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1	The effect of High Altitude on Central blood pressure and arterial stiffness
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15	Key words high altitude, central blood pressure, augmentation index, hypoxia
16	
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21	

22	Central arterial systolic blood pressure (SBP) and arterial stiffness are known to be better
23	predictors of adverse cardiovascular outcomes than brachial SBP. The effect of progressive
24	high altitude (HA) on these parameters has not been examined.

25 Ninety healthy adults were included. Central BP and the augmentation index (AI) were

26 measured at the level of the brachial artery (Uscom BP+ device) at <200m and at 3619m,

4600m and 5140m. The average age of the subjects (70% men) were 32.2 ± 8.7 years.

28 Compared with central arterial pressures, brachial SBP (+8.1±6.4 mmHg; p<0.0001) and

29 pulse pressure (+10.9±6.6 mmHg; p<0.0001) were significantly higher and brachial DBP

30 was lower (-2.8±1.6 mmHg; P<0.0001). Compared <200m, HA led to a significant

31 increase in brachial and central SBP. Central SBP correlated with AI (r=0.50; 95% CI:

32 0.41 to 0.58: p<0.0001) and age (r=0.32; 21to 0.41: p<0.001). AI positively correlated with

age (r=0.39; p<0.001) and inversely with subject height (r=-0.22; p<0.0001) weight (r=-

0.19; p=0.006) and heart rate (r=-0.49: p<0.0001). There was no relationship between

acute mountain sickness scores (LLS) and AI or central BP. The independent predictors of

central SBP were male sex (coefficient, t 4.7; P<0.0001), age (t=3.6; p=0.004) and AI

37 (t=7.5; p<0.0001; overall r2 =0.40; p<0.0001). Subject height (t=2.4; p=0.02), age (7.4;

p<0.0001) and heart rate (t=11.4; P<0.0001) were the only independent predictors of AI

39 (overall r2=0.43; p<0.0001). Central BP and AI significantly increase at HA. This rise was

40 influenced by subject-related factors and heart rate but not independently by altitude, LLS

41 or SpO2.

42

44 Introduction

45

(HA).¹ Despite this fact, there has been limited research into cardiovascular risk 46 assessment at HA.¹ HA exposure leads to an increase in resting heart rate, compared with 47 that at sea level, yet paradoxically, maximal heart rate is reduced.² The stroke volume rise 48 noted with exercise at sea level is blunted at HA.^{2,4} Consequently, whilst resting cardiac 49 output is higher at HA, versus sea level, at peak exercise it is comparatively lower.^{2,4,5} 50 These factors along with the notable reduction in arterial oxygen content act to limit peak 51 exercise capacity and oxygen consumption.^{2,5} Other reported cardiovascular responses 52 include an increase in resting brachial artery systolic blood pressure (SBP) and 24hour 53 54 arterial blood pressure (BP), which along with the increase in resting heart rate could be potential implicating factors in the increased cardiovascular risk.⁶⁻⁹ 55 The effects of HA on central arterial haemodynamics, such as central arterial BP and large 56 57 artery stiffness, are far less well understood and have been barely reported. Central arterial 58 BP and large artery stiffness are known to be more powerful predictors of adverse 59 cardiovascular outcomes, including stroke and cardiovascular death than brachial artery BP 60 as they more closely reflect the haemodynamic loading of vital central organs such as the

Cardiovascular death is a leading cause of non-traumatic deaths in adults at high altitude

61 heart, brain and kidneys.^{10.11} Brachial artery BP does not reliably reflect central BP due to

62 the effects of peripheral amplification which is highly variable between individuals.^{10,11}

Unfortunately, the accurate non-invasive assessment of central BP and large artery
stiffness has been traditionally very difficult. It had required the need for either arterial
catheterisation or less portable and expensive non-invasive equipment limiting its research
utility at HA, explaining the paucity of published research at genuine terrestrial HA.^{5.7}

