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| 14. ABSTRACT For the last 12 months we have been working on data analysis and manuscript preparation. The first 6 papers from this study have been published, and another 6 are in preparation. The overview physiology paper was accepted last week at PLOS ONE, and the others have been accepted in major physiology journals. The next batch of papers will be from the OMICS portion of the study. We hired a new bioinformatics postdoc in October, and he has hit the ground running with comprehensive analyses of the OMICS dataset. The gene expression paper will be complete in another 3-4 weeks, and shortly after that the metabolomics and epigenetics data will be ready for publication. By the end of 2014 all major papers from the study will have been published. So far the project is from our perspective a complete success with identification of new physiological aspects of acclimatization, and first-ever insights into the underlying OMICS mechanisms. We are now looking for one additional year of funding to explore new analyses and integration that is possible because of the high quality of this dataset, and that could lead to even more comprehensive, "big picture" views of the process of human acclimatization to hypoxia at high altitude. | | | | | |
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INTRODUCTION:

The goal of this project is to advance high-altitude medical research by discovering the basic molecular mechanisms of acclimatization and de-acclimatization that protect soldiers from high-altitude illness.

BODY:

All major milestones have been accomplished. Now we are working on papers integrating the findings from the extensive physiological studies and the OMICS studies. Since no one has done that work before, we are inventing the methods and approaches as we go along. A major breakthrough has been the application of an advanced clustering algorithm called WGCNA to our datasets. This will allow us to condense the enormous datasets generated by the gene expression and epigenetics chip studies into a manageable system that can easily be tested for relationships to physiological tests.

Accomplishments to date:

- IRB compliance and continuing review have been completed
- Analyses are completed for all subjects at all time points for epigenetics, gene expression, microRNA and metabolomics.
- All cytokine arrays are done, with follow-up and validation ELISAs completed. Writing of those manuscripts is underway.
- Six papers have been accepted for publication, three in Journal of Applied Physiology, one in Experimental Physiology, one in Acta Scandinavica and the overview paper at PLOS ONE
- A seventh paper is under review at NeuroReports.
- Nitric oxide analyses are done, adenosine and hydrogen sulfide analyses are done. Work has begun on a paper on NO and H₂S with Drs. Roach, Kevil and Gladwin.
- Analysis of ADP, ATP and purigenic receptors is complete, writing that manuscript is underway. Another paper is in the works as well with Drs. Eltzschig, Blackburn, Xia, and Davis on adenosine in AltitudeOmics.
- The Lovering laboratory, home of our collaborators on AltitudeOmics, have two papers in preparation on AMS and intrapulmonary shunts, and one on gas exchange during AltitudeOmics.

KEY RESEARCH ACCOMPLISHMENTS:

1. Completed the first ever measurements of acute mountain sickness, cognitive function and exercise capacity after 7 and 21 days of de-acclimatization. The results suggest near complete retention of acclimatization after 7 days de-acclimatization, and about 70% retention after 21 days. This key finding will be used in the OMICS analyses to help identify factors that occur with acclimatization, and are still present after de-acclimatization.
2. Six research papers have been completed and published on the physiology of human acclimatization to high altitude, and another is under review. Seven additional primary papers will be completed this year. Please see Appendices section for a table showing the “Status of Research Papers” and for a PDF of the published papers.

REPORTABLE OUTCOMES:

1. Completed all regulatory steps to gain approval for this multi-site, multi-nation study.
2. Safely completed data collection on 23 young healthy student volunteers, and safely transported and cared for them and 40 scientists to/from Bolivia.
3. We are 100% in analysis and manuscript writing mode regarding all aspects of the study.

CONCLUSION:

Humans retain acclimatization after 7 and 21 days of de-acclimatization. This was a key hypothesis of the study. Yet to be determined is what are the OMICS responses that can be linked to the process of gaining acclimatization, and its retention on descent to low altitude?

Status of manuscripts from the AltitudeOmics study, 2/15/2014

| # | Title | First/Last Authors | Status | PMID | Link to PDF |
|---|--------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|--------------------|-----------------------------|----------|-------------|
| 1 | Amann M, Goodall S, Twomey R, Subudhi AW, Lovering AT, Roach RC. AltitudeOmics: on the consequences of high-altitude acclimatization on the development of fatigue during locomotor exercise in humans. <i>Journal of applied physiology</i> . 2013;115:634-42. | Amann/Roach | Published, JAPPL | 23813531 | |
| 2 | Goodall S, Twomey R, Amann M, Ross EZ, Lovering AT, Romer LM, Subudhi AW, Roach RC. AltitudeOmics: Exercise-induced supraspinal fatigue is attenuated in healthy humans after acclimatisation to high altitude. <i>Acta Physiologica</i> . 2014. | Goodall/Roach | Accepted, Acta Scandinavica | 24450855 | |
| 3 | Subudhi AW, Fan JL, Evero O, Bourdillon N, Kayser B, Julian CG, Lovering AT, Roach RC. AltitudeOmics: Effect of ascent and acclimatization to 5260 m on regional cerebral oxygen delivery. <i>Experimental Physiology</i> . 2013. | Subudhi/Roach | Accepted, Exp Physiol | 24243839 | |
| 4 | Fan JL, Subudhi AW, Evero O, Bourdillon N, Kayser B, Lovering AT, Roach RC. AltitudeOmics: Enhanced cerebrovascular reactivity and ventilatory response to CO2 with high altitude acclimatisation and re-exposure. <i>Journal of Applied Physiology</i> . 2013. | Fan/Roach | Accepted, JAPPL | 24356520 | |
| 5 | Subudhi AW, Fan JL, Evero O, Bourdillon N, Kayser B, Julian CG, Lovering AT, Panerai RB, Roach RC. AltitudeOmics: Cerebral autoregulation during ascent, acclimatization, and re-exposure to high altitude and its relation with acute mountain sickness. <i>Journal of Applied Physiology</i> . 2013. | Subudhi/Roach | Accepted, JAPPL | 24371013 | |
| 6 | Subudhi AW, Bucher J, Bourdillon N, Davis C, Elliott J, Eutermoster M, Evero O, Fan JL, Jameson-Van Houten S, Julian CG, Kark J, Kark S, Kayser B, Kern JP, Kim SE, Lathan C, Laurie SS, Lovering AT, Paterson R, Polaner D, Ryan BJ, Spira J, Tsao JW, Wachsmuth NB, Roach RC. AltitudeOmics: The Integrative Physiology of the Onset and Retention of Acclimatization to Hypoxia in Humans. <i>PLOS One</i> (In Press) 2014. | Subudhi/Roach | Accepted, PLOSOne | | |

| # | Title | First/Last Authors | Status | PMID | Link to PDF |
|----|----------------------------------------------------------------------------------------------------------------------------------|--------------------|------------------------|------|-------------|
| 7 | AltitudeOmics: hemoglobin mass increases within 7 days of acclimatization to 5260m and is lost within 7 days of descent to 1525m | Ryan/Roach | Under revision | | |
| 8 | AltitudeOmics: Detecting high altitude cognitive impairment with DANA, an Android-based neurocognitive assessment tool | Roach/Roach | Submitted, Neuroreport | | |
| 9 | AltitudeOmics: Gene expression and epigenetics during ascent, acclimatization, and re-exposure to 5,260m | Julian/Roach | In prep | | |
| 10 | AltitudeOmics: Effect of ascent, acclimatization, and re-exposure to 5,260m on pulmonary shunt | Elliot/Roach | In prep | | |
| 11 | AltitudeOmics: Effect of ascent, acclimatization, and re-exposure to 5,260m on muscle mitochondrial respiration | Chicco/Roach | In prep | | |
| 12 | AltitudeOmics: Metabolomics during ascent, acclimatization, and re-exposure to 5,260m | Monte/Roach | In prep | | |
| 13 | AltitudeOmics: Effect of ascent, acclimatization, and re-exposure to 5,260m on AMS and pulmonary shunt | Kern/Roach | In prep | | |
| 14 | AltitudeOmics: Gene Expression and Acute Mountain Sickness | ?/Roach | In prep | | |
| 15 | AltitudeOmics: MicroRNA expression during ascent, acclimatization, and re-exposure to 5,260m | Kern/Roach | In prep | | |

AltitudeOmics: on the consequences of high-altitude acclimatization for the development of fatigue during locomotor exercise in humans

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¹Department of Medicine, University of Utah, Salt Lake City, Utah; ²Faculty of Health and Life Sciences, Northumbria University, Newcastle, United Kingdom; ³School of Sport and Service Management, University of Brighton, Eastbourne, United Kingdom; ⁴Altitude Research Center, Department of Emergency Medicine, University of Colorado Anschutz Medical Campus, Aurora, Colorado; ⁵Department of Biology, University of Colorado, Colorado Springs, Colorado; and ⁶Department of Human Physiology, University of Oregon, Eugene, Oregon

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Amann M, Goodall S, Twomey R, Subudhi AW, Lovering AT, Roach RC. AltitudeOmics: on the consequences of high-altitude acclimatization for the development of fatigue during locomotor exercise in humans. *J Appl Physiol* 115: 634–642, 2013. First published June 27, 2013; doi:10.1152/jappphysiol.00606.2013.—The development of muscle fatigue is oxygen (O₂)-delivery sensitive [arterial O₂ content (C_aO₂) × limb blood flow (Q_L)]. Locomotor exercise in acute hypoxia (AH) is, compared with sea level (SL), associated with reduced C_aO₂ and exaggerated inspiratory muscle work (W_{insp}), which impairs Q_L, both of which exacerbate fatigue individually by compromising O₂ delivery. Since chronic hypoxia (CH) normalizes C_aO₂ but exacerbates W_{insp}, we investigated the consequences of a 14-day exposure to high altitude on exercise-induced locomotor muscle fatigue. Eight subjects performed the identical constant-load cycling exercise (138 ± 14 W; 11 ± 1 min) at SL (partial pressure of inspired O₂, 147.1 ± 0.5 Torr), in AH (73.8 ± 0.2 Torr), and in CH (75.7 ± 0.1 Torr). Peripheral fatigue was expressed as pre- to postexercise percent reduction in electrically evoked potentiated quadriceps twitch force (ΔQ_{tw,pot}). Central fatigue was expressed as the exercise-induced percent decrease in voluntary muscle activation (ΔVA). Resting C_aO₂ at SL and CH was similar, but C_aO₂ in AH was lower compared with SL and CH (17.3 ± 0.5, 19.3 ± 0.7, 20.3 ± 1.3 ml O₂/dl, respectively). W_{insp} during exercise increased with acclimatization (SL: 387 ± 36, AH: 503 ± 53, CH: 608 ± 67 cmH₂O·s⁻¹·min⁻¹; P < 0.01). Exercise at SL did not induce central or peripheral fatigue. ΔQ_{tw,pot} was significant but similar in AH and CH (21 ± 2% and 19 ± 3%; P = 0.24). ΔVA was significant in both hypoxic conditions but smaller in CH vs. AH (4 ± 1% vs. 8 ± 2%; P < 0.05). In conclusion, acclimatization to severe altitude does not attenuate the substantial impact of hypoxia on the development of peripheral fatigue. In contrast, acclimatization attenuates, but does not eliminate, the exacerbation of central fatigue associated with exercise in severe AH.

altitude; respiratory muscle work; arterial O₂ content; cerebral blood flow

THE DEVELOPMENT OF LOCOMOTOR muscle fatigue during whole-body endurance exercise is highly sensitive to the delivery of oxygen [O₂; arterial O₂ content (C_aO₂) × leg blood flow (Q_L)]. Specifically, blunted O₂ delivery exaggerates, and augmented O₂ delivery attenuates the rate of development of locomotor muscle fatigue during exercise (1).

Acute exposure to hypoxia (AH) has a substantial impact on the two determinants of leg muscle O₂ delivery during strenuous locomotor exercise. First, despite a marked hyperventilatory response, arterial partial pressure of O₂ [PO₂ (P_aO₂)] and arterial hemoglobin saturation (S_aO₂) fall below sea level (SL) values and cause a significant reduction in C_aO₂. In addition, inspiratory muscle work (W_{insp}) is increased substantially at any given workload in hypoxia (2, 58), and these high levels of W_{insp} compromise, in a dose-dependent manner, Q_L during exercise (34). Each of these two determinants of leg muscle O₂ delivery, namely C_aO₂ and Q_L, accounts for, substantially and independently, the accelerated development of locomotor muscle fatigue in hypoxia (2).

During prolonged exposure to altitude, a progressive, time-dependent hyperventilation, which increases alveolar PO₂, occurs over the initial hours and days and advances more gradually over the ensuing 1–2 wk of acclimatization (56). This ventilatory acclimatization adds to an accompanying reduction in the alveolar-arterial O₂ gradient, which combined, substantially improves arterial oxygenation during exercise by increasing P_aO₂ and S_aO₂ (9, 13). Furthermore, chronic exposure to hypoxia (CH) is accompanied by erythropoiesis, and the combination of an increased hemoglobin concentration ([Hb]) plus improved oxygenation may serve to restore resting SL C_aO₂ (8, 13). In contrast to this beneficial effect on O₂ delivery, Q_L, during intense leg exercise at a given submaximal absolute workload, has been suggested to decline from SL to CH (8, 49, 64). The net effect of these acclimatization-induced, opposing consequences on leg O₂ delivery depends on the degree to which the increase in C_aO₂ can counterbalance potential reductions in Q_L. It has been documented previously that at a given absolute workload, locomotor muscle O₂ delivery is reduced from SL to AH with no further changes following acclimatization (Pikes Peak, 4,300 m) (8, 64). Therefore, given the critical role of muscle O₂ delivery in the development of fatigue, it could be argued that peripheral fatigue during constant-load endurance exercise is exacerbated in AH (vs. SL) and does not improve further during prolonged acclimatization. On the other hand, studies conducted at the same location as the present experiments [Mt. Chacaltaya (Bolivia), 5,260 m] document a reduction in locomotor muscle O₂ delivery from SL to AH and a full recovery following prolonged exposure, with the net effect of similar values in SL and CH (13). Based on these findings, it could be argued that the development of peripheral fatigue during constant-load endurance exercise is fastened in AH but recovers to SL values in CH.

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In this study, we sought to quantify exercise-induced locomotor muscle fatigue induced by the identical constant-load cycling trial performed at SL, in AH, and in CH (following 14 days at 5,260 m) to clarify the effects of acclimatization. We hypothesized that fatigue is, compared with SL, exacerbated significantly in AH and that altitude acclimatization would alleviate this impact.

METHODS

This study was conducted as part of the AltitudeOmics project, examining the integrative physiology of human responses to hypoxia. All procedures conformed to the Declaration of Helsinki and were approved by the Universities of Colorado, Oregon, and Utah Institutional Review Boards and the U.S. Department of Defense Human Research Protection Program Office. All subjects were born and raised below 1,500 m and had not traveled to elevations >1,000 m for 3 mo before the experiments. Eight subjects (age 21 ± 1 y, body weight 69 ± 11 kg, height 176 ± 10 cm) were studied at SL and following 14 days of altitude acclimatization at 5,260 m on Mt. Chacaltaya. At high altitude, subjects did not follow a systematic exercise-training program but were given the opportunity to participate, on a voluntary basis, in light hikes around the campsite (no significant change in altitude).

Experimental Protocol

All participants were familiarized thoroughly with various experimental procedures involved in this investigation. The SL experiments of the present study were conducted ~130 m above SL [Eugene, OR; barometric pressure (BP) 750.0 ± 2.2 Torr]. The experiments in AH were conducted at the same altitude, while breathing a gas mixture containing 10.5% O₂ balance nitrogen, and experiments in CH were conducted on the 14th day of acclimatization at 5,260 m (BP 408.9 ± 0.7 Torr). Two participants were tested every morning. To assure that all subjects were tested exactly on *day 14* after arrival on the mountain, the groups' transport to the mountain was staged, i.e., two new participants arrived every day. SL peak power output (W_{peak}) was obtained from a maximal incremental exercise test (70, 100, 130, and 160 W for 3 min, each followed by 15 W/min increases thereafter) on a computer-controlled bicycle ergometer (Velotron, Dynafit; RacerMate, Seattle, WA). The experimental trial consisted of the identical constant-load cycling exercise (same absolute workload and duration) in each condition. Preliminary experiments (using different subjects), conducted to identify a workload that causes voluntary exhaustion between 8 and 12 min when acutely exposed to 5,260 m, revealed that a constant workload equal to 50% of SL W_{peak} was required to reach this goal. Based on this, the workload during the experimental trials was set to equal 50% (138 ± 14 W) of the subjects' SL W_{peak} (275 ± 14 W). Since an individual's endurance/aerobic capacity is lowest in AH (vs. SL and CH) (13), the first trial was performed to voluntary exhaustion in AH, and the achieved time (10.6 ± 0.7 min) was then used for all subsequent trials. A 5-min warm-up at 10% W_{peak} (27 ± 8 W) preceded each trial. Throughout exercise, subjects were instructed to maintain their preferred pedal frequency, as determined during the practice sessions (88 ± 3 rpm). Neuromuscular function was assessed before and within 2.5 min after exercise. During these procedures, subjects breathed ambient air at SL and in CH and a gas mixture (10.5% O₂) in AH.

Exercise Responses

Pulmonary ventilation (V_E) and gas exchange were measured at rest and throughout exercise using an open circuit system (Ultima PFX; Medical Graphics, St. Paul, MN, and O2cap; Oxigraf, Mountain View, CA). Arterial O₂ saturation (S_pO_2) was estimated continuously at rest and during exercise using a pulse oximeter (Nellcor N-200;

Pleasanton, CA) with adhesive forehead sensors. A correction factor based on arterial blood gases was used to adjust for the nonlinearity associated with the obtained pulse oximeter values (error between 60% and 80% saturation: 6%; error between >90% saturation: 3%). Heart rate was measured from the R-R interval of an ECG, using a three-lead arrangement. Ratings of perceived exertion were obtained using Borg's modified CR10 scale (10). [Hb] was measured (Radiometer OSM-3) in resting arterial blood samples collected at SL and on the 16th day at 5,260 m. C_aO_2 was estimated as $1.39 [\text{Hb}] \times (S_pO_2/100)$. During all constant workload trials, esophageal pressure (P_{es}) was measured via a nasopharyngeal balloon (Cooper Surgical, Trumbull, CT), using standard procedures (7). To estimate W_{insp} , P_{es} was integrated over the period of inspiratory flow, and the results were multiplied by respiratory frequency (f_R) and labeled the inspiratory muscle pressure-time product. Vastus lateralis oxygenation was assessed using a multichannel near-infrared spectroscopy (NIRS) instrument (Oxymon Mk III; Artinis, Zetten, The Netherlands). As described previously (5), a NIR emitter and detector pair was affixed over the belly of the left vastus lateralis muscle (~15 cm proximal and 5 cm lateral to the midline of the superior border of the patella), using a spacer with an optode distance of 5.0 cm. Probes were secured to the skin using double-sided tape and shielded from light using elastic bandages. The Beer-Lambert Law was used to calculate micrometer changes in tissue oxygenation [oxyhemoglobin (O₂Hb) and deoxyhemoglobin (HHb)] across time, using received optical densities from two continuous wavelengths of NIR light (780 and 850 nm) and a fixed differential path-length factor of 4.95 (26). Total hemoglobin (THb) was calculated as the sum of [O₂Hb] and [HHb] changes to give an index of change in regional blood volume (59). Data were recorded continuously at 10 Hz and expressed relative to the resting baseline recorded in each experimental condition. Mean cerebral blood flow (CBF) was estimated from blood velocity (CBFv) in the left middle cerebral artery (MCA; 50 ± 4 mm deep), determined using a 2-MHz transcranial Doppler (Spencer Technologies, Seattle, WA). An index of cerebral O₂ delivery was calculated as the product of CBFv and C_aO_2 . Changes in CBFv were assumed to reflect changes in CBF, based on evidence that the MCA changes minimally in response to hypoxia and hypocapnia (47, 54). The validity of this assumption at altitude has been challenged recently (62). Evidence of MCA dilation was demonstrated in subjects at altitudes above 6,400 m, but no changes in MCA diameter were observed at altitudes comparable with the present study (<5,300 m) (63). We acknowledge that these measurements must be interpreted with caution until definitive studies of MCA diameter at altitude are conducted.

Expiratory Flow Limitations and Lung Volume Responses

Expiratory flow limitations. Subjects performed three maximal volitional flow-volume (FV) maneuvers before and after exercise (after assessment of neuromuscular function). Exercise tidal FV loops (FVLs) were plotted within the best of the six maximal loops (MFVLs), based on measured inspiratory capacity (IC) maneuvers (rest, 3 min of exercise, and immediately before the termination of exercise). Acceptable IC maneuvers during exercise required that peak inspiratory P_{es} match that obtained at rest. The amount of expiratory flow limitation was defined as the percentage of the tidal volume (V_T) that met the boundary of the expiratory portion of the MFVL (38).

Lung volumes. Functional residual capacity (FRC) was measured in a body plethysmograph (Platinum Elite Series; Medical Graphics), and total lung capacity (TLC) was calculated as the sum of FRC and IC. End-expiratory lung volume (EELV) was determined by subtracting the maximal IC, as measured during exercise from TLC, as measured at rest. End-inspiratory lung volume (EILV) was calculated as the sum of EELV and V_T . Inspiratory reserve volume, during exercise, was calculated by subtracting EILV from TLC, and expiratory reserve volume, during exercise, was determined by subtracting the residual volume from EELV.

Force and Compound Muscle Action Potentials

Knee-extensor force during voluntary and evoked contractions was measured using a calibrated load cell (TedeA, Basingstoke, UK). The load cell was fixed to a custom-built chair and connected to a noncompliant cuff, attached around the participant's right leg, just superior to the ankle malleoli. Participants sat upright in the chair with the hips and knees at 90° of flexion. Compound muscle action potentials (M-waves) were recorded from surface electrodes placed 2 cm apart over the vastus lateralis muscle belly. A reference electrode was placed over the patella. Evoked signals were amplified [gain: 1,000; force: custom-built bridge amplifier; electromyographic (EMG): PowerLab 26T; ADInstruments (Oxfordshire, UK)], band-pass filtered (EMG only: 20–2,000 Hz), digitized (4 kHz; PowerLab 26T, ADInstruments), acquired, and later analyzed (LabChart v7.0; ADInstruments) for peak-to-peak amplitude.

