informed consent obtained from all study participants. Seven healthy male
128	2013, with written informed consent obtained from all study participants. Seven healthy male lowlanders [mean \pm standard deviation (SD): age 46 \pm 15 years; height 173 \pm 6 cm; body
128 129	2013, with written informed consent obtained from all study participants. Seven healthy male lowlanders [mean \pm standard deviation (SD): age 46 \pm 15 years; height 173 \pm 6 cm; body mass 68 \pm 5 kg] voluntarily participated in this study. All participants were free from
128 129 130	2013, with written informed consent obtained from all study participants. Seven healthy male lowlanders [mean \pm standard deviation (SD): age 46 \pm 15 years; height 173 \pm 6 cm; body mass 68 \pm 5 kg] voluntarily participated in this study. All participants were free from cardiovascular disease, were not taking medications, and did not engage in regular exercise.
128 129 130 131	2013, with written informed consent obtained from all study participants. Seven healthy male lowlanders [mean \pm standard deviation (SD): age 46 \pm 15 years; height 173 \pm 6 cm; body mass 68 \pm 5 kg] voluntarily participated in this study. All participants were free from cardiovascular disease, were not taking medications, and did not engage in regular exercise. Additionally, none of the participants had been exposed to an altitude higher than 1,500 m

strenuous physical activity and alcohol for 24 h, and from caffeinated beverages for 12 h.
They also abstained from strenuous exercise, alcohol and caffeine for the following four study
days.

136

137 Study locations and procedures

138 Measurements were taken at sea level followed by three consecutive days at high altitude 139 (3,776 m; day 1, day 2, and day 3). The sea level study was conducted at the Mount Fuji 140 Research Station (450 m above sea level, ambient barometric pressure ca. 720 mmHg), and 141 the high altitude study at the Mount Fuji Summit Hut (3,776 m above sea level, ambient 142 barometric pressure ca. 490 mmHg, Figure 1). All participants were familiarized with the 143 experimental protocol and supine-to-stand maneuver before beginning the study. Sea level measurements were performed 2 weeks before ascent to high altitude. On the day of ascent, 144 145 all participants reached the Self-Defense Forces base camp in the morning by vehicle (1,280 m above sea level, ambient barometric pressure ca. 655 mmHg). Participants ascended to the 146 147 top of Mount Fuji (3,776 m) within 3 h by riding on a bulldozer, arriving at approximately 9:00 AM. All studies were performed between 14:00 and 17:00. After three nights at high 148 149 altitude the participants walked down the mountain and returned to sea level.

151 *Experimental protocol and measurements*

Each participant rested in the supine position for 30 min before they were asked to stand quickly and assume an erect and immobile posture. Participants were requested not to speak, to breathe normally, and to remain as still as possible to reduce any influence of the skeletal muscle pump when in the stand position. Stand position was determined once the participant maintained a stable balance.

157	Heart rate (HR), peripheral arterial oxygen saturation (SpO ₂), and partial pressure of
158	end tidal carbon dioxide (P _{ET} CO ₂) were continuously measured during supine and stand.
159	P _{ET} CO ₂ was measured in 5 participants due to equipment fault. HR was recorded using a
160	commercial HR monitor (Polar RS800CX, Polar Electro Japan, Tokyo, Japan). SpO2 was
161	monitored by finger pulse oximetry (PULSOX-300i; Konica Minolta, Tokyo, Japan) on the
162	right index finger. P _{ET} CO ₂ was measured using a CO ₂ monitor (OLG-2800, Nihon Kohden,
163	Tokyo, Japan). Systolic BP (SBP) and diastolic BP (DBP) were measured at 1-min intervals
164	during supine and stand using the oscillometric method on the upper left arm (HEM-7200;
165	Omron, Tokyo, Japan).
166	Cerebral and peripheral hemodynamics were measured continuously using NIRS

168 maneuver (Al-Rawi et al. 2001). NIRS has been utilized for continuous monitoring of

167

(NIRO-300; Hamamatsu Photonics KK, Hamamatsu, Japan) throughout the supine-to-stand