In the only study to investigate the influence of terrestrial HA on both large arterial
stiffness and central BP Parati et al observed a significant increase in both central SBP and
the arterial augmentation index (AI, marker of arterial stiffness) in untreated subjects
travelling to HA.⁷ However, the altitude gain was very rapid (4559m within 28 hours of
ascent) and only a single altitude was studied. Nevertheless, their findings are potentially
important given the huge numbers exposed to HA worldwide.^{1,2}

The Uscom BP⁺ is a novel device which is able to estimate central blood pressure 73 using a simple oscillometric BP cuff on the upper arm.¹² It has shown excellent agreement 74 with catheter based assessments of central BP and gold standard measures of arterial 75 stiffness.¹³¹⁵ It utilises pulse wave analysis to assess the AI which reflects the enhancement 76 77 (augmentation) of central aortic systolic pressure by reflected arterial pulse waves. It has 78 the advantage over several competing devices. It is highly portable and only requires the 79 use of an upper arm cuff therefore avoiding the need to assess either the radial or digital pulse where the signal to noise ratio may be less favourable. 80

In this study we sought to utilise this available technology to investigate, for the first time the effects of a step-wise increasing terrestrial HA on both central BP and AI during a trek to >5000m.

84

85 Methods

86 Study design and participants

87 Ninety healthy British Military servicemen aged >18years were included. Inclusion was 88 entirely voluntary and represented a large subset of military servicemen who had been selected to take part in a quadrennial military adventure training exercise to HA. 89 90 Significant mountaineering experience was not essential but those with very limited experience were encouraged to attend a winter skills course (<1200m) within 3 months of 91 departure. The subjects were assessed at near sea level (<200m) and during progressive 92 93 ascent in the Dhaulagiri region in the Himalayas in March/April 2016. Health status was confirmed following a detailed baseline questionnaire. All subjects were assessed to be 94 95 medically fit for a high altitude venture by their general practitioner. To be considered fit they were all required to have passed their annual military basic fitness test which includes 96 97 a 1.5 mile timed run. Key exclusion criteria included a history of hypertension and/ or atrial fibrillation. All participants were low altitude dwellers and none had prior exposure 98 99 to >1400m terrestrial altitude in the four weeks prior to this study. The subjects were 100 studied consecutively in groups of 8-10 individuals with a two day stagger between 101 successive groups. HA related symptoms were assessed using the Lake Louis Scoring System (LLS).^{16,17} 102

103

104 High Altitude Ascent and descent profile

The subjects flew from the UK to Kathmandu (1400m day 1-3) where they underwent a short period of local acclimatisation at 1400m. From there they travelled by a staged road move to Darbang (1030m) then on foot with loads of up to 12kg over the ensuing 11 days to HA of 5140m (after passing over French pas at 5360m) (figure 1). From there they

commenced trekking on foot over the ensuing 11 days (to day 14) to an altitude of 5140m
(with an overpass of 5360m) before commencing their decent (day 15) on foot to Marpha
(2719m) and then by road back to Kathmandu. Research assessments were performed at
sea level and at static research camps at 3619m (day 9), 4600m (day 12) and 5140m (day
14) during ascent.

114

115 Physiological assessments and central blood pressure measurement

116 Oxygen saturations (SpO₂) were measured using a Nonin Onyx (Nonin Medical Inc, Plymouth, Minnesota, USA) pulse oximeter. Blood pressure and arterial stiffness 117 assessments were obtained at the same time using an Uscom BP⁺ device (Uscom, Sydney, 118 NSW, Australia) as previously reported.¹³⁻¹⁵ The upper arm cuff was attached to the 119 dominant arm of seated subjects. All subjects were rested for at least five minutes prior to 120 BP assessment and they were not permitted to drink caffeine or smoke for at least three 121 hours and alcohol for >10 hours prior to BP measurements.¹⁸ The subjects were advised 122 123 not to speak during the recordings. The BP⁺ device measures both central and peripheral BP (mmHg) using supra systolic oscillometry. Following an initial inflation-deflation the 124 125 cuff is re-inflated to approximately >30mm Hg above the recorded suprasystolic pressure 126 for 10 seconds, during which suprasystolic BP and pulse wave assessments are recorded 127 via the arm cuff. All recordings were stored on a mini SD card within the device and later 128 exported for data analysis. Only readings with a signal-to-noise ratio of ≥ 6 was were included and tests with a ratio of <6 were repeated. 129