Neuromuscular Function

Force and EMG variables were assessed before and immediately (<2.5 min) after each trial. Before each trial, maximum voluntary contraction (MVC) force was determined from three control contractions. Femoral nerve stimulation was delivered during each 5-s MVC, and an additional stimulus was delivered after the MVC to determine the potentiated quadriceps twitch force ($Q_{tw,pot}$) and voluntary muscle activation (VA) (42). Briefly, the force produced during the superimposed twitch (SIT), delivered within 0.5 s of attaining peak force during the MVC, was to be compared with the force produced by the single twitch, delivered during relaxation, ~2 s after the MVC: $VA (\%) = [1 - (SIT/Q_{tw,pot})] \times 100$. The contraction sets were repeated three times, with 30 s between each set. Visual feedback of the target force was provided via a computer monitor.

Femoral nerve stimulation. Single electrical stimuli (200 μ s pulse width) were delivered to the right femoral nerve via surface electrodes (32 mm diameter; CF3200; Nidd Valley Medical, North Yorkshire, UK) and a constant-current stimulator (DS7AH; Digitimer, Welwyn Garden City, Hertfordshire, UK). The cathode was positioned over the nerve, high in the femoral triangle; the anode was placed midway between the greater trochanter and the iliac crest (32). The site of stimulation that produced the largest resting twitch amplitude and M-wave was located. Single stimuli were delivered, beginning at 100 mA and increasing by 20 mA, until plateaus occurred in twitch amplitude and M-wave. Supramaximal stimulation was ensured by increasing the final intensity by 30% (mean current, 250 \pm 55 mA). Muscle contractility was assessed for each potentiated twitch as twitch amplitude ($Q_{tw,pot}$: peak force – onset force), maximum rate of force development (MRFD), contraction time, maximum relaxation rate (MRR), and one-half relaxation time ($RT_{0.5}$). Sarcolemmal membrane excitability was inferred from the peak-to-peak amplitude of the electrically evoked M-wave (27).

Reliability Measures

On a separate day, measures of neuromuscular function were repeated twice in all subjects at SL. The two assessment procedures were separated by a 2-min walk around the laboratory, followed by a 5-min rest period. Coefficient of variation (CV) and Pearson product-moment correlation coefficients (r) were calculated to evaluate test-retest error (precision) and test-retest reliability of the neuromuscular function-assessment procedure. All correlations were significant and indicated; in combination with the CVs, acceptable degrees of reproducibility include: MVC, CV = 3.1%, $r = 0.97$; $Q_{tw,pot}$, CV = 4.1%, $r = 0.98$; M-wave peak, CV = 4.8%, $r = 0.98$; VA, CV = 3.3%, $r = 0.77$.

Statistical Analysis

A one-way repeated-measures ANOVA was performed to evaluate differences among trials. A least-significance difference test identified

the means that were significantly different with $P < 0.05$. Results are expressed as mean \pm SE.

RESULTS

C_aO_2 and Cerebral O_2 Delivery

C_aO_2 at rest was significantly lower in AH compared with SL and CH (17.3 \pm 0.5, 19.3 \pm 0.7, 20.3 \pm 1.3 ml O_2 /dl, respectively). Acclimatization to altitude significantly increased [Hb] and S_pO_2 , resulting in similar C_aO_2 at SL and in CH ($P = 0.16$). Resting CBFv was similar among SL, AH, and CH (50.5 \pm 3.7, 52.7 \pm 2.3, and 55.7 \pm 3.0 cm/s, respectively; $P = 0.45$). In all three conditions, CBFv increased significantly from rest to the final minute of exercise (22 \pm 3%, 39 \pm 6%, and 28 \pm 5% for SL, AH, and CH, respectively; Table 1). The percent increase was significantly greater in AH compared with that observed at SL and in CH. The cerebral O_2 delivery index during the last minute of exercise was 18 \pm 5% lower in AH vs. SL (Table 1) and 17 \pm 8% greater in CH vs. SL (Table 1).

Ventilatory Effects

Ventilatory response. AH increased W_{insp} work by 34 \pm 8% above that at SL ($P < 0.01$) and dropped S_pO_2 by 36 \pm 3% during the final minute of exercise. Following 14 days of acclimatization, W_{insp} was increased further by 23 \pm 8% from AH, and S_pO_2 , during the final minute of exercise, was 36 \pm 5% higher in CH vs. AH. Breathing frequency and V_E rose

Table 1. Mean responses to the final minute of exercise (138 \pm 14 W, 10.6 \pm 0.7 min)

| | Sea Level | Acute Hypoxia | Chronic Hypoxia |
|-----------------------------------------------------------------------|-----------------|------------------|-------------------|
| HR, beats/min | 152 \pm 5 | 174 \pm 4* | 166 \pm 4*† |
| V_E , l min ⁻¹ | 64 \pm 4 | 113 \pm 8* | 133 \pm 10*† |
| f_R , breaths min ⁻¹ | 32 \pm 2 | 50 \pm 3* | 54 \pm 3* |
| V_T , liter | 2.0 \pm 0.1 | 2.2 \pm 0.2 | 2.6 \pm 0.2*† |
| $\dot{V}O_2$, l min ⁻¹ | 2.58 \pm 0.19 | 2.44 \pm 0.19* | 2.39 \pm 0.16*† |
| $\dot{V}CO_2$, l min ⁻¹ | 2.51 \pm 0.22 | 2.81 \pm 0.21* | 2.40 \pm 0.15*† |
| $V_E/\dot{V}O_2$ | 25 \pm 1 | 50 \pm 4* | 56 \pm 3*† |
| $V_E/\dot{V}CO_2$ | 26 \pm 1 | 41 \pm 2* | 58 \pm 3*† |
| S_pO_2 , % | 94.1 \pm 1.0 | 62.2 \pm 1.8* | 75.6 \pm 1.2*† |
| CBFv, cm/s | 59.1 \pm 4.8 | 74.2 \pm 3.8* | 73.2 \pm 3.4* |
| Cerebral O_2 delivery, a.u. | 1,105 \pm 62 | 895 \pm 40* | 1,289 \pm 42*† |
| T_i/T_{tot} | 0.35 \pm 0.01 | 0.39 \pm 0.01* | 0.39 \pm 0.01* |
| T_e , s | 1.30 \pm 0.08 | 0.74 \pm 0.05* | 0.70 \pm 0.04* |
| W_{insp} , cmH ₂ O · s ⁻¹ · min ⁻¹ | 387 \pm 36 | 503 \pm 53* | 608 \pm 67*† |
| IC, liter | 3.29 \pm 0.22 | 3.13 \pm 0.23 | 3.60 \pm 0.23* |
| V_T/IC | 0.60 \pm 0.03 | 0.68 \pm 0.02* | 0.72 \pm 0.02*† |
| IRV, liter | 1.30 \pm 0.14 | 0.99 \pm 0.13* | 0.96 \pm 0.05* |
| ERV, liter | 1.98 \pm 0.25 | 2.14 \pm 0.29 | 1.67 \pm 0.25*† |
| EILV, %TLC | 80.5 \pm 1.6 | 85.4 \pm 1.7* | 85.2 \pm 0.9* |
| EELV, %TLC | 51.5 \pm 1.8 | 53.8 \pm 2.5 | 46.9 \pm 2.1*† |
| Expiratory flow limitation, n out of 8 subjects | 0/8 | 2/8 | 4/8 |
| RPE | 12.3 \pm 1.0 | 19.8 \pm 0.1* | 17.9 \pm 0.6*† |
| Dyspnea | 11.5 \pm 0.7 | 19.5 \pm 0.2* | 19.3 \pm 0.2* |

HR, heart rate; V_E , minute ventilation; f_R , breathing frequency; V_T , tidal volume; $\dot{V}O_2$, maximum oxygen (O_2) uptake; $\dot{V}CO_2$, carbon dioxide production; S_pO_2 , arterial O_2 saturation; CBFv, cerebral blood flow velocity; T_i , duration of inspiration; T_{tot} , duration of entire breath; T_e , duration of expiration; W_{insp} , inspiratory muscle work; IC, inspiratory capacity; IRV, inspiratory reserve volume; ERV, expiratory reserve volume; EILV, end-inspiratory lung volume; TLC, total lung capacity; EELV, end-expiratory lung volume; RPE, rating of perceived exertion. * $P < 0.05$ vs. sea level; † $P < 0.05$ vs. acute hypoxia, $n = 8$.

substantially over the time of exercise in AH and CH, and V_E was, during the final minute, $79 \pm 13\%$ and $110 \pm 12\%$, respectively, higher compared with SL ($P < 0.01$). Pulmonary V_E during the final minute of exercise was $19 \pm 4\%$ higher in CH vs. AH ($P < 0.01$). Compared with SL, O_2 uptake, during the final minute of exercise, was $5 \pm 2\%$ and $7 \pm 2\%$ lower in AH and CH, respectively (both $P < 0.05$; Fig. 1).

Expiratory flow limitation. At SL, exercise flow rates during tidal breathing were well within the MFVL in all eight subjects. At end-exercise in AH, 6–51% of the V_T in two of the eight subjects reached flow limitation, as lung volume approached end-expiration. As V_E increased further in CH, expiratory flow rate became more limited, and 10–64% of the V_T in four of the eight subjects met the limit imposed by the MFVL.

Membrane Excitability and Contractile Function

M-waves. As a measure of membrane excitability we examined pre- vs. postexercise vastus lateralis M-wave amplitudes in conjunction with the quadriceps muscle mechanical properties. Pre-exercise M-wave amplitudes were similar in all three conditions (10.2 ± 1.0 mV, 9.4 ± 0.7 mV, and 12.9 ± 1.8 mV for SL, AH, and CH, respectively; $P = 0.15$). Postexercise M-wave amplitudes were unchanged from pre-exercise baseline values at SL and in AH (10.2 ± 1.0 mV and 9.6 ± 0.9 mV, respectively; $P > 0.3$). However, following exercise in CH, M-wave amplitudes (7.8 ± 2.1 mV) were reduced significantly from pre-exercise baseline levels (range: 1–18%; $P < 0.01$).

Quadriceps twitch force. Pre-exercise $Q_{tw,pot}$ was similar in all three conditions (106 ± 4 N, 109 ± 4 N, and 110 ± 5 N for SL, AH, and CH, respectively; $P = 0.18$). Exercise in both hypoxic conditions caused a substantial ($P < 0.01$) but similar ($P = 0.14$) reduction in $Q_{tw,pot}$ in all eight subjects. In contrast, exercise at SL did not induce measurable locomotor muscle fatigue; the postexercise $Q_{tw,pot}$ was similar to pre-exercise baseline.

MVC force. Pre-exercise MVC was similar in all three conditions (391 ± 30 N, 394 ± 25 N, and 372 ± 30 N for SL, AH, and CH, respectively; $P = 0.21$). At SL, postexercise MVC was similar to pre-exercise baseline ($P = 0.42$). In

contrast, exercise in AH and CH caused a substantial reduction in MVC in all eight subjects. However, the exercise-induced reduction in MVC was $30 \pm 9\%$ less in CH vs. AH ($P < 0.05$).

Muscle activation. Pre-exercise baseline values were similar in all three conditions ($94 \pm 1\%$, $94 \pm 1\%$, and $93 \pm 1\%$ for SL, AH, and CH, respectively; $P = 0.19$). Following the exercise at SL, muscle activation was unchanged from pre-exercise baseline ($P = 0.88$). In both AH and CH, postexercise muscle activation was significantly lower compared with pre-exercise baseline values. However, the pre- to postexercise decrease in muscle activation was $52 \pm 12\%$ less in CH vs. AH ($P < 0.01$).

Within-twitch measurements. MRFD, MRR, and $RT_{0.5}$ complement the findings reported for $Q_{tw,pot}$. The pre- to postexercise changes in within-twitch measurements of MRFD, MRR, and $RT_{0.5}$ were similar in CH vs. AH.

Vastus Lateralis Tissue Oxygenation

O_2Hb was unchanged from baseline to warm-up at SL ($P = 0.40$) but decreased in AH ($P < 0.05$) and CH ($P = 0.05$). Compared with baseline, O_2Hb was unchanged during the final minute of exercise at SL ($P = 0.73$) but was significantly lower in AH and CH (both $P < 0.01$). This decrease was significantly greater in AH vs. CH. HHb was unchanged from baseline to warm-up at SL ($P = 0.80$) but decreased significantly in AH and CH. Compared with baseline, HHb was unchanged during the final minute of exercise at SL ($P = 0.24$) but similarly increased in AH and CH (both $P < 0.01$). THb was unchanged from baseline to warm-up in all three conditions. In contrast, compared with baseline, THb was increased significantly and similarly ($P = 0.37$) during the final minute of exercise in all three conditions.

DISCUSSION

The purpose of this investigation was to evaluate the effect of altitude acclimatization on the development of fatigue during whole-body endurance exercise. Subjects repeated the identical constant-load cycling exercise at SL, in AH, and in CH. No measurable degree of fatigue was found following the exercise at SL. However, the identical exercise in AH, characterized by a reduced C_aO_2 and increased W_{insp} , resulted in a substantial degree of both peripheral and central fatigue. Two weeks of exposure to 5,260 m restored C_aO_2 to SL values but increased W_{insp} further over that observed in AH. The critical finding was that the rate of development of peripheral locomotor muscle fatigue failed to recover from AH to CH and was similar in both conditions. In contrast, the development of central fatigue was attenuated significantly in CH (vs. AH) but still greater compared with SL. Taken together, our findings suggest that acclimatization to high altitude attenuates the impact of AH on the development of central fatigue but fails to improve the exacerbated development of peripheral fatigue present during exercise in AH.

Peripheral Fatigue

Acute hypoxia. The cycling bout in AH was, compared with SL, characterized by a substantially exaggerated rate of peripheral fatigue (Table 2 and Fig. 2). These observations confirm numerous earlier findings using whole-body (4, 31, 57) and single-muscle exercise (28, 39).

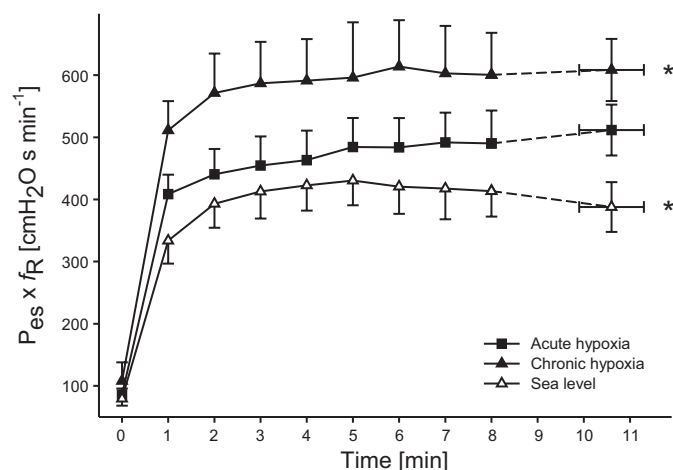


Fig. 1. Inspiratory muscle pressure-time product [esophageal pressure (P_{es}) × respiratory frequency (f_R)] during the identical constant-load cycling exercise performed in all 3 conditions. * $P < 0.05$ vs. acute hypoxia (AH), $n = 8$.

Table 2. Effects of constant-load cycling exercise on quadriceps muscle function

| | Percent Change from Pre- to Immediately Postexercise | | |
|-----------------------------|------------------------------------------------------|-----------------|------------------------|
| | Sea Level | Acute Hypoxia | Chronic Hypoxia |
| $Q_{tw,pot}$ | $-3.1 \pm 1.8^*$ | -20.9 ± 2.4 | -18.8 ± 3.4 |
| MRFD | $-4.1 \pm 2.5^*$ | -21.2 ± 4.2 | -17.9 ± 3.5 |
| MRR | $2.7 \pm 2.8^*$ | -13.2 ± 3.1 | -9.0 ± 2.2 |
| $RT_{0.5}$ | $1.0 \pm 2.2^*$ | 9.2 ± 1.3 | 8.2 ± 1.4 |
| MVC | $-1.3 \pm 1.2^*$ | -12.3 ± 1.2 | $-8.9 \pm 1.3^\dagger$ |
| Voluntary muscle activation | $-0.1 \pm 1.0^*$ | -6.9 ± 1.1 | $-3.7 \pm 1.2^\dagger$ |
| M-wave amplitude | $0.7 \pm 2.7^*$ | $2.5 \pm 2.0^*$ | -7.8 ± 2.1 |

Changes in muscle function are expressed as a percent change from pre-exercise baseline. All exercise trials were performed for the same duration (10.6 ± 0.7 min) and at the same absolute workload (138 ± 14 W). Values are expressed as means \pm SE. $Q_{tw,pot}$, potentiated single twitch; MRFD, maximal rate of force development; MRR, maximal rate of relaxation; $RT_{0.5}$, 1/2 relaxation time; MVC, maximal voluntary contraction force; M-wave, compound muscle action potential. Percent muscle activation is based on superimposed twitch technique. Various variables in acute and chronic hypoxia were, compared with baseline, altered significantly, 2.5 min after exercise ($P < 0.01$). *Not significantly different from pre-exercise baseline. $^\dagger P < 0.05$ vs. acute hypoxia, $n = 8$.

Compared with SL, C_aO_2 was approximately one-third lower and W_{insp} , $\sim 34\%$ higher during exercise in AH. These substantial alterations are known to contribute about equally to the exacerbated development of peripheral fatigue in AH (2). The impact of an acutely lowered C_aO_2 on muscle fatigability is mediated via the facilitating effects of the associated reduc-

tion in muscle O_2 delivery on the intramuscular accumulation of metabolites known to cause peripheral fatigue, i.e., hydrogen ion and inorganic phosphate (37, 61). The W_{insp} -induced exacerbation of peripheral fatigue results from the same intramuscular metabolic consequences associated with reductions in locomotor muscle O_2 delivery. However, in the case of the W_{insp} -related impairment in peripheral fatigue, the compromised O_2 delivery is the consequence of a sympathetically mediated impact on Q_L , secondary to the activation of the respiratory muscle metaboreflex (34). Taken together, the combined effects of a significantly reduced C_aO_2 and a higher W_{insp} has a profound impact on leg O_2 delivery and thus peripheral locomotor muscle fatigue (1).

Chronic hypoxia. Despite 2 wk of acclimatization to altitude, the rate of development of peripheral locomotor muscle fatigue was similar in AH and CH (Table 2 and Fig. 2). Somewhat conflicting data from earlier investigations suggest different mechanisms as a potential explanation of this finding. On the one hand, studies conducted by Reeves and colleagues (8, 64), following 2–3 wk at 4,300 m, report similar locomotor muscle O_2 delivery during submaximal endurance exercise in AH and CH. Given the critical dependency of the development of peripheral fatigue on muscle O_2 delivery, this similarity might explain the nearly identical levels of end-exercise locomotor muscle fatigue in AH and CH. On the other hand, experiments conducted at the same location as the present study (Mt. Chacaltaya, 5,260 m) have documented a significant improvement in leg muscle O_2 delivery from AH to CH, with the net

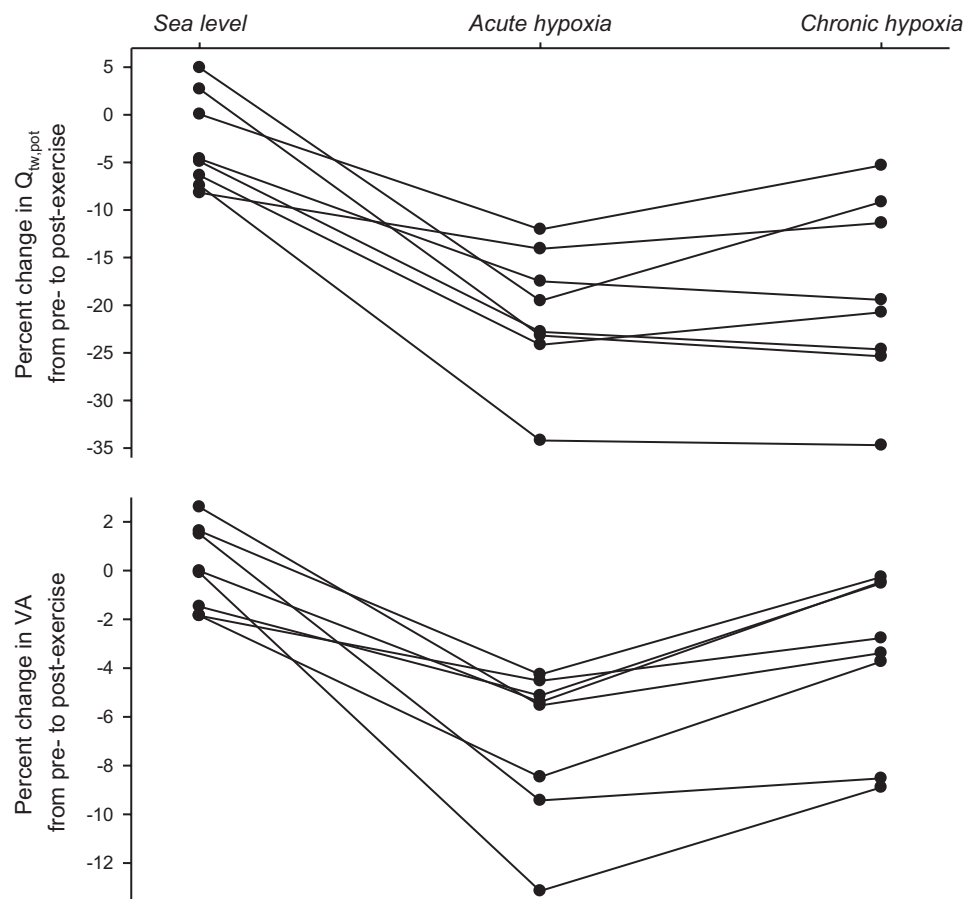


Fig. 2. Individual data illustrating the effects of constant-load bike exercise (138 ± 14 W; 10.6 ± 0.7 min) on potentiated quadriceps twitch force ($Q_{tw,pot}$; top) and voluntary muscle activation (VA; bottom) at sea level [SL; resting arterial oxygen (O_2) content: 19.3 ± 0.7 ml O_2 /dl] and in AH (17.3 ± 0.5 ml O_2 /dl) and chronic hypoxia (CH; 20.3 ± 1.3 ml O_2 /dl).

effect of similar values during submaximal bike exercise at SL and in CH (13). It might be important to emphasize that these latter experiments involved a greater altitude (5,260 m vs. 4,300 m) and a 9–10 wk acclimatization period vs. only a 2–3 wk period, as in the experiments by Reeves and colleagues (8, 64), as well as the present study. Regardless, based on the findings from the earlier Chacaltaya experiments, it appears that the similar degrees of end-exercise fatigue in AH and CH in the present study (Fig. 2) might have occurred in the face of a significant difference in bulk muscle O₂ delivery, i.e., higher in CH vs. AH.