169	cerebral oxygenation by evaluating the concentrations of oxyhemoglobin and
170	deoxyhemoglobin at the measurement site using spatially resolved spectroscopy (Houtman et
171	al. 1999; Mehagnoul-Schipper et al. 2000; Mehagnoul-Schipper et al. 2003). TOI provides a
172	measure of tissue oxygen saturation and is calculated as the ratio of oxygenated to total tissue
173	hemoglobin [TOI = oxyhemoglobin / total hemoglobin (oxyhemoglobin +
174	deoxyhemoglobin)]. A probe holder containing an emission probe and detection probe was
175	attached to the right side of the forehead (with a distance of 3 cm between the probes) to
176	measure TOI at the frontal lobe (cerebral TOI). Two further probes were attached to the lower
177	third of the vastus lateralis muscle (10–12 cm above the knee joint) to measure peripheral
178	oxygenation (leg TOI) (Koga et al. 2007). Pen marks were made on the skin to indicate the
179	margins of the probe holder and electrodes so that the probe could be positioned at exactly the
180	same place each day.
181	

182 Data analysis

Continuous measurements of cerebral TOI, leg TOI, HR, P_{ET}CO₂, and SpO₂ were averaged from the final 5 min of supine rest and from the final 10 s of standing following the maneuver (i.e., 4'50"–5'00" min). Incremental measurements of SBP and DBP were taken every minute at supine rest with the average reported. During standing, one measurement of SBP and DBP

187	was taken 5 min following the maneuver. Mean arterial pressure (MAP) was calculated:
188	[MAP = (SBP-DBP)/3+DBP]. To evaluate the effects of high-altitude acclimatization (i.e.,
189	from day 1 to day 3 at high altitude) to orthostasis, cerebral TOI and leg TOI are presented as
190	the percent change from supine to stand for each day spent at high altitude.
191	
192	Statistical analysis
193	A sample size estimation for the primary analysis (cerebral TOI) indicated that 4 participants
194	were needed to produce an 80% chance of obtaining statistical significance at the 0.05 level
195	(G Power 3.1) (Faul et al. 2009) for a meaningful Cohen's F effect size of 0.92 for a 2×4
196	repeated measures analysis of variance (ANOVA). The effect size was calculated based on a
197	minimum important difference of 4.4% determined by the difference in cerebral TOI between
198	orthostatic symptomatic and asymptomatic persons (Harms et al. 2000), a standard deviation
199	of the difference of 2.4% from the same study (Harms et al. 2000), and a correlation of
200	repeated measures of $r = 0.76$, based on data from a previous study (Al-Rawi et al. 2001). A
201	minimum 6 participants were set as the target for recruitment to account for possible dropout.
202	All data were expressed as mean \pm SD. Statistical analysis was performed using GraphPad
203	Prism 7 commercial software (MDF Co., Ltd, Tokyo, Japan), with statistical significance
204	accepted at $P < 0.05$. Two-way repeated measures ANOVA with Bonferroni post-hoc tests

205	were used to assess differences between day (sea level, day 1, day 2, and day 3) and posture
206	(supine and stand) for all cardiorespiratory and oxygenation measures. One-way repeated-
207	measures ANOVA was used to determine differences in the percentage change in cerebral
208	TOI and leg TOI between days. To assess the effect of acclimatization to high altitude, one-
209	way repeated-measures ANOVA (day 1 to day 3) with linear trend analysis were conducted
210	on cardiorespiratory and oxygenation measures. This is because we sought to observe the
211	overall slope and fit of the response in physiological responses during sojourn at high altitude.
212	This linear trend analysis approach was particularly advantageous to reduce the number of
213	comparisons made (Horiuchi et al. 2016; Horiuchi et al. 2017).

215 **Results**

216 *Cardiorespiratory variables*

217 Supine cardiorespiratory variables demonstrated expected responses to acute hypoxia and

218 provide evidence of altitude acclimatization over the three days. Specifically, compared to sea

219 level, SpO₂ was lower on day 1 at high altitude and then increased on subsequent days at

altitude (**Table 1**). Further, linear trend analysis identified that SpO₂ (P = 0.01) and MAP (P < 0.01)

221 0.001) increased, and $P_{ET}CO_2$ (P = 0.003) decreased, linearly with sojourn at high altitude.