The BP⁺ calculates a number of additional haemodynamic indices that were of
interest to this study, including the AI. Its quoted AI is the arterial augmentation pressure
(difference between the second and first systolic peaks of the central pressure waveform)

133	expressed as a percentage of the pulse pressure and it is an <i>indirect</i> measure of large
134	arterial stiffness. Further parameters that we were specifically interested in for this study
135	were the time to systolic wave Reflection (TR) and the suprasystolic pulse pressure
136	variation (ssPPV). The reflected Wave Transit Time is an indirect measure of pulse wave
137	velocity and large arterial stiffness. The ssPPV is a novel measure of fluid responsiveness
138	and is heavily influenced by respiratory variation and left ventricular stroke volume, both
139	of which can be affected at HA. ^{$19-21$} The BP ⁺ calculates the ssPPV as the difference
140	between maximum and minimum pulse pressures divided by the average pulse pressure
141	over the 10 second rhythm strip.

143 Ethics

Participation was entirely voluntary and all participants underwent detailed written
informed consent. The study was approved by the Ministry of Defence Research and
Medical Ethics Committee (MODREC) and was conducted according to the standards of
the declaration of Helsinki.

148

149 Statistical analysis

150 Data were analysed using GraphPad InStat version 3.05 and with all graphical figures

151 presented using GraphPad Prism version 4.00 for Windows (GraphPad Software, San

152 Diego, CA, USA). Sample size calculations were performed using a proprietary

determined sample- size calculator using (GraphPad StatMate version 2.00 for Windows).

154 The Kolmogorov-Smirnov test was undertaken to assess normality of all continuous data

and all continuous data are presented as mean \pm standard deviations and median \pm

156 interquartile range for parametric and non-parametric data respectively. Comparison of unpaired data was performed using an unpaired T test or the Mann-Whitney Test for 157 158 parametric and non-parametric data respectively and with a paired t test and Wilcoxon 159 matched pairs test for equivalent paired data. Continuous data from ≥ 3 groups were compared using a one-way Analysis of Variance (ANOVA) with either Tukey post-hoc 160 tests or a Kruskal-Wallis test with Dunn post-test for parametric and non-parametric data 161 162 respectively. Correlations were performed using Pearson and Spearman rank correlation (±95% confidence interval, CI) for parametric and non-parametric data respectively. A 163 two tailed P value < 0.05 was considered statistically significant for all comparisons. All 164 165 univariate predictors of central arterial systolic blood pressure were entered into a multiple linear regression analysis model in order to identify its independent predictors. A two 166 167 tailed *P* value <0.05 was considered statistically significant for all comparisons.

168 Sample size calculations

Parati et al studied 44 subjects who travelled form sea level to 4559m within 29 hours.⁸ 169 170 From this group there were 22 subjects who were randomised not to receive prophylactic medication to prevent acute mountain sickness. In this group they observed a non-171 172 significant increase in central systolic blood pressure from 103.7±10.7 to 108.8±8.0 mmHg 173 from sea level to that after 48h at HA. The AI significantly increased at HA versus sea 174 level. Based on this data and the average standard deviation of their central BP readings, 175 we calculated that a sample size of at least 60 subjects would have >80% power to detect a >5mmHg change in central SBP and a >7% change in AI at HA at a significance level (alpha) 176 of 0.05 (two-tailed). 177