Q_L was not measured directly in the present study. However, changes in THb, a NIRS-derived variable, are thought to reflect changes in regional blood volume and potentially Q_L (24, 59). The previously documented similarity in resting Q_L at SL, in AH, and in CH (11, 12, 36, 49, 50) is a critical prerequisite when using THb as an estimate of potential differences in Q_L and O₂ delivery during exercise. Since C_aO_2 was comparable at SL and CH (see RESULTS), the same exercise-induced increase in THb (Fig. 3) suggests a similar degree of O₂ delivery in these conditions. Furthermore, the combination of a lower C_aO_2 in AH vs. CH (and SL; see RESULTS) plus the similar increase in THb during exercise (Fig. 3) insinuates a lower locomotor muscle O₂ delivery in AH vs. CH (and by extension, SL). Both of these observations might support earlier blood flow studies conducted at the same location as the present experiments (13) but might contradict others performed at a lower altitude (8, 64). However, NIRS findings obtained from skeletal muscle need to be interpreted with caution. A significant limitation associated with NIRS is that this measurement is confined to a finite location, and changes in THb might not be representative of the whole muscle. Indeed, significant blood flow heterogeneity has been documented previously in skeletal muscle (35). Whereas heterogeneity diminishes with higher exercise intensities and is not affected by hypoxia (36), the exact location of NIRS probe placement from day to day is a potential source of error. To minimize this risk, we had strict criteria regarding probe placement (see METHODS), and at least two investigators independently assured correct probe positioning before each experiment.

Assuming that the similar degrees of peripheral fatigue in AH vs. CH occurred in the face of a greater O₂ delivery in CH, other, rather disadvantageous adaptations associated with acclimatization must have outweighed this benefit. A potential candidate is the documented impairment in the capacity of skeletal muscle to extract O₂ in CH, i.e., a decreased capillary muscle O₂ conductance (41). This impact might, despite a similar O₂ delivery at SL and in CH, potentially lower extracellular PO₂ to or beyond a previously suggested critical value (~30 Torr) associated with exacerbated development of peripheral fatigue (55). Alternatively, the higher O₂ delivery in CH vs. AH (13), combined with the same degree of peripheral fatigue, might suggest that C_aO_2 and bulk O₂ delivery, per se, might not depict key determinants of the exaggerated fatigability in hypoxia. Important here is the fact that despite the normalized C_aO_2 and bulk O₂ delivery in CH, P_aO_2 only partially recovers with acclimatization and remains fairly low in CH. This could hint toward a key role of P_aO_2 in exacerbating the development of peripheral fatigue at altitude.

In CH, V_E was ~20% higher compared with AH. Given the substantially lower air density at 5,260 m (0.64 kg/m³ vs. 1.18

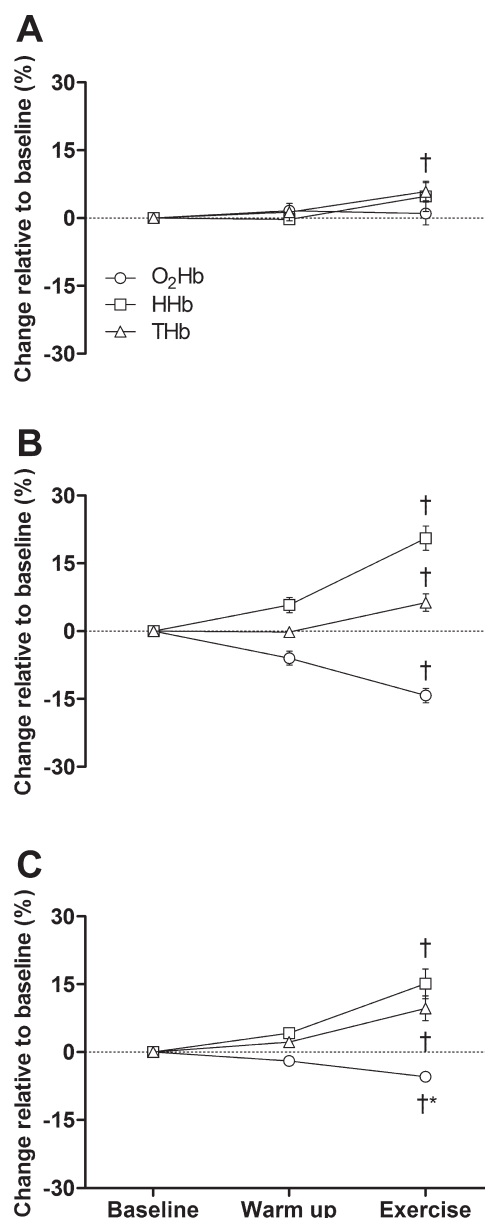


Fig. 3. Vastus lateralis oxygenation at resting baseline, during the final 30 s of a 3-min warm-up (28 W), and during the final 30 s of constant-load exercise (131 W) at SL (A), in AH (B), and in CH (C). † $P < 0.05$ vs. respective baseline; * $P < 0.05$ vs. AH, $n = 8$. O₂Hb, oxyhemoglobin; HHb, deoxyhemoglobin; THb, total hemoglobin.

kg/m³ at 130 m, where AH experiments occurred), it could be argued that in terms of respiratory muscle work, the reduced density might balance the acclimatization-induced increase in V_E , with the net effect of a similar W_{insp} in CH and AH. However, W_{insp} was, similar to V_E , ~20% higher in CH vs. AH. This observation, per se, might suggest that the lower air density at altitude had no effect on the relationship between minute V_E and respiratory muscle work. However, it has been shown that bronchoconstriction, associated with severe hypoxia, increases the resistive component of respiratory work and offsets the theoretical benefit of a reduced air density (22). This results in a similar respiratory muscle work for a given V_E at altitude and at SL (18). Therefore, any increase in W_{insp}

observed in hypobaric CH is attributable to the exaggerated ventilatory response associated with altitude acclimatization.

The increase in minute V_E in the present study was mainly due to the increase in V_T ; f_R was similar in both conditions. The higher V_T was achieved via reductions in EELV (Table 1), which is compared with increasing EILV to raise V_T , more economical, since higher lung volumes are associated with a reduced compliance (38). We therefore conclude that the 23% higher W_{insp} at the same workload in CH vs. AH resulted from the substantially higher V_E following acclimatization. Finally, this exaggerated W_{insp} likely aggravated the respiratory muscle metaboreflex and associated impact on leg vascular conductance (25) and presumably blunted exercise Q_L more in CH compared with AH.

In contrast to our findings, it was suggested previously that acclimatization to high altitude might eliminate the impact of AH on the rate of development of fatigue during single muscle exercise (adductor pollicis) and restore it to that observed at SL (28). However, submaximal, intermittent exercise, including a small muscle mass, does not maximally challenge O_2 delivery and use. Therefore, the observed positive effect could, at least in part, be explained by the use of the available reserve capacity. Specifically, various compensatory mechanisms, including increases in cardiac output and muscle O_2 delivery and extraction, could have reduced the hypoxia-induced impact on the development of fatigue. Such an effective compensation might not—or only to a much smaller degree—be possible during intense, whole-body exercise, performed close to a human's maximal circulatory and ventilatory capacity (14, 15).

CH had a significant impact on the effect of exercise on M-wave amplitude. Reductions in M-wave amplitude have been associated with decreases in sarcolemma excitability (19). The attenuated excitability results from reduced sarcolemma sodium (Na^+)-potassium (K^+)-ATPase activity (46) and can contribute to compromised muscle force output (21). Pre-exercise M-wave amplitudes (and $Q_{tw,pot}$) in our experiments were similar in all three conditions. This suggests that neither severe AH nor CH impairs sarcolemma Na^+ - K^+ -ATPase activity and membrane excitability of resting locomotor muscle. This confirms earlier findings (40); however, it contrasts with others (16) who report decreased resting M-wave amplitudes following 10 days of exposure to severe hypoxia (>4,300 m). Regardless, although M-wave amplitudes did not change from pre- to postexercise at SL and in AH, we observed, in contrast to Garner et al. (30), a significant exercise-induced decrease in CH (Table 2). AH has recently been shown to have no effect on exercise-induced changes in Na^+ - K^+ -ATPase activity, which explains the similar M-wave behavior in SL and AH (51). However, altitude acclimatization causes a downregulation of Na^+ - K^+ -ATPase pump concentration, and although this does not alter resting M-wave characteristics, it likely explains the exercise-induced decrease in M-wave amplitude observed in CH (20, 33).

The lower postexercise M-wave amplitude in CH indicates a failure of the motor nerve/sarcolemma to propagate evoked stimuli to the contractile apparatus and might have masked potential benefits of acclimatization on fatigue resistance. Put simply, postexercise twitch forces might have been larger in CH if M-waves had remained unchanged from pre-exercise. If so, this would have resulted in a smaller exercise-induced reduction in $Q_{tw,pot}$ in CH. Regardless, failure of neuromuscular transmission/sarcolemmal excitability contributes to reduced force output in response to a given central nervous activation and can therefore be

considered a key determinant of the impaired fatigue resistance in CH.

Central Fatigue

Exercise in AH induced a substantial degree of central fatigue, which was attenuated by ~50% when the same trial was repeated in CH (Table 2). This significant improvement, associated with acclimatization, clearly contrasts with the absence of a beneficial effect of CH on peripheral fatigue, as described above. Since the development of central fatigue is highly sensitive to O_2 (1), we attribute this improvement to the effects of high-altitude acclimatization on O_2 availability within the brain. Specifically, the cerebral O_2 delivery index at the end of exercise in CH was improved from AH (Table 1) (65) and may explain the lower degree of central fatigue in CH vs. AH.

Despite the similar CBFv and a slightly higher brain O_2 delivery in CH vs. SL (Table 1), which agrees with earlier Chacaltaya studies using the Kety-Schmidt technique to measure CBF/ O_2 delivery (44), exercise-induced central fatigue was greater in CH. Two considerations discussed previously might account for this observation. First, the significant degree of peripheral fatigue in CH (vs. no fatigue at SL) presumably facilitated central fatigue via increases in inhibitory neural feedback from locomotor muscle (mediated by *group III/IV* muscle afferents), which limit central motor drive (3, 6). Second, although C_aO_2 and brain O_2 delivery were similar/higher in CH vs. SL, the still substantially lower P_aO_2 might have contributed to the greater degree of central fatigue during exercise in this condition. Indeed, a low P_aO_2 was recently suggested to impair cerebral metabolism (48) and alterations in neurotransmitter turnover (23), and both of these factors have been linked to the development of central fatigue (17, 53).

Taken together, the current findings provide a global indication of the positive effects of altitude acclimatization on the development of central fatigue during exercise. However, we cannot comment on the specific sites of the central motor pathway involved or the relative contribution of C_aO_2 and P_aO_2 in mediating these beneficial adaptations.

Implications of Findings for Performance-Related Questions in CH

AH generally impairs endurance exercise performance (60). Prolonged exposure to hypoxia is known to recover some of this impairment (29, 52); however, SL performance is never matched at altitude. Our current findings indicate that the acclimatization-induced partial recovery of endurance performance occurs independent of any improvement of peripheral locomotor muscle fatigue from AH to CH. This insinuates that peripheral locomotor muscle fatigability, per se, does not contribute to the improvement of endurance performance observed from AH to CH. We therefore propose that the significantly attenuated central fatigue during exercise in severe CH likely accounts, at least in part, for the improvement of endurance performance associated with altitude acclimatization.

Mechanisms underlying the hypoxia-induced curtailment of central motor drive (i.e., increase in central fatigue) and endurance exercise performance have been documented previously to differ depending on the severity of arterial hypoxemia. Specifically, peripheral fatigue might depict the dominant determinant of central motor drive and thus the limiting factor

above 70–75% S_pO_2 . At more severe degrees of hypoxemia (<70% S_pO_2), central motor drive and endurance performance might primarily—but not exclusively—be determined/limited by central nervous system (CNS) hypoxia (5). Since peripheral fatigue did not change with acclimatization in the present study, but S_pO_2 increased from below to above the “threshold” described previously (5), reductions in central fatigue might be mediated mainly by improved arterial oxygenation and associated smaller influence of CNS hypoxia on central motor drive.

A recent Point:Counterpoint debate in this journal has focused on the potential existence/relevance of differences in physiological responses to exercise performed in normobaric vs. hypobaric hypoxia (43, 45). Since the present AH and CH experiments were performed in normobaric and hypobaric hypoxia, respectively, these potential differences, if indeed existent, might have influenced our findings.

Conclusion

AH exacerbates central and peripheral fatigue during endurance exercise. Our experiments indicate that acclimatization to high altitude significantly attenuates the development of central fatigue but does not improve the development of peripheral fatigue observed during whole-body endurance exercise in AH.

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DISCLOSURES

The authors declare no conflicts of interest.

AUTHOR CONTRIBUTIONS

Author contributions: M.A., S.G., and A.W.S. conception and design of research; M.A., S.G., R.T., and A.W.S. performed experiments; M.A., S.G., and R.T. analyzed data; M.A., S.G., R.T., and A.W.S. interpreted results of experiments; M.A. and S.G. prepared figures; M.A. and A.W.S. drafted manuscript; M.A., S.G., R.T., A.W.S., A.T.L., and R.C.R. edited and revised manuscript; M.A., S.G., R.T., A.W.S., A.T.L., and R.C.R. approved final version of manuscript.

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AltitudeOmics: Exercise-induced supraspinal fatigue is attenuated in healthy humans after acclimatisation to high altitude

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Abstract

Aims: We asked whether acclimatisation to chronic hypoxia (CH) attenuates the level of supraspinal fatigue that is observed after locomotor exercise in acute hypoxia (AH). **Methods:** Seven recreationally-active participants performed identical bouts of constant-load cycling (131 ± 39 W, 10.1 ± 1.4 min) on three occasions: 1) in normoxia (N, P_iO_2 , 147.1mmHg); 2) in AH (F_iO_2 , 0.105; P_iO_2 , 73.8mmHg); 3) after 14 days in CH (5,260m; P_iO_2 , 75.7mmHg). Throughout trials, prefrontal-cortex tissue oxygenation and middle cerebral artery blood velocity (MCA_v) were assessed using near-infrared-spectroscopy and transcranial Doppler sonography. Pre- and post-exercise twitch responses to femoral nerve stimulation and transcranial magnetic stimulation were obtained to assess neuromuscular and corticospinal function. **Results:** In AH, prefrontal oxygenation declined at rest ($\Delta 7\pm 5\%$) and end-exercise ($\Delta 26\pm 13$) ($P<0.01$); the degree of deoxygenation in AH was greater than N and CH ($P<0.05$). The cerebral O_2 delivery index ($MCA_v\times C_aO_2$) was $19\pm 14\%$ lower during the final minute of exercise in AH compared to N ($P=0.013$) and $20\pm 12\%$ lower compared to CH ($P=0.040$). Maximum voluntary and potentiated twitch force were decreased below baseline after exercise in AH and CH, but not N. Cortical voluntary activation decreased below baseline after exercise in AH ($\Delta 11\%$, $P=0.014$), but not CH ($\Delta 6\%$, $P=0.174$) or N ($\Delta 4\%$, $P=0.298$). A twofold greater increase in motor evoked potential amplitude was evident after exercise in CH compared to AH and N. **Conclusion:** These data indicate that exacerbated supraspinal fatigue after exercise in AH is attenuated after 14 days of acclimatisation to altitude. The reduced development of supraspinal fatigue in CH may have been attributable to increased corticospinal excitability, consequent to an increased cerebral O_2 delivery.

Key words: adaptation, altitude, exercise, transcranial magnetic stimulation

Glossary

C_aO_2 , arterial O_2 content; CSP, cortical silent period; ERT, estimated resting twitch; F_iO_2 , fraction of inspired O_2 ; f_R , respiratory frequency; [Hb], haemoglobin concentration; MCA_v , middle cerebral artery blood velocity; MEP, motor evoked potential; M_{max} , maximum M-wave; MVC, maximum voluntary contraction; P_aO_2 , partial pressure of arterial O_2 ; P_iO_2 , partial pressure of inspired O_2 ; $Q_{tw,pot}$, potentiated quadriceps twitch force; rMT, resting motor threshold; SIT, superimposed twitch; S_pO_2 , arterial O_2 saturation; TMS, transcranial magnetic stimulation; $\dot{V}CO_2$, carbon dioxide output; \dot{V}_E , minute ventilation; $\dot{V}O_2$, oxygen uptake; V_T , tidal volume.

Introduction

The mechanisms underpinning impairments in exercise performance in hypoxia are not fully understood, but multiple peripheral and central mechanisms of fatigue have been proposed (Amann and Calbet, 2008, Nybo and Rasmussen, 2007, Perrey and Rupp, 2009). The rate of development of peripheral fatigue is increased during intense locomotor exercise in acute hypoxia (Amann *et al.*, 2006b, Goodall *et al.*, 2012). This has been documented in numerous human studies as an increased decline in the force response to motor nerve stimulation after exercise and an increased rate of rise in electromyogram (EMG) signals during exercise (Amann and Calbet, 2008). Amann *et al.* (2006a) suggested that the accelerated development of peripheral fatigue and associated intramuscular metabolic changes in acute moderate hypoxia restricts central motor drive preventing excessive end-exercise locomotor muscle fatigue under conditions of attenuated arterial oxygenation. It was subsequently demonstrated that in acute severe hypoxia, peripheral fatigue becomes the less important variable and the primary limitation to exercise transfers to a hypoxia-sensitive central component of fatigue (Amann *et al.*, 2007). Less is known about the mechanism(s) of fatigue during locomotor exercise in chronic hypoxia. We recently reported the accelerated development of peripheral fatigue after locomotor exercise in acute hypoxia to be similar after a period of acclimatisation (14 days) to high altitude; conversely, the level of central fatigue was attenuated (Amann *et al.*, 2013). The measure of central fatigue, however, was determined using peripheral stimulation and the responsiveness of the brain-to-muscle pathway after a period of chronic hypoxia remains unknown.

Transcranial magnetic stimulation (TMS) has been used to specify the site of fatigue within the central nervous system in acute severe hypoxia (Goodall *et al.*, 2012, Goodall *et al.*, 2010). When TMS is delivered over the motor cortex during a maximal voluntary contraction (MVC), it is possible to detect a twitch-like increment in force in the active muscle. That is, despite maximal effort, motor cortical output at the time of stimulation is insufficient to drive the motoneurons maximally. An increase in this increment in force after exercise provides evidence of a reduced cortical voluntary activation, indicative of supraspinal fatigue (Gandevia *et al.*, 1996, Todd *et al.*, 2003). Further, EMG recordings in response to cortical stimuli (motor evoked potential [MEP]) can be monitored to assess changes in excitability of the brain to muscle pathway. Descending volleys evoked from cortical stimulation depend on the stimulus intensity and excitability of corticospinal cells, whereas responses in the muscle depend on transmission through relevant excitatory and inhibitory interneurons and excitability of the motoneuron pool (Taylor and Gandevia, 2001). Hypoxia affects

neuronal function *in-vitro* (Nieber *et al.*, 1999), however, acute hypoxia appears to have negligible effects on resting MEPs elicited by TMS (Goodall *et al.*, 2010, Rupp *et al.*, 2012, Szubski *et al.*, 2006). A MEP evoked during muscular contraction is followed by an interval of EMG silence, the so-called cortical silent period (CSP). The initial phase of the CSP has been attributed to inhibitory spinal mechanisms (Inghilleri *et al.*, 1993), whereas the later period (>100 ms) represents increased cortical inhibition (Chen *et al.*, 1999, Inghilleri *et al.*, 1993, Taylor and Gandevia, 2001). Szubski *et al.* (2006) found a shorter CSP in acute hypoxia, suggestive of a reduced corticospinal inhibition during the exercise.

Responsiveness of the corticospinal pathway and the associated development of central fatigue after locomotor exercise during periods of prolonged hypoxia have not been studied. A recent investigation found an increase in corticospinal excitability (increased resting MEP) after a period of prolonged acute hypoxia (Rupp *et al.*, 2012); however, the mechanisms for this response and the associated effects upon the development of central fatigue during locomotor exercise have not been studied. We have recently related the development of supraspinal fatigue during exercise in severe acute hypoxia to a reduction in cerebral O₂ availability (Goodall *et al.*, 2012). Acclimatisation to altitude not only brings about improvements in arterial oxygenation, but also improvements in cerebrovascular function (Ainslie and Ogoh, 2009, Lucas *et al.*, 2011). It is unknown how haematologic (e.g., hemodynamic and cerebrovascular) adaptations might serve to impact corticospinal excitability and the development of supraspinal fatigue during locomotor exercise in chronic hypoxia. Accordingly, the aim of the present study was to assess corticospinal excitability and supraspinal fatigue after locomotor exercise in chronic hypoxia. We hypothesised that altered cerebrovascular and corticospinal responses after a period of acclimatisation to high altitude would reduce the severity of supraspinal fatigue compared to that observed in acute hypoxia.

Methods

Ethical Approval

All procedures conformed to the Declaration of Helsinki and were approved by the Universities of Colorado Denver, Oregon and Utah Institutional Review Boards and the US Department of Defense Human Research Protection Office.