HR also tended to linearly increase with days at altitude (P = 0.06, **Table 1 and 2**). The

223 supine-to-stand maneuver increased MAP on day 2 at high altitude (P = 0.02), HR at sea level and all days at high altitudes (all P < 0.05), SpO₂ on all days at high altitude (all P < 0.05), 224 225 and decreased $P_{ET}CO_2$ on day 1 (P = 0.02) and day 2 (P = 0.01) at high altitude (**Table 1**). 226 227 Cerebral tissue oxygenation index 228 Compared to sea level, supine cerebral TOI decreased on day 1 at high altitude (P < 0.01, 229 Figure 2A), and did not recover during the sojourn at high altitude (P = 0.36, Table 2). 230 Cerebral TOI was lower after the supine-to-stand maneuver at sea level (supine-to-stand mean Δ % [95% CI]: - 4.5%, [-7.5, -1.5], P < 0.001, Figure 2A and 2C). In contrast, altitude had no 231 232 effect on cerebral TOI after the supine-to-stand maneuver (day 1 Δ %: -2.3%, [-5.3, 0.7], P = 233 0.34; day 2 -2.4%, [-5.4, 0.6], P = 0.22; day 3 -2.1%, [-5.1, 0.9], P = 0.49; Figure 2A and 234 **2C**). Linear trend analysis of day 1 to 3 revealed that short-term acclimatization did not alter 235 cerebral oxygenation change to orthostasis (P = 0.86). 236

237 Peripheral leg tissue oxygenation index

Supine leg TOI was unchanged at high altitude compared to sea level (P = 0.32, Figure 2B

and Table 2). Leg TOI was maintained after the supine-to-stand maneuver at sea level (Δ % -

4.6% [-10.9, 1.7], P = 0.42, **Figure 2B and 2D**). In contrast, leg TOI was lower after the supine-to-stand maneuver on all days at high altitude (day 1 Δ %: -12.0%, [-18.3, -5.7]; day 2 -12.1%, [-18.4, -5.8]; day 3 -10.2%, [-16.5, -3.9]; all P < 0.001, **Figure 2B and 2D**). Linear trend analysis of day 1 to 3 revealed that short-term acclimatization did not alter leg oxygenation change to orthostasis (P = 0.37).

245

246 **Discussion**

247	The principle finding of this study is that after orthostasis at high altitude cerebral TOI was
248	protected against the reduction that was observed at sea level. Indeed, the non-significant
249	mean difference in cerebral TOI after orthostasis at high altitude can be considered trivial as it
250	was less than the minimum important difference (4.4%) that was calculated from the
251	difference in cerebral TOI between orthostatic symptomatic and asymptomatic persons
252	(Harms et al. 2000). A reduction in cerebral TOI (4.5%) was achieved after orthostasis at sea
253	level in this study. Peripheral (leg) TOI was in contrast reduced after orthostasis at high
254	altitude but not at sea level. These data highlight that cerebral oxygenation is preferentially
255	maintained compared to leg oxygenation during orthostasis at high altitude. This reciprocal
256	response was unchanged during the 3-day high altitude sojourn that led to cardiorespiratory
257	altitude acclimatization adaptations including a progressive recovery of SpO ₂ . These results

indicate a different vascular regulation between the cerebral and peripheral circulations toorthostasis during short-term high-altitude acclimatization.

260	In agreement with previous studies (Cheung et al. 2014; Sanborn et al. 2015), high
261	altitude hypoxia reduced cerebral TOI during supine rest compared to sea level (Figure 2A).
262	Since TOI is calculated as the ratio of oxyhemoglobin to total tissue hemoglobin (Al-Rawi et
263	al. 2001), changes in TOI could be due to alterations in cerebral blood flow (oxygen delivery)
264	or oxygen extraction. Oxyhemoglobin is mainly included in the artery and cerebral blood flow
265	(arterial inflow) increases within the first 6-12 hours at high altitude, remaining elevated for
266	several days compared with sea level as first reported (Severinghaus et al. 1966). Subsequent
267	studies have confirmed these results (Jensen et al. 1990; Lucas et al. 2011; Subudhi et al.
268	2014; Willie et al. 2014). Thus, TOI reductions in the present study may be explained by
269	increases in deoxyhemoglobin in the brain that is consistent with a previous research (Cheung
270	et al. 2014). Indeed, it was reported that hypoxia causes an increase in the volume of cerebral
271	deoxyhemoglobin by increasing oxygen extraction (Rasmussen et al. 2007). The absence of a
272	further reduction in cerebral oxygenation to orthostasis suggests that the cerebrovascular
273	perfusion is preferentially maintained compared to peripheral perfusion e.g. leg TOI. There
274	are several possibilities to explain these results. We found significant reductions in the leg
275	TOI 5 min post-orthostasis. This may indicate that sympathetic-induced vasoconstriction
276	occurred at the peripheral arteries, which could be an adaptive response to ensure the