178

179 **Results**

180	Ninety subjects were included. The average age of the subjects were 32.2±8.7 years with
181	70% being male. Heart rate and LLS increased and SpO ₂ fell at HA compared with sea
182	level (table 1). The average 1.5 mile run time for included subjects was 9.9±1.2 minutes.
183	Overall brachial arterial SBP (+8.4 [5.0 to 12.0] mmHg; p<0.0001) and pulse
184	pressure (+11 [7.0 to 15.0] mmHg; p<0.0001) were significantly greater than that observed
185	centrally. Conversely the brachial artery DBP was lower (-2.6 [-3.4 to -2.0] mmHg;
186	P<0.0001) than the equivalent central readings.
187	Compared with baseline sea level values there was a significant increase in both
188	brachial and central SBP and in brachial but not central arterial pulse pressure at HA (table
189	2). The highest increase in both brachial and central SBP was between sea level and 4619m
190	(+7.0 [-5.0 to 16.0] and +7.0 [-4.5 to 18.0] mmHg respectively) (table 2; figure 2).
191	The AI and ssPPV both increased at HA whereas the reflected wave transit time
192	and systolic ejection period decreased versus sea level (table 2; figure 3). Adjusting the AI
193	to an average heart rate of 75 per minute (AI@75) did not alter the findings.
194	There were significant correlations between central SBP and both AI (r=0.50; 0.41
195	to 0.58: p<0.0001) and age (r=0.32; 21to 0.41: p<0.001). Other independent, albeit weak
196	predictors, of central SBP were SpO ₂ (r=-0.14 -0.25 to -0.05: p=0.02), heart rate (r=-0.16;
197	-0.27 to -0.05: p=.003) male sex (r =0.15; 046 to 0.26: p=0.004) ethnicity (r=0.15; 0.04 to
198	0.25: p=0.007) smoking status (r=0.18; -0.28 to -0.07; p=0.001) and altitude (r=0.10;
199	p=0.05). AI positively correlated with age (r=0.39; p<0.001) and inversely with subject
200	height (r-0.22; p<0.0001) weight (r-0.19; p=0.006), and heart rate (-0.49: p<0.0001). There
201	was no relationship between LLS and either AI or central BP.
202	Multivariate analysis was performed to assess the independent predictors of central

203 systolic BP. Only the univariate predictors were included in the model. The independent

predictors of central SBP were male sex (coefficient, t 4.7; P<0.0001), age (t 3.6; p=0.004) and AI (t 7.5; p<0.0001; overall r^2 =0.40; p<0.0001). If AI was removed from the model (overall r^2 =0.29; p<0.0001) then the independent predictors of central systolic BP were age, heart rate and smoking history. Subject height (coefficient 2.4; p=0.02), age (7.4; p<0.0001) and heart rate (11.4; P<0.0001) were the only independent predictors of AI (overall r^2 =0.43; p<0.0001). The order of the trekking groups did not influence the findings when included in the multivariate analysis.

211

212 **Discussion**

To the author's knowledge, this is the first study to assess the effects of stepwise increasing terrestrial HA on arterial stiffness and central BP over a conventional and progressive HA trek. We found that HA exposure led to a significant increase in central SBP and AI. Neither altitude nor the SpO₂ were independent predictors of AI and central SBP. Heart rate was a significant determinant of both AI and central BP.

HA exposure leads to a wide range of complex effects on both the pulmonary and 218 systolic circulation which have been well described.^{2,4,5,22} Hypobaric hypoxia leads to 219 widespread sympathetic activation leading to an increase in resting heart rate.²³⁻²⁵ The 220 221 reported effects on BP are variable and are highly dependent on the degree of hypoxia and speed and duration of exposure. Furthermore, the type of hypoxic environment may be a 222 major confounder.²⁶ Several previously published studies have used simulated hypoxia 223 (using either a normobaric or hypobaric chamber) in an attempt to replicate the degree of 224 hypoxia observed at genuine HA.^{4,22,25,26} Whilst they are very useful surrogates for HA 225 exposure, u simulated hypoxia does not fully reproduce the environmental and 226 geographical effects genuine terrestrial HA such as the cold or the exercise burden. The 227