Participants

This study was conducted as part of the AltitudeOmics project examining the integrative physiology of human responses to hypoxia (Subudhi *et al.* under review at PLoSOne). After written informed consent, seven (five male) recreationally active sea level inhabitants participated in the study (mean \pm SD age, 21 ± 1 yr; stature, 1.78 ± 0.10 m; body mass, 69 ± 11 kg; maximum O₂ uptake [$\dot{V}O_{2max}$], 46.4 ± 8.2 ml·kg⁻¹·min⁻¹ [participant IDs: 1,2,3,5,6,7,10]). The participants were non-smokers, free from cardiorespiratory disease, born and raised at <1500 m, and had not travelled to elevations >1000 m in the 3 months prior to investigation. Participants arrived at the laboratory in a rested and fully hydrated state, at least 3 h postprandial, and avoided strenuous exercise in the 48 h preceding each trial. They also refrained from caffeine for 12 h before each test, while alcohol and prophylactic altitude medication were prohibited for the entire duration of the investigation. All of the subjects participated in a companion study investigating the acclimatisation-induced effects on peripheral measures of neuromuscular fatigue (Amann *et al.*, 2013); while the data were obtained from the same protocol described below, the primary TMS and cerebral oxygenation related outcome measures in the current study do not overlap with previous analyses.

Experimental design

Participants completed a preliminary trial and three experimental trials. Each trial was conducted at the same time of day, and separated by at least 5 d during a 12 wk period. During the preliminary trial, participants were thoroughly familiarized with the methods used to assess neuromuscular function and performed a maximal incremental exercise test in normoxia for the determination of $\dot{V}O_{2max}$ and peak workload (W_{peak}); further maximal incremental tests were performed in AH and CH (Subudhi *et al.* under review at PLoSOne). During the experimental trials, participants performed constant-load exercise at a workload equal to 50% W_{peak} obtained in the preliminary trial: 1) to the limit of tolerance in acute normobaric hypoxia (AH: F_iO₂ = 0.105; Eugene, Oregon, barometric pressure [BP] = 750 ± 2 mmHg; P_iO₂ = 73.8 ± 0.2 mmHg); 2) for the same absolute intensity and duration as in trial 1, but in normoxia (N: Eugene, Oregon, BP = 750 ± 2 mmHg; P_iO₂ = 147.1 ± 0.5 mmHg); and 3) for the same absolute intensity and duration as in trial 1, but after 14 d at 5,260 m above sea level (CH: Mt. Chacaltaya, Bolivia, BP = 409 ± 1 mmHg; P_iO₂ = 75.7 ± 0.1 mmHg). Participants were flown to La Paz, Bolivia where they spent two nights at low altitude (Coroico, 1,525 m), before being driven to the Chacaltaya Research Station at 5,260 m. Before and within 2.5 min after each exercise trial, twitch responses to supramaximal femoral nerve stimulation and TMS were obtained to assess fatigue. During AH, the post-exercise measurements were made while

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participants continued to breathe the hypoxic gas. Cerebrovascular, cardiorespiratory and perceptual responses, as well as EMG activity of the vastus lateralis (VL), were assessed throughout each trial.

Force and EMG recordings

Knee-extensor force during voluntary and evoked contractions was measured using a calibrated load cell (Tedea, Basingstoke, UK). The load cell was fixed to a custom-built chair and connected to a non-compliant cuff attached around the participant's right leg just superior to the right ankle. Participants sat upright in the chair with the hips and knees at 90° of flexion. EMG activity was recorded from the VL and biceps femoris (BF). Surface electrodes were placed 2 cm apart over the muscle bellies and a reference electrode was placed over the patella. The electrodes were used to record the compound muscle action potential (M-wave) elicited by electrical stimulation of the femoral nerve and the MEP elicited by TMS. Signals were amplified (gain 1000; Force: custom-built bridge amplifier; EMG: PowerLab 26T, ADInstruments Inc, Oxfordshire, UK), band-pass filtered (EMG only: 20-2000 Hz), digitised (4 kHz; PowerLab 26T, ADInstruments Inc), acquired and later analysed (LabChart v7.0, ADInstruments Inc).

Neuromuscular function

Force and EMG variables were assessed before and immediately after each exercise trial. Prior to each trial, MVC force was determined from three, 3 s contractions. Femoral nerve stimulation was delivered at rest ~2 s after the MVC to determine the potentiated quadriceps twitch force ($Q_{tw,pot}$). TMS was delivered during brief (~5 s) maximal and submaximal voluntary contractions for the determination of cortical voluntary activation. Each set of contractions comprised 100, 75, and 50% MVC efforts separated by ~5 s of rest. The contraction sets were repeated three times, with 15 s between each set. Visual feedback of the target force was provided via a computer monitor.

Femoral nerve stimulation

Single electrical stimuli (200 μ s) were delivered to the right femoral nerve via surface electrodes (CF3200, Nidd Valley Medical Ltd, North Yorkshire, UK) and a constant-current stimulator (DS7AH, Digitimer Ltd, Welwyn Garden City, Hertfordshire, UK). The cathode was positioned over the nerve high in the femoral triangle; the anode was placed midway between the greater trochanter and the iliac crest. The site of stimulation that produced the largest resting twitch amplitude and M-wave (M_{max}) was located. Single stimuli were delivered beginning at 100 mA and increasing by 20 mA until

plateaus occurred in twitch amplitude and M_{\max} . Supramaximal stimulation was ensured by increasing the final intensity by 30% (mean current 253 ± 60 mA).

Transcranial magnetic stimulation

TMS was delivered via a concave double cone coil (110 mm diameter; maximum output 1.4 T) powered by a mono-pulse magnetic stimulator (Magstim 200, The Magstim Company Ltd, Whitland, UK). The coil was held over the vertex to preferentially stimulate the left hemisphere (postero-anterior intracranial current flow), and was placed in an optimal position to elicit a large MEP in the VL and a small MEP in the antagonist (BF). The optimal coil position was marked on the scalp with indelible ink to ensure reproducibility of the stimulation. Resting motor threshold (rMT) was determined at the beginning of each experimental trial. Briefly, TMS was first delivered with the coil placed over the optimal site of stimulation at a sub-threshold intensity of 35% maximum stimulator output. Stimulus intensity was then increased in 5% steps until consistent motor evoked potentials (MEPs) with peak-to-peak amplitudes of more than $50 \mu\text{V}$ were evoked. Thereafter, stimulus intensity was reduced in 1% steps until an intensity was reached that elicited an MEP of at least $50 \mu\text{V}$ in 5 out of 10 trials (Groppa *et al.*, 2012). The stimulation intensity that elicited rMT was increased by 30%; thus, the experimental stimulation intensity was 130% of rMT. This stimulation intensity elicited a large MEP in the VL (area between 60 and 100% of M_{\max} during knee-extensor contractions $\geq 50\%$ MVC; Figure 1); indicating the TMS stimulus activated a high proportion of knee extensor motor units, while causing only a small MEP in the BF (amplitude $< 20\%$ of MEP during knee-extensor contractions).

Constant-load exercise

Participants sat on an electromagnetically-braked cycle ergometer (Velotron Dynafit Pro, Racermate, Seattle, WA) while baseline cardiorespiratory and cerebrovascular data were collected for 3 min. The participants warmed-up for 5 min at 10% W_{peak} (26 ± 8 W) before the workload was increased to 50% normoxic W_{peak} (131 ± 39 W). This intensity was chosen to maximise the tolerable duration of exercise in the hypoxic conditions. The participants remained seated throughout exercise and maintained a target pedal cadence equivalent to that chosen during the incremental exercise test (88 ± 3 rpm). Task-failure was reached when cadence dropped below 60% of the target rpm for > 5 s. Constant load exercise was performed firstly in AH; the achieved time (10.1 ± 1.4 min) was then replicated in N and CH.

Tissue oxygenation and cerebrovascular responses

Cerebral oxygenation was assessed using a multi-channel NIRS instrument (Oxymon III, Artinis) (Subudhi *et al.*, 2009, Subudhi *et al.*, 2011). Changes in oxygenated, deoxygenated and total cerebral haeme concentrations (μM) were expressed relative to the resting baseline recorded in each experimental condition. Arterial oxygen saturation was estimated using forehead pulse oximetry ($S_p\text{O}_2$; Model N-595, Nellcor, Pleasanton, CA). Excellent agreement between the pulse oximeter and arterial O_2 saturation across the range of values in the present study has been published (Romer *et al.*, 2007). Hemoglobin concentration [Hb] was measured (OSM-3, Radiometer, Copenhagen, Denmark) in resting arterial blood samples. Samples were collected during the primary physiological protocols at sea level (2-4 d prior to the first exercise trial in the present study) and on the 16th day at 5,260 m (2 d following the constant load exercise trial in the present study) (Subudhi *et al.* under review at PLoSOne). Arterial O_2 content ($C_a\text{O}_2$) was estimated using the equation: $([\text{Hb}] \times 1.39 \times S_p\text{O}_2 / 100)$. Resting [Hb] in combination with the measured $S_p\text{O}_2$ during the exercise protocol were used to obtain $C_a\text{O}_2$ throughout exercise in all conditions. Blood velocity in the left middle cerebral artery (MCA_v) was determined using transcranial Doppler (Spencer Technologies, Seattle, WA). The custom-made NIRS headset was modified to hold a 2 MHz probe positioned over the left temporal window. Measurements were optimised at an average penetration depth of 50 ± 3 mm. An index of cerebral O_2 delivery was calculated as the product of MCA_v and $C_a\text{O}_2$. It was assumed that changes in MCA_v would reflect changes in cerebral blood flow based on evidence that the middle cerebral artery diameter changes minimally in response to hypoxia and hypocapnia (Poulin and Robbins, 1996).

Cardiorespiratory and perceptual responses

Ventilatory and pulmonary gas exchange indices were assessed using an online system (in AH & N Medical Graphics PFX, St. Paul, MN, USA; & in CH Oxigraf O_2cap , Mountain View, CA, USA). Heart rate was identified from the peak MCA_v envelopes. Ratings of perceived exertion for dyspnea and limb discomfort were obtained using the CR10 scale at baseline and every minute throughout exercise (Borg, 1982). In CH, symptoms of acute mountain sickness were assessed on the day of a trial using the Lake Louise Score (Roach *et al.*, 1993).

Data analysis

Cortical voluntary activation was assessed by measuring the force responses to motor-cortex stimulation during submaximal and maximal contractions. Corticospinal excitability increases during

voluntary contraction (Rothwell *et al.*, 1991); thus, we estimated the amplitude of the resting twitch evoked by TMS (ERT; Goodall *et al.*, 2009, Sidhu *et al.*, 2009a). Cortical voluntary activation (%) was subsequently quantified using the equation: $(1 - [\text{SIT} / \text{ERT}] \times 100)$.

The peak-to-peak amplitude and area of evoked MEPs and M_{max} were measured offline. To ensure the motor cortex stimulus activated a high proportion of the knee-extensor motor units, the area of vastus lateralis MEP was normalised to that of M_{max} elicited during the MVC at the beginning of each trial (Taylor *et al.*, 1999) (Figure 1). The duration of the CSP evoked by TMS during MVC was quantified as the duration from stimulation to the continuous resumption of post-stimulus EMG exceeding ± 2 SD of pre-stimulus EMG (>50 ms prior to stimulus). VL EMG signals during exercise were rectified and smoothed (15 ms), then quantified as the mean integrated area during each cycle revolution and averaged over each minute of exercise. A computer algorithm identified the onset and offset of activity where the rectified EMG signals deviated >2 SD from baseline for >100 ms.

Reliability coefficients

On a separate day, the responses to TMS, femoral nerve stimulation and MVC were repeated twice in all participants. The two assessment procedures were separated by a 2 min walk followed by 5 min of rest. Coefficient of variation (CV) and intraclass correlation coefficient (ICC) were calculated to evaluate test-retest reliability. All correlations were statistically significant and indicated, in combination with the CVs, a high level of reproducibility: cortical voluntary activation, CV = 1.4%, ICC = 0.82; CSP, CV = 7.1%, ICC = 0.93; ERT, CV = 10.2%, ICC = 0.84; MEP/ M_{max} , CV = 9.6%, ICC = 0.66; M_{max} , CV = 11.4%, ICC = 0.98; 100% MVC MEP, CV = 14.1%, ICC = 0.96; 75% MVC MEP, CV = 10.2%, ICC = 0.98; 50% MVC MEP, CV = 7.2%, ICC = 0.99; MVC, CV = 4.7%, ICC = 0.94; $Q_{\text{tw,pot}}$, CV = 4.8%, ICC = 0.97.

Statistical analysis

Data are presented as means \pm SD in the text and means \pm SE in the figures. A 3×2 repeated measures ANOVA on condition (3 [AH, N, CH]) and time (2 [pre, post]) was used to test for within-group differences. When ANOVA revealed significant interactions, post-hoc comparisons were made using the least significant differences test. Statistical significance was set at $P < 0.05$. All analyses were conducted using SPSS (v19, IBM Corporation, New York, USA).

Results

Exercise responses

The exercise workload was 131 ± 39 W (50% N W_{peak}), which equated to 83% W_{peak} in AH and 74% W_{peak} in CH. Cerebral oxygenation data are shown in Figure 2. During N, oxyhaemoglobin was unchanged from baseline to warm up and total haemoglobin was increased during the final minute of exercise ($P = 0.658$ and 0.007 , respectively). During AH, deoxygenated haemoglobin increased from baseline to warm up ($P = 0.006$); this response was exaggerated towards end exercise ($P < 0.001$). During CH, deoxygenated haemoglobin increased at end exercise ($P = 0.015$) in line with increased total haemoglobin ($P = 0.043$). Overall, these results demonstrate that the degree of cerebral deoxygenation (Δ deoxygenated haemoglobin) in AH was greater than that observed in N and CH ($P < 0.05$).

S_pO_2 and MCA_v data are shown in Figure 3. Acute exposure to hypoxia decreased S_pO_2 at rest ($\Delta 7 \pm 4\%$; $P = 0.009$) and during the final minute of exercise ($\Delta 34 \pm 10\%$; $P < 0.001$). Resting S_pO_2 in CH was $85 \pm 2\%$ ($P < 0.001$ vs. N; $P = 0.330$ vs. AH), and in the final minute of exercise had fallen to $78 \pm 5\%$ ($P < 0.001$ vs. N; $P = 0.002$ vs. AH). No changes in S_pO_2 were apparent in N ($P > 0.702$). Resting MCA_v did not differ between conditions at baseline (pooled average, 54 ± 9 $\text{cm}\cdot\text{s}^{-1}$; $P = 0.544$). MCA_v did not increase from rest at any time point in N ($P > 0.108$). MCA_v increased from rest to the final minute of exercise in AH ($40 \pm 15\%$; $P < 0.001$) and CH ($25 \pm 14\%$; $P = 0.016$), but did not differ between conditions (Figure 3).

Hemoglobin concentration was 1.42 ± 0.03 $\text{g}\cdot\text{L}^{-1}$ in N and 1.63 ± 0.31 $\text{g}\cdot\text{L}^{-1}$ in CH ($P = 0.005$). Resting P_aO_2 was reduced in AH compared to N (39.1 ± 4.8 vs. 103.3 ± 8.7 mmHg, $P < 0.001$), was increased in CH relative to AH (58.8 ± 3.2 mmHg, $P < 0.001$), but was still lower than N ($P < 0.001$). C_aO_2 was lower at rest in AH vs. N (19.8 ± 1.9 vs. 21.5 ± 2.9 $\text{ml}\cdot\text{dl}^{-1}$; $P = 0.013$); during the final minute of exercise C_aO_2 in AH was $36 \pm 8\%$ lower than N ($P < 0.001$) and $22 \pm 9\%$ lower than in CH ($P = 0.001$). C_aO_2 was lower at rest in CH vs. N (19.4 ± 2.6 vs. 21.5 ± 2.9 $\text{ml}\cdot\text{dl}^{-1}$; $P < 0.001$) and during the final minute of exercise (17.6 ± 2.9 vs. 21.2 ± 2.9 $\text{ml}\cdot\text{dl}^{-1}$; $P = 0.725$). Consequently, cerebral O_2 delivery index ($MCA_v \times C_aO_2$) was $19 \pm 14\%$ lower during the final minute of exercise in AH compared to N ($P = 0.013$) and $20 \pm 12\%$ lower compared to CH ($P = 0.040$). No differences were evident between N and CH at rest ($P = 0.783$) or during the final minute of exercise ($P = 0.797$) (Figure 3).

Cardiorespiratory data are shown in Table 1. Respiratory frequency and minute ventilation (\dot{V}_E) rose substantially over time in all conditions. $\dot{V}_E/\dot{V}CO_2$ during the final minute of exercise in AH and CH was approximately twofold greater than in N ($P < 0.001$); $\dot{V}_E/\dot{V}CO_2$ during the final minute of exercise was 28% higher in CH compared to AH ($P < 0.001$). During the final minute of exercise, whole body $\dot{V}O_2$ was not different across the three conditions ($P = 0.411$). Dyspnea and limb discomfort at end-exercise were higher in AH compared to N ($P < 0.001$ and $P = 0.048$, respectively), but were not different compared to CH ($P = 0.714$ and 0.549 , respectively). Integrated EMG activity at end exercise was higher in AH compared to N (32%; $P = 0.029$), but not CH (16%; $P = 0.303$). There were no reported symptoms of acute mountain sickness during CH.

Pre- and post-exercise responses

Peripheral and central measures of excitability are shown in Table 2.

Neuromuscular responses

MVC did not differ between conditions at baseline (AH, 392 ± 77 N; N, 386 ± 90 N; CH, 376 ± 39 N; $P = 0.942$). MVC was reduced post-exercise in AH (339 ± 77 N, $P = 0.011$) and CH (346 ± 93 N, $P = 0.032$), but not N (387 ± 87 N, $P = 0.684$). The reductions in MVC were not different between conditions ($P \geq 0.119$). $Q_{tw,pot}$ did not differ between conditions at baseline (AH, 107 ± 13 N; N, 105 ± 12 N; CH, 110 ± 16 N; $P = 0.752$). $Q_{tw,pot}$ was reduced post-exercise in AH (84 ± 14 N, $P = 0.005$) and CH (90 ± 18 N, $P = 0.011$), but not N (102 ± 12 N, $P = 0.692$). On average, resting M_{max} in CH displayed a twofold increase compared to AH and N ($P < 0.019$); however, the change in M_{max} during MVC was not statistically significant ($P > 0.058$). Neither measure of M_{max} changed pre- to post-exercise in any condition ($P \geq 0.610$). Pooled across conditions, pre-exercise ERT (mean $r^2 = 0.95$) was 70% of the pre-exercise $Q_{tw,pot}$ and did not differ between conditions (mean ERT 75 ± 25 N; $P = 0.811$). Post-exercise ERT was reduced in AH (52 ± 27 N, $P = 0.049$), but was unchanged in N and CH ($P \geq 0.107$).

Corticomotor responses

rMT in AH, N and CH was 54 ± 5 , 53 ± 3 and $51 \pm 6\%$ maximum stimulator output ($P = 0.276$), respectively. During CH, resting MEP amplitude was twofold greater compared to AH ($P = 0.014$) and N ($P = 0.014$). Exercise elicited a reduction in resting MEP amplitude in CH ($P = 0.022$), but not AH ($P = 0.346$) or N ($P = 0.369$). MEPs evoked during brief knee extensor contractions at 100, 75 and 50% MVC pre-exercise were higher in CH compared to AH ($P < 0.020$) and N ($P < 0.030$) (see also Figure

4). MEPs evoked during the brief knee-extensor contractions (50-100% MVC) post-exercise were not significantly different from pre-exercise values in any condition. MEP amplitude, however, was higher post-exercise during CH compared to AH (50% MVC, $P = 0.018$; 75% MVC, $P = 0.030$) and N (50% MVC, $P = 0.034$). The MEP/ M_{\max} ratio increased for within contraction responses during CH (vs. AH 50 and 75% MVC; $P \leq 0.014$ and N 50% MVC; $P = 0.019$) (Table 2). The CSP did not differ between conditions pre-exercise (pooled average, 186 ± 47 ms; $P = 0.880$) or post-exercise (pooled average, 185 ± 50 ms; $P = 0.760$). Baseline cortical voluntary activation did not differ between conditions (AH, $93 \pm 5\%$; N, $97 \pm 3\%$; CH, $93 \pm 6\%$; $P = 0.310$) (Figure 5). Cortical voluntary activation was reduced post-exercise in AH ($\Delta 11\%$, $P = 0.014$), but not in N ($\Delta 4\%$, $P = 0.298$) or CH ($\Delta 6\%$, $P = 0.174$); the decrease in AH was greater compared to N ($P = 0.022$) (Figure 5).

Discussion

The aim of the present study was to assess corticospinal excitability and supraspinal fatigue after locomotor exercise in chronic hypoxia. The main finding was that exercise-induced supraspinal fatigue, as quantified via changes in cortical voluntary activation, was attenuated after two weeks of acclimatisation to high altitude whereas it was exacerbated in AH vs. N. Importantly, the diminished level of central fatigue in CH occurred in parallel with improvements in cerebral haemodynamics and arterial oxygenation (increased C_aO_2 and S_pO_2) brought about by the two weeks at altitude. Moreover, the attenuated development of central fatigue occurred in line with a substantial increase in corticospinal excitability. This latter finding suggests that a period of acclimatisation modifies the integrity of the corticospinal tract. We confirm our hypothesis that acclimatisation to altitude reduces the level of exercise-induced central fatigue and that this is attributable, at least in part, to an increased overall excitability of the brain to muscle pathway.

Supraspinal Fatigue

A key aim of the present study was to determine the effect of acclimatisation on the development of central fatigue assessed after exercise. We hypothesised that improvements in cerebral oxygenation known to occur after a prolonged stay at altitude would bring about positive modifications on the development of central fatigue. We show that the development of supraspinal fatigue during locomotor exercise is recovered after 2 weeks at high altitude and similar to that observed in normoxia. Thus, the adaptive processes that take place during acclimatisation to high altitude seemingly protect healthy humans against the development of supraspinal fatigue.