277	maintenance of cerebral TOI. Indeed, during 3-day acclimatization at high altitude, MAP,
278	which is observed along with increases in muscle sympathetic nerve activity (Hansen et al.
279	1996), progressively increased irrespective of posture ($P < 0.05$, respectively, Table 1 and 2).
280	Combined, this suggests the maintenance of cerebral TOI to orthostasis at high altitude is
281	dependent on peripheral vasoconstriction to maintain MAP, with compromising consequences
282	for oxygenation of peripheral tissues. These interpretations are supported by a previous study
283	that has reported reductions in peripheral oxygenation with enhanced muscle sympathetic
284	nerve activity during lower body negative pressure (i.e., simulated orthostasis) in hypoxia
285	(Hansen et al. 2000).
286	The absence of a further decrease in cerebral oxygenation after orthostasis at high
287	altitude may be due to a redistribution of cardiac output (cardiac output = stoke volume \times HR)
288	since changes in cerebral oxygenation to orthostasis is also cardiac output dependent (van
289	Lieshout et al. 2001). Stroke volume decreases during short-term (5 days) exposure high
290	altitude (Kanstrup et al. 1999), and hence, increased HR compensates to maintain cardiac
291	output for sufficient oxygen delivery to peripheral tissues. Indeed, HR at high altitude was

significantly higher than sea level and the supine-to-stand maneuver significantly increased
HR throughout the days. Thus, it is also possible that the increase in HR from supine to stand
could compensate to maintain cardiac output and support the maintenance of cerebral TOI.

295 However, we acknowledge that this hypothesis is speculative and warrants future

296 investigation with a measurement of stroke volume and cardiac output.

In the present study, cerebral TOI did not recover, but SpO₂ progressively increased irrespective of posture during the sojourn at high altitude. While SpO₂ assessed by pulse oximeter has been widely used to evaluate systemic hypoxemia (Kohyama et al. 2015), our data and that of others (Sanborn et al. 2015) demonstrate that SpO₂ may not represent cerebral oxygenation. The finding that cerebral TOI responses are divergent from peripheral (leg) TOI and SpO₂ not only has implications for our understanding of physiological responses to high altitude, but also for future research design in the field.

304

305 Methodological considerations

NIRS was used to provide a non-invasive measure of tissue oxygenation. As near infrared light passes through skin before absorption into the tissue the potential exists for blood flow outside of the tissue to influence NIRS derived measurements. A limitation of the present study is that skin blood flow was not measured. Nevertheless, previous research using the same NIRS device as in the present study, reported that a change in cerebral TOI was predominantly associated with internal carotid artery blood flow, and not external carotid artery or skin blood flow during carotid vessel clamping (Al-Rawi et al. 2001). Further, a

313	more recent study demonstrated that cerebral oxygenation during acute hypotension periods
314	in hypoxia (simulated orthostasis) was not associated with skin blood flow (Horiuchi et al.
315	2020). To aid clarity of interpretation future studies should measure where practically
316	possible tissue oxygenation and skin blood flow simultaneously. The present study was
317	completed in a field environment to enable the investigation of high-altitude acclimatization.
318	A limitation of this scenario was that we were not able for logistical reasons to measure other
319	cardiac and cerebrovascular responses, including skin blood flow. These additional measures
320	would have provided a more complete assessment and understanding of the physiological
321	mechanisms that underpin the divergent cerebral and leg oxygenation responses observed.