reported literature has tended to focus on the effects of HA on brachial artery BP and 228 largely following a relatively short period brief (<6 hours) of simulated hypoxia.^{22,26} 229 230 Available data at terrestrial HA has shown that HA exposure typically leads to an increase in both resting systolic and 24 hour blood pressure which may be more pronounced in 231 those with background hypertension.⁹ The effects of HA on central BP and arterial 232 stiffness have been barely examined at HA, yet they are well recognised to be better 233 predictors of cardiovascular risk than brachial BP.^{10,11} Given the vast numbers of patients 234 with known hypertension and cardiovascular disease who undergo recreational HA 235 exposure annually the ability to better define cardiovascular risk in these individuals would 236 237 be important. This has added importance given that cardiovascular death is a leading cause of non-traumatic death at HA.¹ An improved understanding of the effects of HA on central 238 239 BP and other non-invasive measures of cardiovascular risk such as arterial stiffness might 240 allow for tailored medical therapy at HA to reduce the cardiovascular risk to individuals. 241 We observed a significant increase in brachial but not central pulse pressure suggesting 242 differences in BP behaviour in the peripheral versus the central circulation. Indeed whilst 243 the brachial SBP was higher than that observed centrally the increase in central SBP was greater and was significant across all three altitudes studied (table 2). 244

245 There has only been one previous study to investigate the effects of HA on 246 measures of both arterial stiffness and central BP at terrestrial altitude. Parati et al studied 247 44 subjects who were randomised to placebo or to oral acetazolamide prior to and during HA exposure.⁸ Following sea level assessment the subjects ascended to 4559m within 28 248 249 hours by road to 1130m, then cable car to 3647m before completing the rest of the ascent 250 on foot. Measurements at HA were obtained within 4-6 hours of arrival at 4559m and 251 again after two days at this altitude. They observed a non-significant increase in both 252 central and peripheral SBP but an even greater and significant increase in DBP. AI@75

253	significantly increased from Sea level to HA. However, whereas the SBP continued to
254	increase from 4-6 hours to two days at HA there was no further increase in the AI@75
255	beyond the early increase. In our study we noted a similar sized increase in both brachial
256	and central SBP to that in this previous study and the significance in our current study
257	likely relate to our much larger sample size. Our data would seem to suggest that the
258	increase in heart rate is a significant independent predictor of the increase in AI at HA
259	which was not directly related to either the SpO ₂ or altitude. The observed increase in heart
260	rate, AI, brachial and central SBP would strongly suggest that these increases relate to
261	sustained sympathetic activation at HA as has been well described rather than a genuine
262	increase in large artery stiffness. ²³
263	In one of the only previously published studies to assess the effects of HA on
264	arterial stiffness and brachial BP during a conventional trek Rhodes et al studied 17
265	subjects over an ascent from 80m to 4770m over 11 days. ⁶ They found that HA led to a
266	transient increase in large artery stiffness index (using finger photoplethysmography) noted
267	at day four at 3450 m before returning to baseline levels. A significant rise in both systolic
268	and diastolic BP were observed at 3450m and the increase was sustained throughout the
269	HA exposure. ⁶ Interestingly, they observed that the increase in BP was not related to
270	changes in arterial stiffness nor was there a link between the increase in arterial tone and
271	the presence of AMS. We did not identify a relationship between LLS, SpO2 and either AI,
272	which is an indirect measure of large artery stiffness and central systolic BP at HA.
273	Consistent with previous research we found that the AI related to the
274	subjects age and inversely correlated with height and heart rate. ^{27,28} This is explained by
275	the fact that the time of the reflected wave is related to the dimensions of the body and
276	heart rate. In shorter individuals, a reduced return time for reflected waves leads to an
277	increase in central pressure augmentation. ²⁷ As a result of the noted influence of heart rate