Corticomotor responses

The present study found no change in corticospinal excitability (Δ resting MEP) in AH, a finding which is in line with literature utilising varying severities of hypoxia ($F_{I}O_2 = 0.14 - 0.10$; resting $S_pO_2 = 93 - 74\%$) for as little as 10 min to 1 h (Goodall *et al.*, 2010, Rupp *et al.*, 2012, Millet *et al.*, 2012). However, Szubski *et al.* (2006) reported increased corticospinal excitability, expressed as a reduced rMT (not Δ MEP), after ~ 30 min of breathing hypoxic air ($F_{I}O_2 = 0.12$; resting $S_pO_2 = 75\%$). Moreover, the present study found a twofold increase in corticospinal excitability after 14 d acclimatisation to severe altitude (5,260 m, equivalent to $F_{I}O_2 = 0.105$; resting $S_pO_2 = 91 \pm 2\%$) with accompanying increases in the MEP/ M_{max} ratio, suggesting that the increases in MEP size were due to adaptive mechanisms within spinal and/or supraspinal sites. Similarly, Rupp *et al.* (2012) found a 26% increase in corticospinal excitability (Δ MEP amplitude) after 3 h of exposure to normobaric hypoxia ($F_{I}O_2 = 0.12$; resting $S_pO_2 = 86\%$), demonstrating a time-dependent, hypoxia-induced modification in the brain-to-muscle pathway. Thus, a prolonged stay at altitude modifies the integrity of the corticospinal pathway which may contribute to reduce the level of central fatigue; however, a duration-dependent adaptation cannot yet be established with certainty.

TMS over the motor cortex preferentially activates corticospinal neurons trans-synaptically through excitatory interneurons and corticocortical axons (Di Lazzaro *et al.*, 1998). The response to TMS critically depends on membrane excitability of motor cortical neurons and ion-channel function (Borojerdj *et al.*, 2001, Rothwell *et al.*, 1991). *In vitro* investigations using isolated cerebral neurons from rats demonstrate that ion-channel function is affected by O_2 availability and that neuronal hyper-excitability is the consequence of chronic hypoxia (Donnelly *et al.*, 1992). A heightened neural response is necessary to maintain membrane integrity and ionic homeostasis that occur from a period of insufficient metabolic activity (Nieber *et al.*, 1999). Thus, the twofold increase in MEP observed in the present study might be due to facilitated cortical neurons acting to restore the loss of neuronal activity associated with a prolonged exposure to altitude. Additionally, an increased level of muscle sympathetic nerve activity (peroneal microneurography) has been reported during a prolonged stay at the same altitude as in the present study (Hansen and Sander, 2003). That study showed a significant increase in muscle sympathetic nerve activity just 3 days after exposure to high altitude, suggesting that the prolonged stay induced a striking and long-lasting sympathetic over-activity. More recently, Buharin *et al.* (2013) found that a transient increase in sympathetic nerve activity (induced via lower body negative pressure) enhances corticospinal excitability as identified using TMS. The mechanism responsible for the increase in corticospinal excitability was postulated

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to be due to an elevated concentration of noradrenaline, a monoamine that is known to increase exponentially during sustained periods at altitudes exceeding 4,000 m (Cunningham *et al.*, 1965, Mazzeo *et al.*, 1994). Thus, the increased corticospinal excitability observed following 2 weeks of acclimatisation in the present study might be attributable, at least in part, to a heightened sympathetic nerve activity and associated increases in corticospinal excitability as well as hyper-excitable cerebral neurons. The increased corticospinal excitability in this investigation occurred in line with no symptoms of mountain sickness, a finding that opposes that of Miscio *et al.* (2009). Miscio *et al.* (2009) found that exposure to high altitude changes cortical excitability by affecting both inhibitory and excitatory circuits and that this is reflected in acute mountain sickness symptoms. This conclusion was based on a group of participants who resided at 4,554 m for only 3-5 days, a time frame in which acute mountain sickness is said to be most prominent (Hackett and Roach, 2001) and much shorter than the present study.

Despite substantial differences in end-exercise peripheral fatigue, CSP duration immediately after exercise (i.e., pre-to post-exercise change) was similar in all conditions. This suggests that locomotor exercise in N, AH and CH does not influence intracortical inhibition. These findings are in agreement with investigations using locomotor exercise in N and AH (Goodall *et al.*, 2012, Sidhu *et al.*, 2009b). However, Oliviero *et al.* (2002) reported decreased intracortical inhibition and CSP duration in chronic hypoxemic patients with COPD. These changes, mediated by cerebral GABA receptors, were reversed after 3-4 months of O₂ therapy, demonstrating that the changes were O₂ sensitive. However, factors other than chronic hypoxaemia might influence intracortical inhibition in patients with COPD making it difficult to quantify the influence that chronic hypoxaemia has on cortical inhibition.

On balance, we judge the increased corticospinal excitability in CH noted in the present study to be the result of adaptations in ion-channel function and elevations in circulating catecholamines serving to facilitate neurotransmission rather than mechanisms related to intracortical inhibition (Buharin *et al.*, 2013, Nieber *et al.*, 1999, Palange, 1998).

Hematological and cerebrovascular responses

Upon initial exposure to high altitude, acute hypoxia dilates cerebral arterioles thereby overriding the vasoconstrictive effect of hyperventilation-associated hypocapnia (Iwasaki *et al.*, 2011). During a prolonged stay at altitude, hypocapnia further develops and arterial hypoxaemia is ameliorated, as

reflected by increases in arterial [Hb], PO₂ and O₂ saturation (Figure 3). Furthermore, the increase in P_aO₂ and further decrease in P_aCO₂ with acclimatisation causes relative vasoconstriction reducing CBF down to SL values (Subudhi *et al.* 2013). We estimated an index of cerebral O₂ delivery using the product of MCA_v and C_aO₂. Our data demonstrate a reduced cerebral O₂ delivery index during exercise in AH compared to N; however, an improved cerebral O₂ delivery index was evident after two weeks of acclimatisation (Figure 3). The data in AH support a relationship between cerebral O₂ delivery and supraspinal fatigue (Goodall *et al.*, 2012). The calculation of C_aO₂ during exercise from resting [Hb] should be interpreted with caution as a hemoconcentration could have impacted this measure. At sea level, the hemoconcentration accompanying maximal exercise for approximately 10 min is counterbalanced by the concomitant exercise-induced arterial hypoxemia with the net effect of similar C_aO₂ at rest and during exercise (Amman *et al.*, 2006a). At altitude, despite significant hemoconcentration, C_aO₂ actually falls from rest to submaximal/maximal exercise by 10-25% (Calbet *et al.*, 2003). This would suggest that exercise C_aO₂ calculations, based on a resting C_aO₂ measure, might actually overestimate C_aO₂ measured during exercise at altitude. Furthermore, we assumed that MCA diameter would remain constant in hypoxia (Poulin and Robbins, 1996, Serrador *et al.*, 2000). While there is evidence of MCA dilatation at rest in hypoxia (Willie *et al.*, 2012, Wilson *et al.*, 2011), there is currently no evidence of MCA dilatation during intense exercise accompanied with substantial exercise-induced hyperventilation and associated hypocapnia. We acknowledge, however, that our measurements of blood velocity (rather than flow) must be interpreted with caution.

We found acclimatisation-induced increases in O₂ saturation and content (Figure 3). Furthermore, arterial O₂ tension increased from AH to CH (~39 mmHg to ~59 mmHg). Subudhi *et al.* (2013) has shown resting cerebral O₂ delivery to be maintained at levels observed in N during AH and CH, although it is presumed that the delivery of O₂ to the mitochondria within the parenchyma will be reduced because the driving gradient for diffusion from capillary to tissue is the PO₂ difference between capillary and tissue (Xu and Lamanna, 2006). The tissue PO₂ would be close to zero; thus, the driving force is essentially the P_aO₂. In the present study the P_aO₂ increased in line with acclimatisation, thereby improving the gradient for diffusion and perhaps restoring brain tissue O₂ tension to pre-hypoxic levels (Dunn *et al.*, 2000). Thus, we postulate that the lack of central fatigue in chronic hypoxia may be related to increases in brain tissue O₂ tension. However, the link between increases in P_aO₂ and C_aO₂ and the reduction in central fatigue that occurs after a period of acclimatisation warrants further investigation.

Technical Considerations

Exercising in a hypobaric environment was not feasible for the trials in AH. Thus, the two modes of hypoxia (normobaric [AH] vs. hypobaric [CH]) differed. The literature concerning the responses in normobaric and hypobaric hypoxia is equivocal and readers are directed elsewhere to a point-counterpoint debate (Girard *et al.*, 2012). Briefly, it was proposed that evidence is growing, suggestive that hypobaric hypoxia affects responses (ventilation, fluid balance, acute mountain sickness and performance) to a greater extent than normobaric hypoxia (Girard *et al.*, 2012). However, this argument was opposed by the fact that in terms of O₂ sensing, hypobaric hypoxia does not induce different responses compared to normobaric hypoxia (Mounier and Brugniaux, 2012). Moreover, it is unknown how any such differences which might exist between hypobaric and normobaric hypoxia may affect indices of exercise-induced fatigue. We set the F_IO₂ (0.105) at sea level to obtain the same P_IO₂ (~74 mmHg) that was expected at the subsequent altitude in Bolivia (5,260 m).

In line with other investigations that have measured exercise-induced fatigue of the knee extensors (Goodall *et al.*, 2012, Goodall *et al.*, 2010, Sidhu *et al.*, 2009b, Rossman *et al.*, 2013), measurements were made within 2.5 min after exercise termination. Corticospinal excitability associated with maximal single muscle contractions recovers within 1 min post-exercise (Taylor *et al.*, 1999). Thus, the present experimental design, utilising whole body exercise, might not have captured all elements of central fatigue. However, the methods and time to assess fatigue after exercise in all three conditions were identical and even though our measurements were made more than 1 min post-exercise, significant differences were observed, testifying to the strength of our data.

Conclusion

The novel finding was that supraspinal fatigue, present after exercise in acute hypoxia, was attenuated after a period of acclimatisation to high altitude. Importantly, the reduced development of central fatigue in chronic hypoxia occurred in parallel with an increase in the excitability of the brain to muscle pathway consequent to an increased cerebral O₂ delivery. The attenuated rate of development of central fatigue in chronic hypoxia might explain, at least in part, the improvements in locomotor exercise performance that are commonly observed after acclimatisation to high altitude.

Author Contributions

SG, RT, and MA contributed to conception and design of the experiments, data collection, data analysis, data interpretation, manuscript drafting and editorial process. ER contributed to conception and design of the experiments, data interpretation and manuscript revision. AL contributed to data collection. LR contributed to conception and design of the experiments, data interpretation, manuscript drafting and revision. AL, AS and RR conceived, designed and executed the AltitudeOmics study of which the present study was a part, and contributed to manuscript revision. AS also contributed to data collection and data interpretation. All authors approved the final version of the manuscript.

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Conflict of Interest

Nothing to declare

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Figure Legends

Figure 1. Mean area of motor evoked potentials (MEP) recorded from the vastus lateralis (VL) in response to stimulation over the motor cortex during varying contraction intensities pre- (○) and post-exercise (●) (mean for all conditions). The TMS responses were compared to the area of the maximal M-wave (M_{max}) evoked by peripheral stimulation of the femoral nerve. Data are means \pm SE for 7 participants.

Figure 2. Cerebral oxygenation at resting baseline, during the final 30 s of a 3 min warm up (28 W) and during the final 30 s of constant-load exercise (131 W) in normoxia (N; panel a), acute hypoxia (AH; panel b) and chronic hypoxia (CH; panel c). Data are means \pm SE for 7 participants. † $P < 0.05$ vs. respective baseline; ‡ $P < 0.05$ vs. respective warm up; * $P < 0.05$ vs. AH; # $P < 0.05$ vs. CH.

Resting baseline in AH denotes the value after 10 min wash in of the hypoxic gas. O₂Hb, oxygenated haemoglobin; HHb, deoxygenated haemoglobin and THb, total haemoglobin.

Figure 3. Arterial oxygen saturation (S_pO_2) (a), cerebral blood flow velocity (MCA_v) (b) and middle cerebral artery O₂ delivery index ($MCA_v \times C_aO_2$) during constant-load exercise (131 W) in normoxia (N), acute hypoxia (AH), and chronic hypoxia (CH). Values are plotted for the duration of the shortest trial (8 min) and extrapolated to the group mean exercise time (10.1 min). Data are means \pm SE for 7 participants. † P < 0.05 vs. rest; * P < 0.05 vs. N; # P < 0.05 vs. CH.

Figure 4. Representative MEPs evoked during knee extensor contractions at 50% MVC before exercise in each condition. Traces are shown from a representative participant in each condition; 8 stimuli were delivered from which an average value was obtained. Note the increase in MEP amplitude (corticospinal excitability) after acclimatisation.

Figure 5. Cortical voluntary activation measured before (open bars) and immediately after (<2.5 min; closed bars) constant-load exercise (131 W) in normoxia (N), acute hypoxia (AH), and chronic hypoxia (CH). * P < 0.05 pre- vs. post-exercise.

Table 1. Cardiorespiratory and perceptual responses at rest and during the final minute of constant-load cycling (131 W) in normoxia, acute hypoxia and chronic hypoxia.

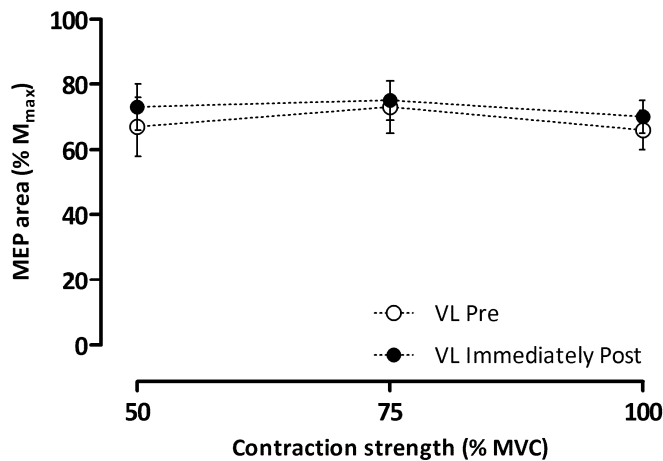
| | | Normoxia | Acute Hypoxia | Chronic Hypoxia |
|--------------------------------------|-----------|------------------|-------------------|------------------|
| HR (beats min ⁻¹) | Rest | 81 \pm 7† | 90 \pm 9 | 104 \pm 16 |
| | Final Min | 150 \pm 16* | 173 \pm 14 | 167 \pm 16 |
| \dot{V}_E (l min ⁻¹) | Rest | 14.3 \pm 2.4 | 20.0 \pm 2.6 | 24.5 \pm 5.4 |
| | Final Min | 60.0 \pm 9.6*† | 108.8 \pm 24.7† | 128.5 \pm 30.0 |
| f_R (breaths min ⁻¹) | Rest | 15.6 \pm 3.6 | 17.5 \pm 4.5 | 13.0 \pm 3.4 |
| | Final Min | 31.4 \pm 4.9*† | 51.6 \pm 8.7† | 54.8 \pm 9.9 |
| V_T (l) | Rest | 1.07 \pm 0.37 | 1.30 \pm 0.34 | 1.47 \pm 0.63 |
| | Final Min | 2.00 \pm 0.45 | 2.07 \pm 0.43 | 2.41 \pm 0.58 |
| $\dot{V}O_2$ (l min ⁻¹) | Rest | 0.49 \pm 0.10 | 0.45 \pm 0.08 | 0.45 \pm 0.12 |
| | Final Min | 2.45 \pm 0.51 | 2.34 \pm 0.58 | 2.07 \pm 0.50 |
| $\dot{V}CO_2$ (l min ⁻¹) | Rest | 0.44 \pm 0.09 | 0.55 \pm 0.09 | 0.39 \pm 0.08 |
| | Final Min | 2.32 \pm 0.51 | 2.69 \pm 0.62† | 1.94 \pm 0.50 |
| $\dot{V}_E / \dot{V}O_2$ | Rest | 30.7 \pm 2.7*† | 47.4 \pm 6.5† | 55.9 \pm 14.9 |
| | Final Min | 25.2 \pm 2.4*† | 51.2 \pm 15.0† | 62.9 \pm 9.2 |
| $\dot{V}_E / \dot{V}CO_2$ | Rest | 33.9 \pm 2.7† | 37.9 \pm 6.5† | 63.4 \pm 6.8 |
| | Final Min | 26.2 \pm 2.6*† | 41.7 \pm 6.9† | 67.1 \pm 9.1 |
| RPE, dyspnoea | Rest | 7.0 \pm 0.0 | 7.3 \pm 0.5 | 7.1 \pm 0.4 |
| | Final Min | 11.4 \pm 2.4*† | 19.4 \pm 0.8 | 19.1 \pm 0.7 |
| RPE, limb | Rest | 7.1 \pm 0.4 | 7.1 \pm 0.4 | 7.0 \pm 0.0 |
| | Final Min | 12.3 \pm 3.3* | 19.9 \pm 0.4 | 17.6 \pm 1.7 |

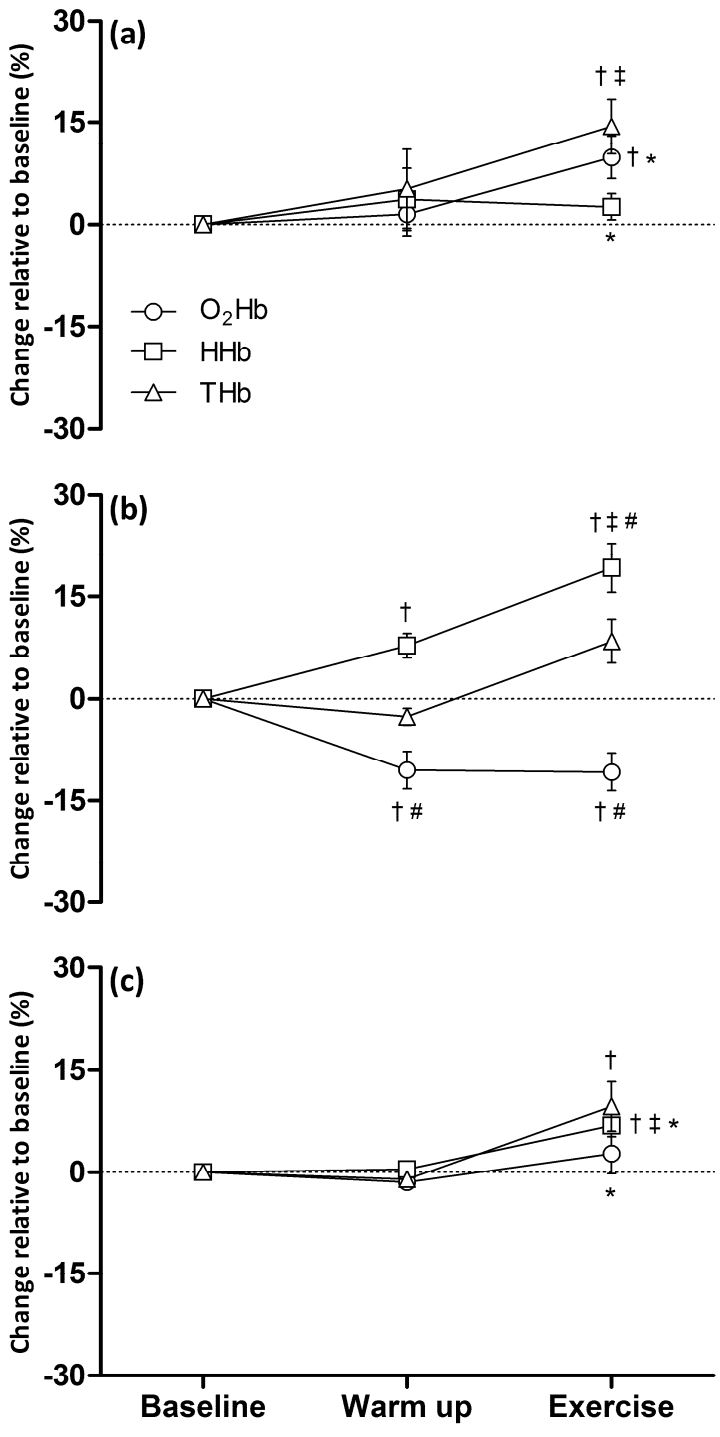
Values are means \pm SD for 7 participants. Resting values were measured during the 5th minute of breathing the test gas mixture. HR, heart rate; \dot{V}_E , minute ventilation; f_R , respiratory frequency; V_T , tidal volume; $\dot{V}O_2$, oxygen uptake; $\dot{V}CO_2$, carbon dioxide output; RPE, ratings of perceived exertion. * P < 0.05 vs. acute hypoxia; † P < 0.05 vs. chronic hypoxia.