323 Conclusions

Cerebral oxygenation post-orthostasis at high altitude was protected against the reduction observed at sea level, whereas peripheral (leg) oxygenation was only reduced post-orthostasis at high altitude. This reciprocal response highlights divergent vascular regulation in cerebral and peripheral circulations and may suggest an adaptive response to preferentially maintain cerebral oxygenation during orthostasis at high altitude. Short-term acclimatization to high altitude did not alter the cerebral and peripheral oxygenation response to orthostasis.

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Figure captions

Figure 1. Illustration of the study procedure. TOI; tissue oxygenation index, P_{ET}CO₂; partial

443 pressure of end-tidal carbon dioxide, BP; blood pressure, HR; heart rate, SpO₂; peripheral

444 arterial oxygen saturation.



447	Figure 2. Cerebral total oxygenation index (TOI [%, panel A]), leg TOI (%, panel B), and
448	change in cerebral TOI (Δ %, panel C) and leg TOI (Δ %, panel D) after the supine-to-stand
449	maneuver at sea level (SL) and each day at high altitude (day 1, day 2, and day 3). Data are
450	presented as resting supine (blue circles), standing 5 min after the supine-to-stand maneuver
451	(red squares) and change from supine to stand (green triangles). * $P < 0.05$ supine versus
452	stand for that day, $\dagger P < 0.05$ versus another day for stand posture only, $\ddagger P < 0.05$ versus
453	another day for both postures. Values are presented as mean \pm standard deviation.



		Sea level	Day 1	Day 2	Day 3	Two-way A	NOVA P	values
						Day	Posture	Interaction
MAP, mmHg	Supine	87 (8)	87 (8)	95 (8)	98 (9)*	< 0.001	0.002	0.26
(n=7)	Stand	92 (9)	89 (7)	103 (10)*#	105 (10)*			
HR, bpm	Supine	64 (10)	72 (11)	75 (10)*	76 (15) [*]	0.01	0.001	0.39
(n=7)	Stand	81 (12)#	86 (20)#	87 (16)#	89 (18)#			
SpO ₂ , %	Supine	96 (1)	82 (5)*	84 (6)*	86 (4)*	< 0.001	0.02	0.002
(n=7)	Stand	96 (2)	86 (4)*#	86 (5)*#	88 (4)*#			
P _{ET} CO ₂ , mmHg	Supine	36.7 (3.3)	35.6 (2.2)	35.2 (2.3)	30.6 (4.6)*	0.001	0.001	0.54
(n=5)	Stand	33.6 (1.5)	31.4 (3.0)#	30.4 (2.3) [#]	27.4 (3.4)*			

Table 1. Supine and 5 min post-orthostasis standing cardiorespiratory responses at sea level and for 3 days whilst acclimatizing to high altitude.

Values are mean (standard deviation). MAP, mean arterial pressure; HR, heart rate; bpm, beats per minute; SpO₂, peripheral oxygen saturation; $P_{ET}CO_2$, partial pressure of end-tidal carbon dioxide ^{*}, P < 0.05 vs. sea level at each position. [#], P < 0.05 between supine and stand at that day.

		F values (degree of freedom)	P value
Cerebral TOI	Supine	F (1,12) = 0.92	0.36
	Stand	F (1,12) = 1.64	0.22
Leg TOI	Supine	F (1,12) = 1.09	0.32
	Stand	F (1,12) = 3.48	0.09
MAP	Supine	F (1,12) = 21.53	< 0.001
	Stand	F (1,12) = 17.99	0.001
HR	Supine	F (1,12) = 4.16	0.06
	Stand	F (1,12) = 0.52	0.49
SpO ₂	Supine	F (1,12) = 9.34	0.01
	Stand	F (1,12) = 7.41	0.02
P _{ET} CO ₂	Supine	F (1,8) = 17.04	< 0.01
	Stand	F (1,8) = 9.80	0.01

Table 2. Linear trend analysis of each variable for 3 days whilst acclimatizing to high altitude

TOI, tissue oxygenation index; MAP, mean arterial pressure; HR, heart rate;

458

SpO2, peripheral arterial oxygen saturation; PETCO2, partial pressure of end tidal carbon dioxide.