on AI it has been suggested that AI should be adjusted for the effects of heart rate and this 278 has traditionally been to an average of 75 per minute (AI@75).²⁹ Adjusting the AI@75 to 279 account for heart rate did not alter our findings. It has also been more recently suggested 280 that adjusting for heart rate on multivariate analysis of AI is more appropriate and this has 281 been additionally done in our analysis.³⁰ Our data has shown that heart rate was the 282 independent variable with the greatest impact on AI. Indeed augmentation of central BP is 283 284 influenced by heart rate and therefore the duration of systole and shifting the reflected arterial wave to diastole and reducing the time to wave reflection as has been observed in 285 our study.²⁹ Therefore it is reasonable to assume that the increase in AI at HA is largely 286 287 related to the associated increase in heart rate leading to a rise in arterial augmentation and 288 central BP rather than actual changes in large artery stiffness over only 14 days HA 289 exposure.

In this study we were also interested in the effects of HA on the ssPPV. This is a 290 measure of the variation in the pulse pressure averaged over the 10 second arterial 291 waveform recording using the BP⁺ device. The beat to beat variation in pulse pressure is 292 known be influenced by a number of factors including left ventricular preload, stroke 293 volume and ventilation, which are all known to be affected at HA.²² Clinically, probably 294 295 the most widespread use of ssPPV has been to assess fluid responsiveness in mechanically ventilated patients intra-operatively and on intensive care.^{20,21} During inspiration negative 296 297 intrathoracic pressure leads to an increase in venous return and ultimately an increase in ventricular filling. Its effect on left ventricular stroke volume is influenced by hydration 298 and intravascular filling, which is dependent on the relative position on the Frank-Starling 299 curve.¹⁹HA-related hypoxia has been shown to affect both right and left ventricular stroke 300 volume with variable effects on ventricular filling.^{4.22.25} The mechanisms to explain these 301 302 changes are complex and include the known hypoxia mediated pulmonary vasoconstriction

leading to an increase in pulmonary artery systolic pressure and right ventricular afterload.⁵
HA acclimatisation is known to lead to relative dehydration and hypoxia-mediated
hyperventilation all of which may affect biventriuclar ventricular stroke volume. Whilst
the ssPPV cannot be used in isolation serial measurements can be used to assess filling and
fluid responsiveness. In our study the ssPPV was very susceptible to the effects of HA
exposure but was not related to LLS. HA led to a marked increase in the ssPPV, despite no
significant increase in the central arterial pulse pressure.

310 This study has a number of limitations that require acknowledgement. The subjects 311 were studied in groups two days apart. This was done to accommodate the large sample 312 size of the study and ensure excellent reproducibility of the measures and ensure that 313 subject BP measurements were conducted robustly at each individual research station by 314 trained researchers. The environmental factors, such as temperature and barometric 315 pressure would not have been identical for the study groups at the time of their data collection which could have potentially influenced the findings. However, we did not 316 317 observe any significant influence of the trekking group order of study on either AI or 318 central systolic blood pressure. Unfortunately, we did not measure hormonal markers of 319 sympathetic activation, such as circulating catecholamines, to better investigate the 320 mechanism for the increase in SBP and AI, however, we did note that the increases did not 321 relate to the degree of hypoxia (SpO2) or LLS.

322

In conclusion in this study we found that HA exposure led to an increase in brachial and central SBP and a rise in AI compared with near sea level baseline levels. The increase in central SBP and AI was not related to the degree of hypoxia and SpO2 at HA nor to LLS.

326	The observed	changes l	likelv relate	e to increase	d sympathetic	e activation ra	ther than any
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327 genuine change in large artery stiffness.