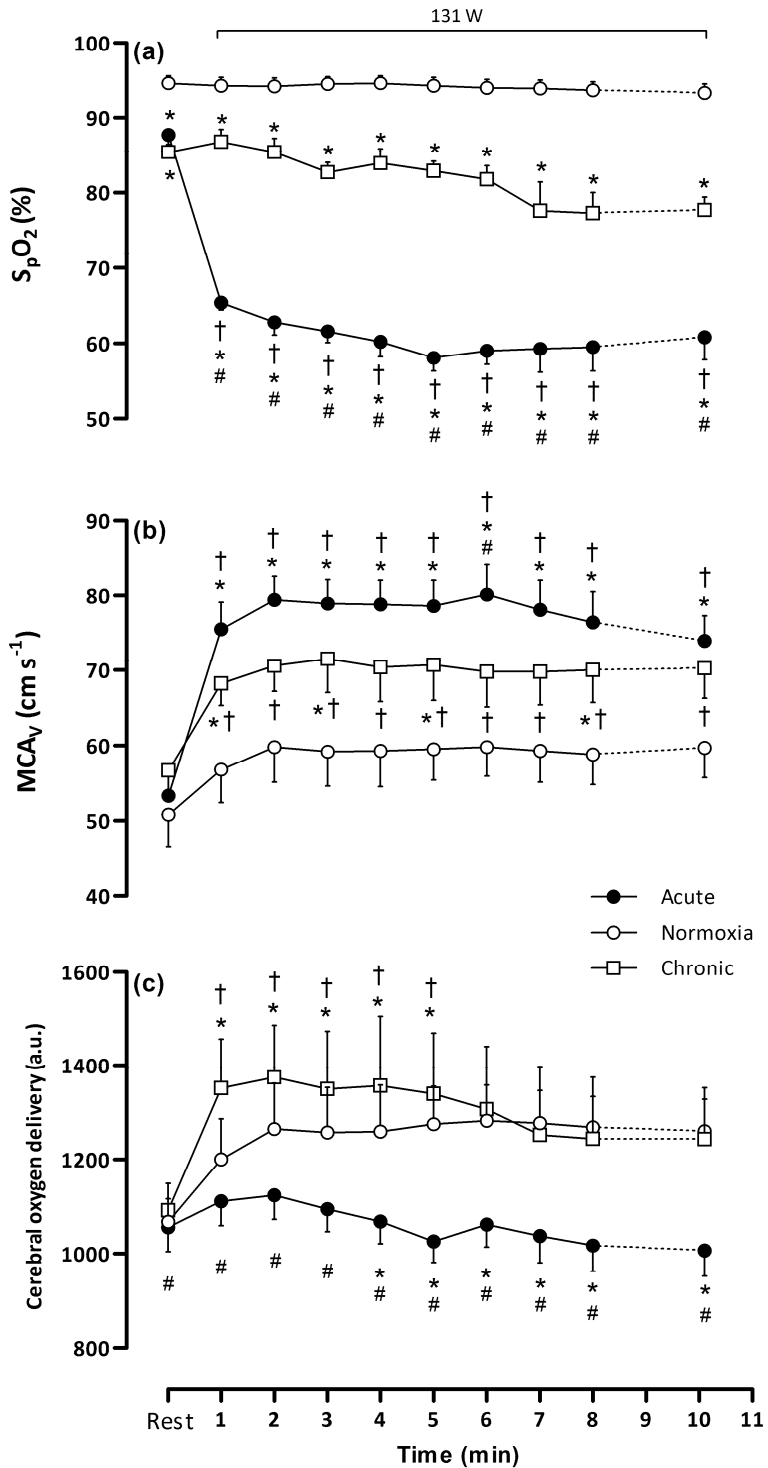
Table 2. Peripheral and central measures of excitability assessed before and after constant-load cycling (131 W) in normoxia, acute hypoxia and chronic hypoxia.

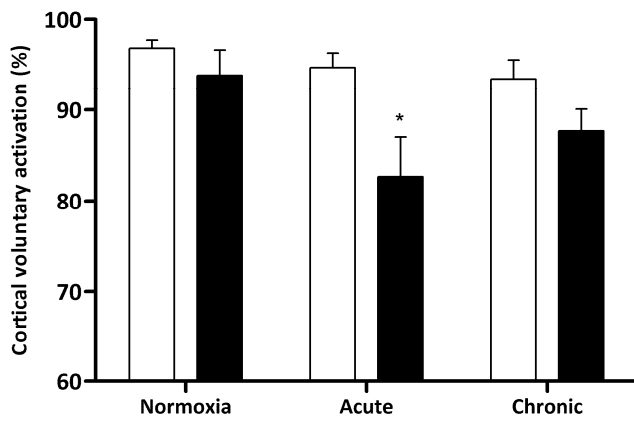
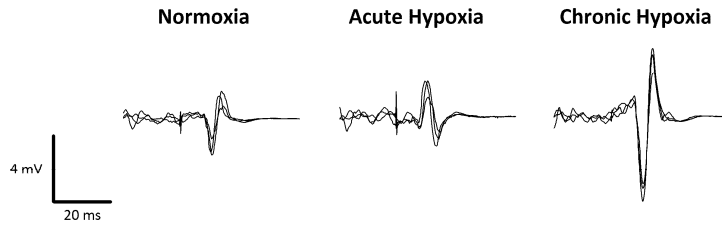
| | | Normoxia | Acute Hypoxia | Chronic Hypoxia |
|--------------------------------------|------|--------------------------|--------------------------|--------------------------|
| Rest | | | | |
| M _{max} amplitude (mV) | Pre | 6.9 ± 2.0 [†] | 8.6 ± 3.7 [†] | 14.9 ± 8.3 |
| | Post | 6.7 ± 1.7 | 9.0 ± 4.1 | 14.0 ± 8.2 |
| MEP amplitude (mV) | Pre | 0.19 ± 0.12 [†] | 0.19 ± 0.11 [†] | 0.41 ± 0.28 |
| | Post | 0.11 ± 0.06 | 0.11 ± 0.10 | 0.21 ± 0.18 [#] |
| MEP/M _{max} (%) | Pre | 2.6 ± 1.3 | 2.7 ± 1.9 | 4.1 ± 4.2 |
| | Post | 1.8 ± 1.2 | 1.5 ± 1.3 | 2.6 ± 3.4 |
| Within contraction | | | | |
| M _{max} amplitude 100% (mV) | Pre | 8.9 ± 1.7 | 9.9 ± 3.2 | 13.0 ± 6.1 |
| | Post | 9.0 ± 1.9 | 10.0 ± 3.3 | 11.9 ± 5.4 |
| MEP amplitude 100% (mV) | Pre | 3.8 ± 1.5 | 3.1 ± 1.0 [†] | 7.1 ± 4.7 |
| | Post | 4.0 ± 2.7 | 3.2 ± 1.0 | 6.5 ± 4.4 |
| MEP amplitude 75% (mV) | Pre | 3.9 ± 1.5 [†] | 2.9 ± 1.4 [†] | 7.6 ± 4.9 |
| | Post | 4.3 ± 2.6 | 3.3 ± 1.2 [†] | 6.9 ± 3.9 |
| MEP amplitude 50% (mV) | Pre | 2.54 ± 0.87 [†] | 2.16 ± 0.52 [†] | 6.5 ± 4.8 |
| | Post | 2.99 ± 2.01 [†] | 2.56 ± 0.95 [†] | 6.4 ± 4.5 |
| MEP/M _{max} (%) 100% MVC | Pre | 35 ± 17 | 33 ± 14 | 52 ± 17 |
| | Post | 39 ± 20 | 37 ± 15 | 52 ± 19 |
| MEP/M _{max} (%) 75% MVC | Pre | 40 ± 15 | 34 ± 19 [†] | 58 ± 18 |
| | Post | 42 ± 17 | 38 ± 18 [†] | 57 ± 13 |
| MEP/M _{max} (%) 50% MVC | Pre | 28 ± 14 [†] | 26 ± 10 [†] | 50 ± 21 |
| | Post | 30 ± 15 [†] | 31 ± 17 [†] | 54 ± 23 |
| CSP (ms) | Pre | 198 ± 58 | 174 ± 46 | 186 ± 36 |
| | Post | 188 ± 64 | 171 ± 35 | 196 ± 51 |

Values are means ± SD for 7 participants. M_{max}, maximal motor response; MEP, motor evoked potential; CSP, cortical silent period. [†] P < 0.05 vs. chronic hypoxia; [#] P < 0.05 vs. Pre.









AltitudeOmics: Effect of ascent and acclimatization to 5260 m on regional cerebral oxygen delivery

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What is the central question of this study?

Hypoxia associated with ascent to high altitude may threaten cerebral oxygen delivery.

We sought to determine if there are regional changes in the distribution of cerebral blood flow that might favor oxygen delivery to areas associated with basic homeostatic functions to promote survival in this extreme environment.

What is the main finding and its importance?

We show evidence of a “brain sparing” effect during acute exposure to high altitude, in which there is a slight increase in relative oxygen delivery to the posterior cerebral circulation. This may serve to support basic regulatory functions associated with the brain stem and hypothalamus.

Abstract

Cerebral hypoxemia associated with rapid ascent to high altitude can be life threatening; yet, with proper acclimatization, cerebral function can be maintained well enough for humans to thrive. METHODS: We investigated adjustments in global and regional cerebral oxygen delivery (DO_2) as 21 healthy volunteers rapidly ascended and acclimatized to 5260m. Ultrasound indices of cerebral blood flow (CBF) in internal carotid and vertebral arteries were measured at sea level (SL), upon arrival at 5260m (ALT1; $P_{bar} = 409\text{mmHg}$), and after 16 days of acclimatization (ALT16). Cerebral DO_2 was calculated as the product of arterial oxygen content (CaO_2) and flow in each respective artery and summed to estimate global CBF. Vascular resistances were calculated as the quotient of mean arterial pressure and respective flows. RESULTS: Global CBF increased $\sim 70\%$ upon arrival at ALT1 ($P < 0.001$) and returned to SL values at ALT16 as a result of changes in cerebral vascular resistance. A reciprocal pattern in CaO_2 maintained global cerebral DO_2 across acclimatization, although DO_2 to the posterior cerebral circulation was increased by $\sim 25\%$ at ALT1 ($P = 0.032$). CONCLUSIONS: Cerebral DO_2 is well maintained upon acute exposure and acclimatization to hypoxia, particularly in the posterior and inferior regions of the brain associated with vital homeostatic functions. This tight regulation of cerebral DO_2 was achieved through integrated adjustments in local vascular resistances to alter cerebral perfusion during both acute and chronic exposure to hypoxia.

Introduction

Although the brain represents only about 2% of body weight, it is a highly metabolic tissue that receives ~15% of cardiac output and accounts for ~20% of total body oxygen consumption at rest (Wade & Bishop, 1962). Maintenance of cerebral oxygen delivery (DO_2) is essential for vital cerebral functions associated with homeostasis. In the face of severe hypoxemia, such as experienced during rapid ascent to extreme altitudes ($> 8,000$ m), reduction in cerebral DO_2 results in loss of consciousness within seconds (Luft *et al.*, 1951; Luft & Noell, 1956) and death within minutes (Bert, 1943). However, with staged acclimatization to progressively higher elevations, cerebral DO_2 can be maintained well enough for humans to reach the summit of Mount Everest (8,848 m) without supplemental oxygen. The mechanisms responsible for this remarkable plasticity in cerebral DO_2 are complex and not completely understood.

Cerebral DO_2 is the product of cerebral blood flow (CBF) and arterial oxygen content (CaO_2). It is well established that CBF rises upon acute exposure to high altitude and returns to near sea-level values with acclimatization (Severinghaus *et al.*, 1966; Huang *et al.*, 1987; Jensen *et al.*, 1990), while CaO_2 decreases in acute hypoxia and returns to sea-level values with acclimatization. These opposing CBF and CaO_2 responses to altitude appear to offset one another and maintain cerebral DO_2 across acclimatization (Severinghaus *et al.*, 1966; Wolff *et al.*, 2002). The pattern of CBF

change in response to hypoxia has been attributed to the relative balance of hypoxic vasodilation and hypocapnic vasoconstriction in the brain (Xu & Lamanna, 2006; Brugniaux *et al.*, 2007). During acute, severe hypoxia, vasodilation typically exceeds vasoconstriction, resulting in greater CBF (Mardimae *et al.*, 2012; Willie *et al.*, 2012). With acclimatization, increased ventilatory drive reduces PaCO₂ and improves PaO₂, tipping the balance in favor of vasoconstriction and restoring CBF to pre-exposure values. Changes in the PaO₂/PaCO₂ ratio have been shown to account for ~40% of the variation in global CBF over acclimatization (Lucas *et al.*, 2011), with other biochemical (e.g. pH, HCO₃⁻, nitric oxide) and hematological (e.g. hemoglobin, hematocrit, blood viscosity) factors presumably accounting for the rest of the response (Todd *et al.*, 1994; Tomiyama *et al.*, 1999; Severinghaus, 2001) to maintain global cerebral DO₂.

Recent data demonstrate that acute normobaric hypoxia (i.e. breathing hypoxic gas) affects the regional distribution of CBF within the brain. Data from positron emission tomography (PET) studies show greater perfusion of the brain stem, hypothalamus, thalamus and cerebellum during acute hypoxia, with (Binks *et al.*, 2008) or without (Buck *et al.*, 1998) controlled levels of PaCO₂. Regional differences in cerebrovascular reactivity to O₂ and CO₂ have been postulated to control the distribution of CBF. Vascular Doppler studies of the major tributary vessels of the brain suggest that a greater percentage of blood flow may be directed towards the posterior cerebral circulation, including the brain stem, in response to controlled levels of hypoxia and hypocapnia (Sato *et al.*, 2012). From a teleological perspective,

this could help preserve vital homeostatic functions at the expense of higher cognitive processing; however, it is unclear whether regional distribution of CBF is similarly affected in hypobaric hypoxia (i.e. high altitude) or if it changes with acclimatization, as not all studies report significant regional differences (Huang *et al.*, 1987; Willie *et al.*, 2012; Willie *et al.*, 2013).

Despite the importance of O₂ supply for cerebral function, longitudinal studies of cerebral DO₂ at high altitude are sparse. In a secondary analysis of data from Severinghaus *et al.*'s original study of CBF at high altitude, global cerebral DO₂ in four subjects appeared stable and in excess of oxygen demand after 6-12 hours and 3-5 days of exposure to 3,810m (Severinghaus, 2001; Wolff *et al.*, 2002). Using similar methodology (Kety-Schmidt technique), no differences were found in global cerebral DO₂ measured after 5 weeks at 5,260 m and return to sea level (Moller *et al.*, 2002). Unfortunately, these two studies were based on a limited number of observations, which makes it difficult to detect small differences if they existed (type II error), and utilized methodology that can only measure global cerebral DO₂. A more recent MRI study with a larger sample size reported a tendency towards elevation of cerebral DO₂ after subjects returned from 2 days at 3,800 m (Smith *et al.*, 2013), but no measurements of regional cerebral DO₂ were made. Based the limited data to date, it is uncertain if global or regional cerebral DO₂ varies over time at high altitude.

In this study we used vascular Doppler technology in conjunction with arterial blood sampling to allow us to quantify global and regional changes in CBF and cerebral DO_2 in the field as healthy people rapidly ascended and acclimatized to high altitude (5,260 m). We tested the hypothesis that upon acute exposure cerebral DO_2 would be maintained to regions of the brain associated with homeostasis at the expense of other tissues, but that these changes would normalize with acclimatization.

Methods

Subject recruitment and screening

This study was conducted as part of the AltitudeOmics project, for which a detailed description of the protocol is published elsewhere (Subudhi *et al.*, In Review). Briefly, following institutional ethics approval from the Universities of Colorado and Oregon and the US Department of Defense Human Research Protection Office, young, healthy sea-level residents were recruited from the greater Eugene, Oregon area (elevation 128 m). Potential subjects were screened to exclude anyone who was born or had lived at altitudes $>1,500$ m for more than one year or had traveled to altitudes $>1,000$ m in the past 3 months. After obtaining written consent, physical exams and the Army Physical Fitness Test (push ups, sit ups and 3.2 km run) were performed to verify health and fitness status.

Study overview

To evaluate effects of altitude acclimatization on cerebrovascular hemodynamics, subjects were studied on 3 occasions: 1) at sea level (SL, 130 m), 2) upon acute exposure to 5,260 m (ALT1), and 3) after 16 days of acclimatization (ALT16). Specifically, ~4 weeks following SL measurements in Eugene, Oregon, subjects were flown to La Paz, Bolivia. They spent two nights at low altitude (Coroico, Bolivia, 1,525 m) before being driven to the Chacaltaya Research Station at 5,260 m while breathing supplemental oxygen. Acute responses to high altitude were assessed 2 to 4 hours after arrival and cessation of supplemental oxygen (ALT1). Subjects acclimatized to altitudes ranging from 3,800 to 5,260 m over the next 15 days, with a majority of the time (75%) spent at 5,250 m. Measurements were repeated on ALT16.

Instrumentation

Subjects were studied in an upright, seated position with feet on the floor. Arterial blood pressure (ABP) was monitored via a fluid filled pressure transducer (Utah Medical, Salt Lake City, UT, USA) positioned at heart level and attached to a 22-gauge catheter in a radial artery. Blood flow velocity in the left middle cerebral artery was measured by transcranial Doppler (MCA_{velocity}: 2MHz probe, Spencer Technologies, Seattle, WA, USA, affixed to a custom-made headset) at depths ranging from 43 to 54 mm. Signal quality was optimized and an M-mode screen shot was recorded to facilitate subsequent probe placements. Arterial saturation was measured on the right side of the forehead by pulse oximetry (Nellcor N-200,

Mansfield, MA, USA). Limb lead electrodes were used to measure ECG (ADInstruments BioAmp, Colorado Springs, CO, USA and Sonosite Micromaxx, Bothell, WA, USA). Metabolic variables, including expired ventilation and gas concentrations were assessed via breath-by-breath (Medgraphics PFX, St. Paul, MN, USA and Vacumed UVM, Ventura, CA, USA) and mixing chamber (Oxigraf O₂cap, Mountain View, CA, USA) systems, calibrated with the same 3-L syringe and known concentrations of O₂ and CO₂ prior to each test. Additionally, core temperature was monitored by telemetry pill (CorTemp HQInc., Palmetto, FL, USA) Analog data were sampled and recorded at 200Hz (ADInstruments Powerlab 16/30, Colorado Springs, CO, USA).

Cerebral Blood Flow

After verification of signal quality, resting data were recorded for 10 min while subjects breathed room air. At 6 min, 2 ml of arterial blood was drawn anaerobically for blood gas analysis (described below). During the last 4 min of the resting period, diameter and blood flow velocity in the left internal carotid (ICA: 1.5 cm distal to the carotid bifurcation) and vertebral arteries (VA: between spinous processes of C4 and C5) were recorded over a minimum of 5 cardiac cycles by a registered diagnostic sonographer (SonoSite Micromaxx L25 probe, Bothell, WA, USA). Briefly, vessel diameter from a longitudinal view was identified and measured with digital calipers in synchronization with the ECG tracing to identify systole and diastole. Velocity was measured in the center of the vessel with an insonation angle < 60 degrees and a sample volume maximized for vessel diameter. The peak velocity tracing across cardiac cycles was used for calculation of mean velocity (time

averaged peak) and volumetric flow. This procedure was used to verify accurate tracing of the spectral envelop during data collection and results in higher values than the time averaged mean method (Schoning *et al.*, 1994). All data were downloaded in DICOM format for verification of measurements offline (Sante DICOM Editor, Athens, Greece).

Regional blood flow (ml/min) in the ICA and VA (ICA_{flow} and VA_{flow}) was determined using standard, validated ultrasound techniques (Hoskins, 2008), where:

$$X_{Flow} = \pi * (\text{diameter in cm}/2)^2 * \text{time averaged peak velocity in cm/s} * 60 \text{ s.}$$

Average coefficients of variation determined from three repeated measurements of ICA and VA flow measurements in 7 subjects at SL were $4.0 \pm 2.6\%$ and $4.0 \pm 2.1\%$, respectively. Global CBF (gCBF) was estimated assuming symmetrical bilateral flow in the major tributary arteries of the brain (Ogoh *et al.*, 2013; Willie *et al.*, 2013) as:

$$gCBF = (ICA_{flow} + VA_{flow}) * 2.$$

Regional and global measurements of CBF were also expressed relative to estimates of cardiac output (%Q) derived from simultaneous intra-arterial blood pressure tracings (Bogert *et al.*, 2010). Cerebral vascular resistance index (CVRi) was calculated as:

$$CVRi = \text{mean ABP}/X_{flow}$$

Cerebral Oxygen Delivery

Arterial blood was immediately analyzed for PaO₂, PaCO₂ (Siemens RAPIDLab 248, Erlangen, Germany), [Hb], SaO₂ (Radiometer OSM3, Copenhagen, Denmark) and Hct (M24 Centrifuge, LW Scientific, Lawrenceville, GA, USA). Blood gases were

temperature corrected (Kelman & Nunn, 1966; Severinghaus, 1966). CaO_2 (vol%) was calculated as:

$$\text{CaO}_2 = 1.39 \times [\text{Hb}] + \text{PaO}_2 * 0.003$$

Regional and global cerebral DO_2 were calculated as the products of CaO_2 and ICA_{flow} , VA_{flow} , and gCBF .

Data Analysis

After verification of normality, mixed repeated measures ANOVA's were used to analyze the interaction of time by sex for each variable of interest ($\alpha = 0.05$).

Subsequent estimation-maximization and multiple-imputation (5 trials) analyses verified negligible effects of missing values (SPSS 20, IBM, Chicago, IL, USA). Paired t-tests (without imputation of missing values) were used for *post hoc* comparisons with the Holm procedure to control for Type I error. A priori power calculations ($\alpha = 0.05$, $\beta = 0.20$) were integrated into the study design to limit Type II error. Pearson product moment correlations were used to describe shared variance between variables. Data are presented as mean \pm SD.

Based on the hypothesis that increased CBF may play a role in the pathogenesis of acute mountain sickness [AMS (Jensen *et al.*, 1990; Baumgartner *et al.*, 1994; Baumgartner *et al.*, 1999)], a secondary analysis was performed to evaluate potential relationships (Spearman correlations) between changes in CBF and DO_2 with the severity of Lake Louise Questionnaire (LLQ) symptoms scores reported in these subjects on ALT1 (Subudhi *et al.* – in review). Paired t-tests were used to

evaluate differences in CBF and DO₂ between those with severe AMS (LLQ ≥ 6 including headache) and those remaining healthy.

Results

Subject Characteristics

Detailed baseline characteristics of the 21 (12 males and 9 females; 21 ± 1 years old) subjects participating in AltitudeOmics are presented elsewhere (Subudhi *et al.*, In Review). Males exhibited greater [Hb], CaO₂ and DO₂ than females over the course of the study (all P < 0.05), but since no interactions in CBF or DO₂ were detected across acclimatization, combined data are presented below.

Cerebral Blood Flow and Oxygen Delivery

Acute exposure to 5,260 m (Pbar = 408 ± 1 mmHg) decreased PaO₂, SaO₂ and CaO₂ by 66.1 ± 5.4 mmHg, 22 ± 6%, and 4.1 ± 1.2 ml/dl, respectively (all P < 0.001; Table 1). This severe degree of hypoxia increased heart rate 14 ± 11 bpm (P < 0.001) without affecting mean ABP (P = 0.380). CBF increased 74 ± 81% in the ICA (P = 0.018), 59 ± 54% in the VA (P = 0.001), and 69 ± 57% globally (P = 0.003). Respective CVR_i values fell (all P < 0.001; Table 2), allowing a larger percentage of cardiac output to perfuse the brain (P = 0.010). Increased ICA_{flow} was characterized by increased ICA velocity (P = 0.004) without a change in diameter (P = 0.068), while increased VA_{flow} was explained by an increase in VA diameter (P = 0.005) without a change in velocity (P = 0.120). MCA_{velocity} was unchanged (P = 0.953). Increased gCBF offset the decrease in CaO₂ to maintain global cerebral DO₂ (Figure 1), although a

small increase in VA DO₂ was observed (P=0.039, Figure 2). Observed changes in measures of regional and global CBF and DO₂ were not correlated with LLQ scores of AMS (r = -0.07 to -0.23, P = 0.38 to 0.78), nor were they different between those reporting severe AMS and those remaining healthy (P = 0.57 to 0.97).

Following acclimatization, a $32 \pm 36\%$ rise in ventilation was accompanied by a 5.5 ± 2.7 mmHg decrease in PaCO₂ and 9.2 ± 4.1 mmHg increase in PaO₂ (ALT1 vs. ALT16; all P < 0.001). SaO₂ and [Hb] rose $6 \pm 5\%$ and 1.8 ± 0.9 g/dL, respectively, improving CaO₂ by 3.1 ± 1.2 ml/dl (all P < 0.001; Table 1). ABP was unaffected by acclimatization (ALT1 vs. ALT16; P=0.211). ICA_{flow}, VA_{flow} and gCBF returned to SL values (SL vs. ALT16; P = 0.810, 0.977, 0.620, respectively; Table 2). Respective CVR_i values increased as both ICA and VA diameters decreased from ALT1 to ALT16 (all P < 0.020) and restored the relative distribution of cardiac output back to SL values (SL vs. ALT16; P = 0.121). Cerebral DO₂ fell from ALT1 to ALT16 (ICA DO₂ P = 0.028, VA DO₂ P = 0.020, global DO₂ P = 0.011) as the reductions in CBF outweighed the increase in CaO₂ (Figure 1); however, neither global nor regional cerebral DO₂ values fell below that measured at SL (all P > 0.420; Figures 1 & 2).

Discussion

This is the first study to assess regional cerebral oxygen delivery in the field over a period of acclimatization to high altitude. Our findings confirm that global cerebral DO₂ was preserved across acclimatization through a changing balance between CBF and CaO₂, but there was slight increase in relative DO₂ to the posterior cerebral

circulation during acute exposure. Although changes in CBF and DO_2 were not associated with the incidence or severity of AMS, regional regulation of CBF may serve to support vital homeostatic cerebral functions in hypoxia.

Preservation of Cerebral Oxygen Delivery

The increase in CBF upon arrival at high altitude and decrease back to sea level values with acclimatization was opposed by changes in CaO_2 (Figure 2). These responses preserved cerebral DO_2 close to sea level values and affirm that components of CaO_2 (PaO_2 , SaO_2 , [Hb]) outweigh the influence PaCO_2 in regulating CBF in severe hypoxia. Increased CBF upon arrival at high altitude resulted from reduced cerebral vascular resistance rather than increased blood pressure (Tables 1&2). Although reduction in vascular resistance is commonly attributed to dilation of pial and parenchymal arterioles in the brain (Fog, 1938), we observed increased diameter of larger tributary arteries, supporting a global vascular response to this degree of hypoxia (Heistad *et al.*, 1978; Faraci & Heistad, 1990; Willie *et al.*, 2012). Mechanisms governing hypoxic vasodilation are complex, involving local (e.g. astrocyte regulation, nitric oxide) and diffuse (e.g. central chemoreception, autonomic nervous system) mechanisms, but all stem from a reduction in PaO_2 (Severinghaus, 2001; Xu & Lamanna, 2006). When PaO_2 is above 60 mmHg, little vasodilation is evident (Mardimae *et al.*, 2012; Willie *et al.*, 2012). Below this threshold, the degree of vasodilation increases exponentially and outweighs the degree of hypocapnic vasoconstriction (Mardimae *et al.*, 2012; Willie *et al.*, 2012) - presumably to provide greater blood flow in a time of need. While the correlation between changes in gCBF and CaO_2 was not significant, the change in CaO_2 from SL

to ALT1 was similar among all subjects and may not have afforded an appropriate range of values to detect the relation that has previously been shown with progressive hemodilution (Korosue & Heros, 1992). Qualitatively, the ~70% increase in gCBF was within the expected range during acute hypocapnic hypoxia (Severinghaus, 1966; Jensen *et al.*, 1990; Severinghaus, 2001; Brugniaux *et al.*, 2007) and proportional to the ~60% reduction in PaO₂ that was responsible for the reduction in CaO₂. This reciprocal relationship, whether evolved or serendipitous, is advantageous for survival in these extreme conditions as it mitigates negative consequences of cerebral hypoxemia.

Although increased CBF has been suggested to play a role in the pathogenesis of AMS (Baumgartner *et al.*, 1994), our results were more similar to those refuting the hypothesis (Jensen *et al.*, 1990; Baumgartner *et al.*, 1999). Regional and global CBF and DO₂ measurements were not correlated with AMS symptoms scores and did not differentiate between those with severe AMS and those who remained healthy after rapid ascent to high altitude. Nonetheless, our data should be interpreted with caution since it is possible that increased CBF contributes to the development of AMS when other, yet to be described, factors are present.

Increased PO₂ and decreased PCO₂ after 16 days at high altitude are hallmarks of ventilatory acclimatization that are addressed elsewhere (Fan *et al.* in review). As a result, PaO₂-mediated dilation was reduced and PaCO₂-mediated vasoconstriction was increased, thereby lowering CBF. Assuming a cerebral O₂ reactivity of 3% CBF /

% SaO₂ and a CO₂ reactivity of 4% CBF / mmHg CO₂ from a previous duplex ultrasound study (Willie *et al.*, 2012), we could account for the entire decrease in gCBF across acclimatization. Specifically, the 5% increase in SaO₂ could be expected to reduce CBF by ~15% and the 5.5 mmHg decrease in PaCO₂ could be expected to reduce CBF by ~22%, thus accounting for the 36% decrease in gCBF we observed from ALT1 to ALT16 (Table 2). We acknowledge that increased cerebrovascular CO₂ reactivity with acclimatization in our subjects (Fan *et al.* in review) may account for an even greater proportion of the net effect on CBF at ALT16. Also, the relative influence of other hematological factors, such as increased hematocrit and blood viscosity (Sorensen *et al.*, 1974; Todd *et al.*, 1994; Tomiyama *et al.*, 1999) from erythropoiesis and plasma volume contraction, may have contributed to the reduction of CBF across acclimatization (data to be presented elsewhere). Yet our data suggest that the inherent vascular reactivities to O₂ and CO₂ are sufficient to maintain tight control over cerebral DO₂ in hypoxia. Consistent delivery of oxygen may help offset the decreased PO₂ gradient (plasma to mitochondria) and support the cerebral metabolic demand for oxygen at this altitude (Severinghaus *et al.*, 1966; Moller *et al.*, 2002) to preserve cerebral function. Together, our data demonstrate that integrated mechanisms controlling cerebral blood flow are well suited to preserve global cerebral oxygen delivery at 5,260 m.

Regional Cerebral Oxygen Delivery

We observed a small increase in DO₂ through the posterior cerebral circulation upon arrival at high altitude (Table 2) that dissipated with acclimatization. The acute increase in DO₂ was characterized by an increase in VA diameter and supports

recent findings of greater VA (vs. ICA) vasoreactivity during acute hypoxia (Willie *et al.*, 2012; Ogoh *et al.*, 2013). Of note, Ogoh *et al.* (Ogoh *et al.*, 2013), showed that acute hypoxia (~15 min) increased VA, but not ICA, blood flow. Since the areas perfused by the VA include the brainstem, and posterior aspects of the thalamus and hypothalamus, increased blood flow and DO_2 to these regions during acute hypoxia (Buck *et al.*, 1998; Binks *et al.*, 2008) may be seen as necessary to maintain vital homeostatic functions (Sheldon *et al.*, 1979; Bilger & Nehlig, 1993). Since increased cardiorespiratory drive with acclimatization was not associated with a continued elevation of VA DO_2 , we speculate that the increased VA DO_2 during acute hypoxia was protective, to defend against a potential threat in oxygen supply, rather than to merely support neuronal metabolic activity associated with heightened autonomic activity (i.e. neurovascular coupling). Although such hypothetical explanations for regional differences in the regulation of CBF and DO_2 are intriguing, our results must be interpreted with caution since measured differences were small and are not consistently reported in the literature (Huang *et al.*, 1987; Willie *et al.*, 2013). Future studies with more focal measurements of DO_2 (e.g. PET and MRI) and neuronal activity in key regulatory regions of the brain, as well as measurements of neurovascular coupling (as an index of neuronal plasticity) during acute and prolonged hypoxia are needed to yield further insight into this question.

Brain Sparing

Reduced cerebral vascular resistance associated with vasodilation upon arrival at altitude can explain the proportional increase in CBF and greater allocation of cardiac output. This effect could be magnified if there is net constriction in other

vascular beds at rest. Previous studies have shown that superior mesenteric and renal artery blood flow decrease in acute hypoxia and could allow for greater perfusion of the brain (Greene & Roach, 2004). With acclimatization, cerebral vascular resistance and blood flow returned to sea-level values. These results are similar to fetal 'brain sparing' effects (Campbell *et al.*, 1967; Peeters *et al.*, 1979; Sheldon *et al.*, 1979) that are presumed to preserve vital homeostasis during hypoxia in utero (Pearce, 2006; Salihagic-Kadic *et al.*, 2006). Similar effects have also been shown in newborn dogs (Cavazzuti & Duffy, 1982), piglets (Goplerud *et al.*, 1989), and premature infants (Daven *et al.*, 1983). The largest response to hypoxia tends to occur in the brainstem during the early postnatal period and decreases with age (Bilger & Nehlig, 1993). We are the first to demonstrate that such a 'brain sparing' reaction exists in healthy human adults exposed to acute hypoxia and recedes with acclimatization. Preferential distribution of cardiac output to the brain upon acute altitude exposure may represent a conserved mechanism that protects against hypoxic brain damage in mammals, particularly in regions associated with basic cardiovascular and respiratory control during periods of acute hypoxia. Measurements of regional cerebral metabolism are needed to determine if 'brain sparing' effectively matches DO_2 , or if the increase in CBF represents a protective form of overcompensation.

Limitations

Our rapid ascent profile in combination with supplemental oxygen during transport from low to high altitude was designed to induce an abrupt change in PaO_2 , similar to that which can be achieved in laboratory studies with hypoxic gas or hypobaric

chambers. As such, our results must be interpreted in this context and thus may be expected to be different from other field studies that have followed more traditional progressive ascents (Huang *et al.*, 1987; Jensen *et al.*, 1990; Baumgartner *et al.*, 1994; Willie *et al.*, 2013).

We used duplex sonography primarily because it is a non-invasive technique that can be utilized in field settings. This technique yields volumetric measurements, in terms of ml/min, which, based on first principles, can be multiplied by CaO_2 to yield DO_2 . Our low CVs were in line with a previous study showing similarity between duplex sonography and both PET and xenon inhalation methods of measuring gCBF (Schoning & Scheel, 1996). Nevertheless, we acknowledge that all these techniques are limited by the lack of an absolute standard for validating CBF. Our gCBF measurements were based on unilateral, left-sided measurements of the ICA and VA – the main arteries perfusing the brain. While left VA flow has been reported to be ~20% higher than the right (Schoning *et al.*, 1994), this was not expected to have an effect on global measurements since ICA flow represents the majority of gCBF (Schoning & Scheel, 1996). Yet, unilateral VA measurements may have influenced our finding of increased VA DO_2 . Future studies are needed to determine if ‘brain sparing’ effects are attenuated when independent measurements of left and right VA flow are summed.

Since the ICA feeds the MCA, we expected that changes in ICA flow would be reflected in $\text{MCA}_{\text{velocity}}$. This was not the case: ICA flow increased ~70% while

MCA_{velocity} was unchanged throughout the study. A similar discrepancy between ICA flow and MCA_{velocity} has been previously described by Willie *et al.* (Willie *et al.*, 2012) and argued to support dilation of the MCA in hypoxia (Wilson *et al.*, 2011). We calculated that a 12% increase in MCA diameter could explain the measured discrepancy between ICA_{flow} and MCA_{velocity} . This exact degree of vasodilation has recently been demonstrated at high altitude with a color-coded ultrasound technique (Willie *et al.*, 2013), yet because additional studies are needed to clarify artery-specific responses to hypoxia and validate MCA-diameter measurement techniques, we chose to refrain from further interpretation of MCA_{velocity} .

Summary & Implications

Overall, our findings highlight the integrative nature of responses that preserve oxygen delivery to the brain at high altitude. Regional cerebral vasoreactivity to O_2 and CO_2 may favor oxygen delivery to posterior and inferior regions of the brain during acute hypoxia to sustain vital cerebral functions associated with homeostasis. Whether these mechanisms evolved to promote survival in conditions provoking cerebral hypoxia is not clear at present, but further research in this area may yield important insights into human tolerance and adaptation to chronic states of hypoxemia.

Competing Interests

The authors have no conflicts or competing interests to disclose.

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Table 1. Cardiopulmonary and hematological values [mean \pm SD (n)]

| Variable | | SL | ALT1 | ALT16 |
|-------------------|-------|-----------------------|-----------------------|-------------------------|
| VE | l/min | 12.05 \pm 2.50 (21) | 11.93 \pm 2.92 (17) | 14.88 \pm 2.65 (21)*# |
| PaO ₂ | mmHg | 102.2 \pm 5.5 (21) | 36.1 \pm 2.8 (18)* | 45.3 \pm 3.2 (20)*# |
| PaCO ₂ | mmHg | 38.1 \pm 4.4 (21) | 26.5 \pm 3.1 (18)* | 20.9 \pm 2.5 (20)*# |
| SaO ₂ | % | 98 \pm 1 (21) | 76 \pm 6 (18)* | 82 \pm 3 (20)*# |
| [Hb] | g/dl | 13.9 \pm 1.4 (21) | 14.2 \pm 1.5 (18)* | 16.0 \pm 2.0 (20)*# |
| CaO ₂ | ml/dl | 19.4 \pm 1.9 (21) | 15.2 \pm 2.1 (18)* | 18.4 \pm 2.4 (20)*# |
| HR | bpm | 76 \pm 12(21) | 90 \pm 16 (16)* | 96 \pm 13 (20)* |
| SV | ml | 91 \pm 27 (21) | 85 \pm 20 (16) | 83 \pm 21 (20) |
| Mean ABP | mmHg | 79 \pm 8 (21) | 76 \pm 13 (16) | 80 \pm 10 (20) |

* Different from SL

Different from
ALT1

Table 2. Cerebrovascular values [mean \pm SD (n)]

| Variable | | SL | ALT1 | ALT16 |
|-----------|-------------|----------------------|-----------------------|-----------------------|
| ICA Dia | cm | 0.51 \pm 0.08 (21) | 0.54 \pm 0.07 (16) | 0.50 \pm 0.07 (20)# |
| ICA Vel | cm/s | 29.8 \pm 8.2 (21) | 38.9 \pm 8.1 (16)* | 32.1 \pm 5.4 (20)# |
| ICA Flow | ml/min | 384 \pm 197 (21) | 556 \pm 203 (16)* | 379 \pm 97(20)# |
| ICA CVRi | mmHg/ml/min | 0.25 \pm 0.12 (21) | 0.16 \pm 0.09 (16)* | 0.23 \pm 0.07 (19)# |
| VA Dia | cm | 0.36 \pm 0.06 (20) | 0.41 \pm 0.06 (16)* | 0.36 \pm 0.06 (19)# |
| VA Vel | cm/s | 21.4 \pm 4.4 (20) | 24.4 \pm 6.4 (16) | 19.3 \pm 7.1 (19)# |
| VA Flow | ml/min | 133 \pm 47 (20) | 206 \pm 98 (16)* | 122 \pm 55 (19)# |
| VA CVRi | mmHg/ml/min | 0.66 \pm 0.24 (20) | 0.46 \pm 0.28 (16)* | 0.84 \pm 0.58 (19)# |
| gCBF | ml/min | 1057 \pm 413 (20) | 1524 \pm 456 (16)* | 981 \pm 223 (19)# |
| gCBF CVRi | mmHg/ml/min | 0.09 \pm 0.03 (20) | 0.05 \pm 0.02 (16)* | 0.08 \pm 0.02 (19)# |
| DO2 ICA | ml/min | 75 \pm 37 (21) | 84 \pm 32 (16) | 68 \pm 19 (19)# |
| DO2 VA | ml/min | 26 \pm 10 (20) | 31 \pm 16 (16)* | 22 \pm 11 (19)# |
| DO2 gCBF | ml/min | 206 \pm 79 (20) | 230 \pm 74 (16) | 181 \pm 51 (19)# |
| MCAv | cm/s | 59.5 \pm 10.3 (21) | 61.1 \pm 13.3 (17) | 57.7 \pm 7.1 (21) |
| MCA CVRi | mmHg/cm/s | 1.36 \pm 0.25 (21) | 1.28 \pm 0.32 (17) | 1.41 \pm 0.24 (20) |
| ICA %Q | % | 5.4 \pm 2.7 (21) | 7.6 \pm 2.7 (15)* | 4.8 \pm 1.4 (18)# |
| VA %Q | % | 1.9 \pm 0.8 (20) | 2.6 \pm 1.1 (15)* | 1.5 \pm 0.7 (18)# |
| gCBF %Q | % | 15.0 \pm 5.8 (20) | 20.4 \pm 6.2 (15)* | 12.6 \pm 3.4 (18)# |

* Different from SL

Different from ALT1

Figure 1. Reciprocal changes in global cerebral blood flow (gCBF) and arterial oxygen content (CaO₂) maintained global cerebral oxygen delivery (DO₂) across the study. * Different from sea level (SL). # Different from arrival at altitude (ALT1).

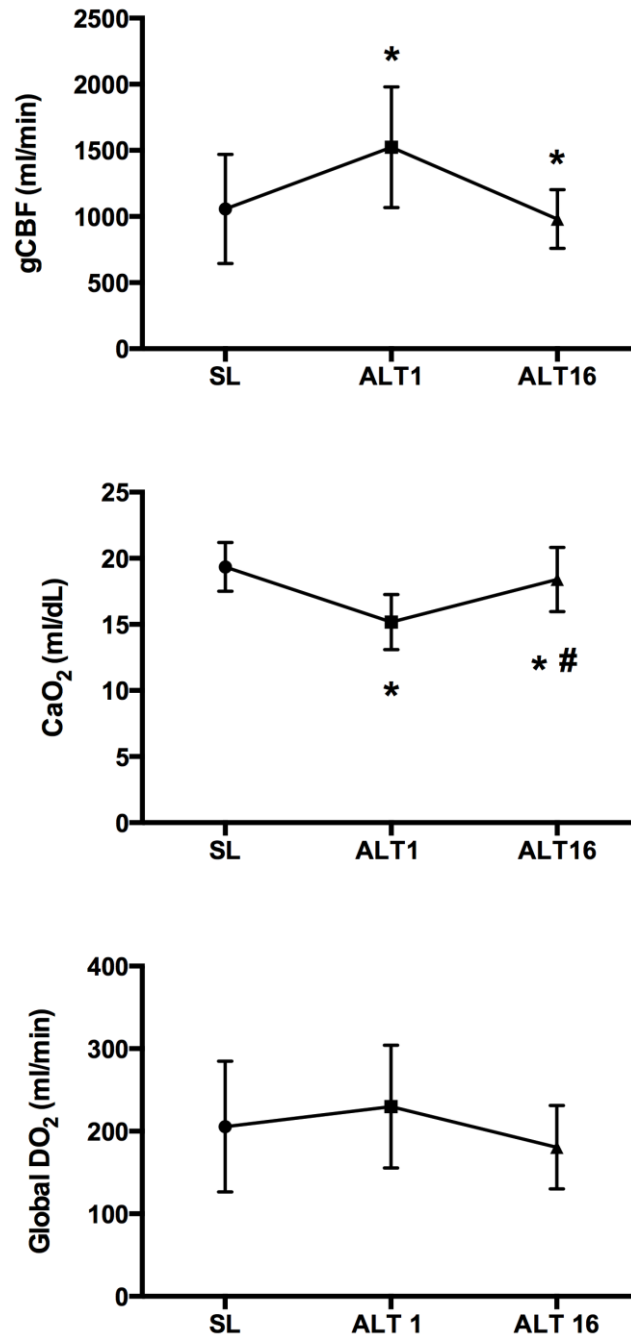
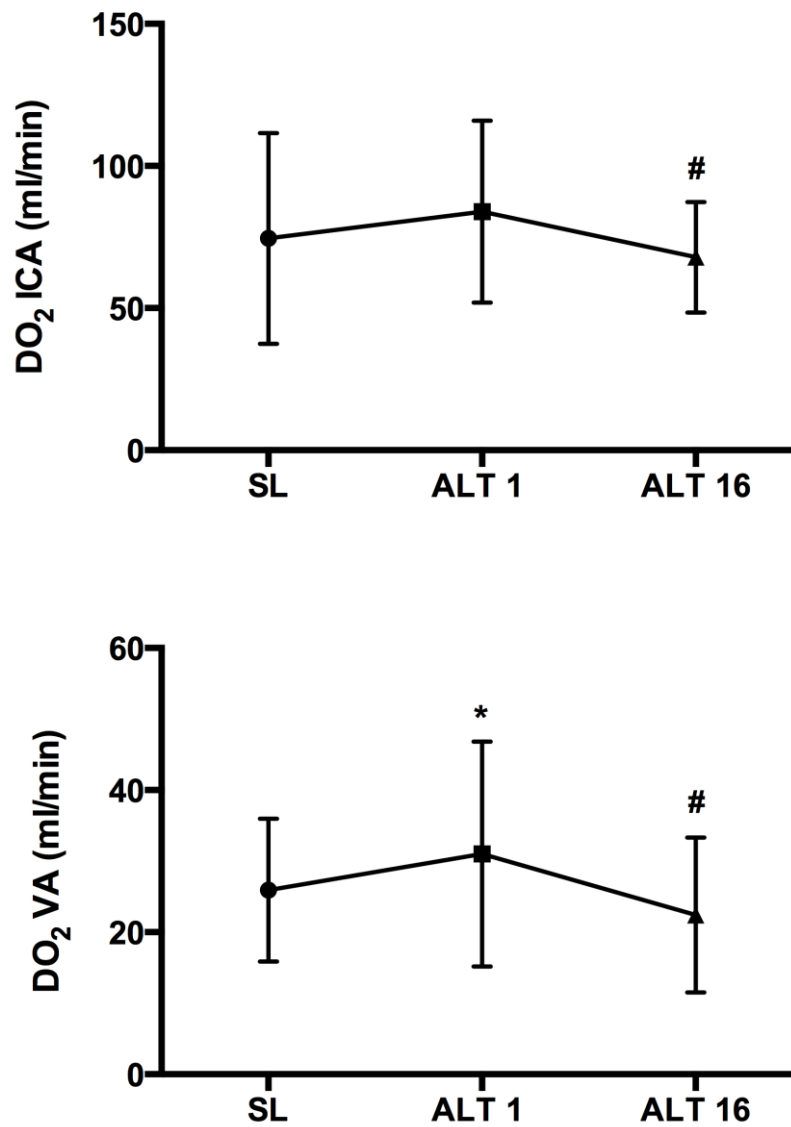


Figure 2. Regional oxygen delivery (DO_2) increases in the vertebral artery (VA), but not internal carotid artery (ICA) at ALT1. Regional DO_2 is reduced with acclimatization, but not below sea level (SL) values. * Different from SL. # Different from arrival at altitude (ALT1).