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330	Acknowledgments
331	The authors would also like to acknowledge and thank the staff in the Department of
332	Cardiology at Poole Hospital for their support. We are extremely grateful to the subjects
333	for their time and for volunteering to take part in this study.
334	
335	Conflict of Interest
336	The authors have no conflict of interest to declare.
337	
338	What is known about the topic?
339	• HA exposure leads to an increase in heart rate and there is evidence from a single study
340	of rapid largely cable car ascent to 4559m that it leads to an increase in central SBP and
341	arterial AI.
342	
343	What this study adds?
344	• This is the first study to examine the effects of stepwise increasing terrestrial HA on
345	arterial stiffness and central BP over a conventional and progressive HA trek to
346	>5000m.
347	• We have discovered that the HA exposure led to a significant increase in central SBP
348	and AI.
349	• Neither altitude nor the SpO ₂ were independent predictors of AI and central SBP.

- The increase in AI related to the increase in heart rate at HA and did not reflect a
- 351 genuine change in large artery stiffness.

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447 Legends for Figures

- **Figure 1**Ascent Profile the altitude and timing of data collection
- **Figure 2** Changes in systolic blood pressure with HA exposure. Symbol* denotes
- 451 significant difference vs baseline level
- **Figure 3** Change in Augmentation Index with high altitude

Table 1: Baseline Demographics

Demographic	Result
Age, years (range)	32.2±8.7 (18-56)
Males n, %	63 (70%)
Height, cm	173.5±9.1
Weight, kg	73.4±12.3
Body mass index kg/m ²	24.38±2.7
Ethnicity, %	
- Caucasian	78 (86.7%)
- Nepalese	11 (12.2%)
- South Asian	1 (1.1%)
Smoking status (N, %)	
- Current	9 (10%)
- Ex	11 (12.2%)
- Never	70 (77.8%)

Table 2 Effect of high altitude on measured vascular parameters including central blood

Parameter	Sea level	3619m	4600m	5140m	P value
Heart rate/ minute	65.2±12.8	69.6±11.8	77.3±15.3	78.2±13.6	<0.0001*†‡
Oxygen Saturations, %	97.7±1.4†	91.9±3.4	82.8±6.3	80.4±5.3	<0.0001*†‡
Lake louis Scores	0.23 (0.64)	1.1 (1.9)	1.4 (1.6)	1.3 (1.4)	<0.0001*†‡
Brachial artery	$132.8{\pm}14.0$	136.9±13.4	138.8±13.3	138.6±13.9	0.04†‡
systolic BP, mmHg					
Brachial artery diastolic	81.8±11.7	84.7±9.4	83.7±9.8	83.9±9.7	0.28
BP, mmHg					
Mean brachial arterial	99.3±12.9	102.0±9.9	102.1±9.9	102.2±9.8	0.23
BP, mmHg					
Brachial artery pulse	51.6±11.3	52.1±9.7	55.5±10.9	54.7±11.3	0.02†
pressure, mmHg					
Central systolic BP,	$124.7{\pm}14.8$	130.1±14.2	131.4±15.4	129.4±14.3	0.02*†‡
mmHg					
Central diastolic BP,	84.0±11.6	87.5±9.6	86.8±9.6	87.3±9.5	0.09
mmHg					
Central artery pulse	40.7±9.5	42.6±9.6	44.6±13.4	42.1±9.9	0.26
pressure, mmHg					
Augmentation	55.3±34.9	71.1±34.1	61.8±36.7	56.6±32.7	0.001†
index, %					
Reflected wave transit	0.16±0.02	0.16±0.02	0.14 ± 0.02	0.14 ± 0.01	<0.000*†‡
time, s					
Systolic ejection period, s	0.30±0.03	0.31±0.02	0.29±0.03	0.28±0.02	<0.0001†‡
Supra Systolic pulse	0.23±0.13	0.28±0.15	0.37±0.20	0.34±0.19	<0.0001*†‡
pressure variation					

pressure and augmentation index

BP, blood pressure; results of post hoc tests vs baseline sea level, *3880m, † 4400m, ‡ 5140m



3 4 5 6 7 8 9 10 11 12 13 14 Time in Days



Altitude

Brachial

5140m

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		Sea level

Altitude

3619m 4600m





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5140m

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