1 **AltitudeOmics: Enhanced cerebrovascular reactivity and ventilatory response to CO₂ with high**
2 **altitude acclimatisation and re-exposure**

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14

15 Running head: Cerebral function at altitude

16 Key words: cerebral blood flow, cerebral CO₂ reactivity, rebreathing, altitude acclimatisation

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25 **Abstract**

26 The present study is the first to examine the effect of high altitude acclimatisation and re-
27 exposure on the responses of cerebral blood flow and ventilation to CO₂. We also compared the
28 steady-state estimates of these parameters during acclimatisation with the modified rebreathing
29 method. We assessed changes in steady state responses of middle cerebral artery velocity
30 (MCAv), cerebrovascular conductance index (CVCi) and ventilation ($\dot{V}E$) to varied levels of CO₂ in
31 21 lowlanders (9 females; 21 ± 1 years), at sea-level (SL), during initial exposure to 5,260m (ALT1),
32 after 16 days of acclimatisation (ALT16) and upon re-exposure to altitude following either 7
33 (POST7) or 21 days (POST21) at low altitude (1,525m). In the non-acclimatised state (ALT1), MCAv
34 and $\dot{V}E$ responses to CO₂ were elevated compared to SL (by 79±75% and 14.8±12.3 L/min,
35 respectively, P=0.004 & P=0.011). Acclimatisation at ALT16 further elevated both MCAv and $\dot{V}E$
36 responses to CO₂ compared to ALT1 (by 89±70% and 48.3±32.0 L/min, respectively, P<0.001). The
37 acclimatisation gained for $\dot{V}E$ responses to CO₂ at ALT16 was retained by 38% upon re-exposure to
38 altitude at POST7 (P=0.004 vs. ALT1), while no retention was observed for the MCAv responses
39 (P>0.05). We found good agreement between steady-state and modified rebreathing estimates of
40 MCAv and $\dot{V}E$ responses to CO₂ across all three time points (P<0.001, pooled data). Regardless of
41 the method of assessment, altitude acclimatisation elevates both the cerebrovascular and
42 ventilatory responsiveness to CO₂. Our data further demonstrates that this enhanced ventilatory
43 CO₂ response is partly retained after 7 days at low altitude.

44

45 **Introduction**

46 The ability to maintain adequate oxygen transport to the brain by cerebral blood flow
47 (CBF) in hypoxic environments is vital. The CBF responsiveness to CO₂, termed cerebrovascular
48 CO₂ reactivity, provides a useful, non-invasive index of cerebrovascular function (3, 19). To date,
49 only a handful of studies have investigated the effect of acclimatisation to high altitude on
50 cerebrovascular CO₂ reactivity (1, 16, 17, 24, 30, 49). The interpretation of findings from these
51 studies is difficult due to the timing of measurements at high altitude (1, 16, 17, 24, 25), the
52 confounding effects of previous high-altitude exposure (1), artificial normobaric hypoxia (28, 46),
53 and the method used to assess reactivity (24, 30, 49). Data from Fan et al., (16, 17), obtained on
54 subjects at different stages of altitude acclimatisation, suggest that cerebrovascular CO₂ reactivity
55 is elevated with prolonged exposure to high altitude when using a modified rebreathing
56 technique. In contrast, Lucas et al., (30) reported a reduced cerebrovascular CO₂ reactivity in the
57 same subjects that at the end of a 14 day stay at 5,050 m, when assessed with a steady-state
58 technique (poikilocapnic hypoxia). More recently, Rupp et al., (49) reported a reduced
59 cerebrovascular CO₂ reactivity during steady-state hypoxic hypercapnia following 5 days at 4,350
60 m. Thus, the effect of altitude acclimatisation on cerebrovascular CO₂ reactivity remains unclear.

61 In addition, it is unknown if and for how long changes in cerebrovascular CO₂ reactivity
62 from acclimatisation persist after descent. Repetitive seven-month exposures to high altitude
63 were reported to improve arterial O₂ saturation (SaO₂), lower resting heart rate (HR) and decrease
64 susceptibility to acute mountain sickness (AMS) upon subsequent re-exposures (59). Remarkably,
65 these prior-exposure adaptations persisted despite a five-month deacclimatisation period. The
66 specific effect of high altitude re-exposure on cerebrovascular and ventilatory responsiveness to
67 CO₂ has yet to be examined.

68 Changes in cerebrovascular CO₂ reactivity with high-altitude acclimatisation depend on the
69 method of assessment. At sea level, the steady-state method results in higher cerebrovascular CO₂

70 reactivity (40-42) and lower ventilatory CO₂ sensitivity (6, 18, 23, 55) compared to the modified
71 rebreathing test. These differences have been attributed to the presence of a PCO₂ gradient
72 (between alveolar, arterial, and cerebrospinal fluid compartments) during the steady-state
73 method, which is supposedly abolished or minimised during rebreathing (6). Meanwhile, elevated
74 basal $\dot{V}E$ and subsequent underestimation of the ventilatory CO₂ sensitivity has been proposed as
75 one possible explanation for lower steady-state estimates (34). No studies have directly compared
76 the steady-state and modified rebreathing test estimates of cerebrovascular and ventilatory CO₂
77 responsiveness following ascent or acclimatisation to high altitude.

78 The purpose of the present study was therefore two-fold: first, we wished to assess the
79 effect of altitude exposure on cerebrovascular and ventilatory responsiveness to CO₂ in acute
80 conditions, after acclimatisation and upon re-exposure to high altitude after a period spent at low
81 altitude; second, we wished to compare the steady-state and modified rebreathing methods for
82 assessing the ventilatory and cerebrovascular responsiveness to CO₂ at high altitude.

83

84

85 **Methods**

86 *Subject recruitment and screening*

87 This study was conducted as part of the AltitudeOmics project. Following institutional
88 ethics approval, young (19-23 years old), healthy, sea level residents were recruited from the
89 greater Eugene, Oregon area (elevation 130 m). Potential subjects were screened to exclude
90 anyone who was born or had lived at altitudes >1500 m for more than one year or had travelled to
91 altitudes >1000 m in the past 3 months. A detailed description of subject recruitment procedures,
92 including inclusion and exclusion criteria have been presented elsewhere (54).

93

94 *Ethical approval*

95 The study was performed according to the *Declaration of Helsinki* and was approved by the
96 Institutional Review Boards of the Universities of Colorado and Oregon and by the Human
97 Research Protection Office of the U.S. Department of Defense. All participants were informed
98 regarding the procedures of this study, and written informed consent was given prior to
99 participation.

100

101 *Experimental Design*

102 After familiarisation with the experimental procedures outlined below (visit one), the
103 subjects underwent experimental trials near sea level (SL: 130 m, barometric pressure: 749
104 mmHg) and three times at high altitude (5,260 m, Mt Chacaltaya, Bolivia; barometric pressure 406
105 mmHg); on the 1st and 16th days at high altitude (ALT1 and ALT16) and again after either 7 (POST7;
106 n=14) or 21 (POST21; n=7) days at low altitude (1,525 m, barometric pressure: 639 mmHg). An
107 overview of the entire experimental design and protocol has been described in detail elsewhere
108 (54).

109

110 *Experimental protocol*

111 For each subject, all ALT measurements were carried out around the same time of day to
112 minimise any confounding effect of circadian rhythm. Measurements were taken upon arrival at
113 ALT1 to minimise the influence of AMS. Likewise, no symptoms of AMS were observed at ALT16 or
114 POST7.

115 For this study, following 10-15 min of quiet rest in a seated position, each experimental
116 testing session comprised of: a) instrumentation; b) 10 min room air baseline; and c)
117 cerebrovascular CO₂ reactivity tests. The cerebrovascular CO₂ reactivity tests consisted of: i) 10
118 min with end-tidal PCO₂ (PETCO₂) clamped at 40 mmHg; ii) 3 min voluntary hyperventilation to
119 lower PETCO₂ to ~20 mmHg; iii) the modified rebreathing test (details below); and iv) 3 min with

120 PETCO₂ clamped at 50 mmHg. The entire cerebrovascular CO₂ reactivity protocol was carried out
121 in background of hyperoxia (end-tidal PO₂ [PETO₂] > 250 mmHg).

122

123 *Experimental setup*

124 Throughout the protocol, the subjects sat upright and breathed through a mouthpiece
125 attached to a two-way non-rebreathing valve (Hans-Rudolph 2700, Hans-Rudolph Inc., Shawnee,
126 KS, USA). The breathing circuit allowed switching from room air to either an end-tidal clamping
127 system or a rebreathing system. The end-tidal clamping setup used in the present study is a
128 modified version of the system previously described by Olin et al., (39). The setup allowed
129 stabilising PETCO₂ at 40 and 50 mmHg. Throughout the end-tidal PCO₂ clamping, we maintained
130 PETO₂ at >250 mmHg by titrating 50% or 100% O₂ into the inspiratory reservoir at SL and ALT,
131 respectively.

132

133 *Modified rebreathing method*

134 The modified rebreathing method is a well-established method for assessing both
135 ventilatory and cerebrovascular CO₂ reactivities (14, 16, 34, 41). By using hyperoxia (PETO₂ > 250
136 mmHg) the test minimises peripheral chemoreceptors' output (11, 21) and the ventilatory
137 response to the modified rebreathing method can thus be interpreted as the ventilatory CO₂
138 sensitivity primarily from the central chemoreflex. The details of the modified rebreathing method
139 have been previously described in Fan et al., (16, 17). The rebreathing bag was filled with gas to
140 achieve inspired PCO₂ and PO₂ of 0 mmHg and 300 mmHg, respectively, at each altitude. Subjects
141 were instructed to hyperventilate for 3 min (part ii) to lower and then maintain PETCO₂ at 20
142 mmHg at both sea level and 5,260 m (in background PETO₂ > 250 mmHg). Subjects were then
143 switched to the rebreathing bag, and following two initial deep breaths to mix the gas from the
144 bag with that in the respiratory system, they were instructed to breathe *ad libitum* (part iii). The

145 rebreathing tests were terminated when PETCO₂ reached 50 mmHg, PETO₂ dropped below 200
146 mmHg or the subject reached the end of his/her hypercapnic tolerance.

147

148 *Measurements*

149 *Cerebrovascular variables:* Middle cerebral artery velocity (MCAv, an index of cerebral
150 blood flow) was measured in the left middle cerebral artery using a 2-MHz pulsed Doppler
151 ultrasound system (ST3, Spencer technology, Seattle, WA, USA). The Doppler ultrasound probe
152 was positioned over the left temporal window and held in place with an adjustable plastic
153 headband (Marc 600 Headframe, Spencer technology, Seattle, WA, USA). The signal was acquired
154 at depths ranging from 43 to 54 mm. Signal quality was optimised and an M-mode screen shot
155 was recorded to facilitate subsequent probe placements. Peripheral saturation was measured on
156 the right side of the forehead by pulse oximetry (N-200, Nellcor Inc., Hayward, CA, USA).

157 *Cardiovascular variables:* Beat-to-beat mean arterial blood pressure (MAP) was measured
158 from an arterial catheter inserted in a radial artery, and connected to a calibrated, fluid-filled,
159 disposable pressure transducer positioned at the level of the heart (DELTRAN II, Utah Medical, Salt
160 Lake City, UT, USA). Heart rate (HR) was determined with a three-lead ECG (ADInstruments
161 BioAmp & Micromaxx, SonoSite Inc., Bothell, WA, USA). Cerebrovascular conductance index (CVCi)
162 was calculated using the equation $CVCi = MCAv/MAP$ and normalised to values obtained at a
163 PETCO₂ of 20 mmHg and expressed as percentage change.

164 *Respiratory variables:* $\dot{V}E$ was measured using a pneumotachograph (Universal Ventilation
165 Meter, Vacu•Med, Ventura, CA, USA; Ultima™ series, Medgraphics CPX, Minneapolis, MN, USA)
166 and expressed in units adjusted to BTPS. PETO₂ and PETCO₂ were measured using fast responding
167 gas analysers (O₂Cap Oxygen analyser, Oxigraf, Mountain View, CA, USA). The pneumotachograph
168 was calibrated using a 3-L syringe (Han-Rudolph 5530, Kansas City, KS, USA) and the gas analysers

169 were calibrated using gas mixtures of known concentrations of O₂ and CO₂ prior to each testing
170 session.

171 *Arterial blood gas variables:* A 20-22 gauge arterial catheter was placed into a radial artery
172 and blood samples (2 mL) were taken over approximately 5 cardiac cycle periods. Core body
173 temperature was telemetrically recorded from an ingestion pill (CorTemp, HQInc, Palmetto, FL,
174 USA). All samples were analysed immediately for arterial pH, PO₂ (PaO₂), PCO₂ (PaCO₂) (Rapidlab™
175 248, Siemens Healthcare Diagnostics Inc., Henkestrasse, Germany), haemoglobin concentration
176 ([Hb]) and O₂ saturation (SaO₂) (Radiometer OSM3, Radiometer Medical ApS, Copenhagen,
177 Denmark). The blood gas values were analysed in triplicate and temperature corrected (26, 53).
178 Arterial bicarbonate concentration ([HCO₃⁻]) was subsequently calculated using the Henderson-
179 Hasselbalch equation.

180

181 *Data acquisition*

182 All analog data were sampled and recorded at 200Hz on a PC for off-line analysis
183 (ADInstruments Powerlab 16/30, Bella Vista, Australia).

184

185 **Data analysis**

186 *Steady-state responses*

187 Since the subjects could not tolerate PETCO₂ clamping at 50 mmHg at ALT16, the steady-
188 state MCAv-CO₂, MAP-CO₂ and CVCi-CO₂ slopes were estimated from the difference in mean
189 MCAv, MAP and CVCi at the end of 20 and 40 mmHg PETCO₂ clamp (20 sec averages), plotted
190 against the change in PaCO₂ between these two conditions across all time points (SL, ALT1, ALT16,
191 POST7 and POST21). The absolute value of $\dot{V}E$ at clamp 40 mmHg was used as an estimate of
192 steady-state $\dot{V}E$ responsiveness to CO₂, since voluntary hyperventilation was necessary to reduce
193 PETCO₂ to 20 mmHg.

194

195 *Modified rebreathing*

196 The rebreathing data were first reduced to one-second averages across the entire
197 rebreathing period. The $\dot{V}E$ -CO₂ slopes were analysed using a specially-designed programme
198 (Analyse $\dot{V}E$ Rebreathing programme rev11, University of Toronto, Toronto, Canada), as previously
199 described (15, 16, 34). The MCAv-CO₂ slopes were analysed using a commercially available
200 graphing programme (Prism 5.0d, GraphPad Software Inc., San Diego, CA, USA), whereby
201 segmental linear regression (least squares fit) was used to estimate the MCAv-CO₂ slope during
202 the modified rebreathing. For comparison, we plotted the MCAv-CO₂ slopes using a sigmoid curve
203 as described by Battisti-Charbonney et al., (4), using the Prism programme. To minimise the sum
204 of squares for non-linear regression (Levenberg-Marquardt algorithm) we used the equation:

205
$$MCAv = a + (b/(1 + \exp(-(PETCO_2 - c)/d)))$$

206 Where MCAv is the dependent variable in cm/s, PETCO₂ is the independent variable in mmHg, *a* is
207 the minimum MCAv determined from the mean MCAv of the hypocapnic (hyperventilation)
208 region, *b* is the maximum MCAv value, *c* is the mid-point value of MCAv, and *d* is the range of the
209 linear portion of the sigmoid (inverse reflection of the slope of the linear portion).

210 We found good agreement in the MCAv-CO₂ slope obtained from these two models
211 ($R^2=0.71$). However, due to the range of PETCO₂ used in this study, segmental linear regression
212 generally provided better fit across all conditions, whereas the sigmoidal curve model was the
213 preferred model for only 12 out of 58 trials. As such, only the MCAv-CO₂ slopes obtained using the
214 segmental linear model are presented.

215

216 *Statistical analysis*

217 Due to logistics impacts on planning and transportation, not all subjects were able to
218 participate in all high-altitude studies, please see the Figures and Table for complete sample size

219 reporting for each procedure. Most data are reported as the improvement over the time of
220 acclimatization (change from ALT1 to ALT16) and as the amount of that improvement that was
221 retained after time at low altitude, calculated as % retention = $(\text{POST7 or POST21} - \text{ALT1}) / (\text{ALT16}$
222 $- \text{ALT1}) * 100$ (5). The effects of altitude acclimatisation and re-exposure (between SL, ALT1, ALT16,
223 POST7 and POST21) on the steady-state MCAv-CO₂ slope, CVCi-CO₂ slopes and $\dot{V}E$ at 40 mmHg,
224 were analysed using mixed model linear regression (IBM® SPSS® Statistics version 21, IBM®
225 Corporation, Armonk, NY, USA). To assess the effects of altitude acclimatisation (between SL, ALT1
226 and ALT16) on the rebreathing estimates of MCAv-CO₂ and $\dot{V}E$ -CO₂ slopes, we used mixed model
227 linear regression analysis (Diagonal repeated covariance assumed). The interactions between
228 variables of interest were assessed using correlational (Pearson) analysis (IBM® SPSS®, Statistics
229 version 21). Data are shown as mean ± SD. Results were considered significant at the alpha level
230 <0.05. Trends were consider at the alpha <0.10 level. A priori power calculations ($\alpha = 0.05$, $\beta =$
231 0.20) were used to determine sample size and limit Type II error.

232

233

234 **Results**

235 Detailed baseline characteristics of the 21 (9 females; 21 ± 1 years old) subjects
236 participating in AltitudeOmics are presented elsewhere (54). All 21 subjects completed the
237 protocol at SL. Due to logistical issues, 4 of 21 subjects were unable to complete the entire
238 experimental protocol at ALT1. Upon re-exposure to altitude, 14 of 14 subjects completed the
239 protocol at POST7 and 5 of 7 at POST21. Due to low n, no comparison was carried out between
240 ALT1 and POST21

241

242 *Resting variables*

243 The resting variables across acclimatisation and re-exposure have already been reported in
244 detail elsewhere (54) and will not be reproduced in this paper.

245

246 *Steady-state method (Table 1)*

247 Acclimatisation: Compared to SL, the steady-state MCAv-CO₂ slope was elevated at ALT1
248 (by 79 ± 70%, P<0.001), and was further elevated at ALT16 (by 89 ± 70% vs. ALT1, P=0.001).
249 Similarly, the steady-state MAP-CO₂ slope was elevated at ALT1 (by 256 ± 265%, P=0.013) and
250 further elevated at ALT16 (by 164 ± 1370% vs. ALT1, P<0.001). The steady-state CVCi-CO₂ slope
251 was elevated at ALT1 (by 82 ± 79%, P<0.001), and remained higher at ALT16 (by 93 ± 81%, P<0.001
252 vs. SL, no difference with ALT1). $\dot{V}E$ at 40 mmHg was elevated at ALT1 compared to SL (by 14.8 ±
253 12.3 L/min, P=0.011), and further elevated at ALT16 (by 48.3 ± 32.0 L/min vs. ALT1, P<0.001).

254 Re-exposure: Upon re-exposure to altitude, it appears that the acclimatisation gained in the
255 steady-state MCAv-CO₂ slope was not retained at POST7 (P=0.145 vs. ALT1). Compared to ALT16,
256 the steady-state MCAv-CO₂ slope was lowered at both POST7 and POST21 (P=0.029 & P=0.003,
257 respectively), but nevertheless remained higher compared to SL (P<0.001 & P=0.024,
258 respectively). Similarly, 49% of the acclimatisation gained in the MAP-CO₂ slope was retained at
259 POST7. Specifically, the MAP-CO₂ slope remained higher at POST7 compared to ALT1 (P=0.005).
260 When compared to ALT16, the MAP-CO₂ slope was lowered at both POST7 and POST21 (P<0.001
261 for both). Nevertheless, MAP-CO₂ slope were higher at POST7 and POST21 compared to SL
262 (P<0.001 & P=0.020, respectively). In contrast, no difference was observed in the CVCi-CO₂ slope
263 at POST7 when compared to ALT1 or ALT16 (P=0.980 & P=0.804, respectively), but it remained
264 higher compared to SL (P<0.001). Likewise, CVCi-CO₂ slope tended to remain higher at POST21
265 compared to SL (P=0.058), but was not different from ALT16 (P=0.715).

266 Upon re-exposure, the effect of acclimatisation on the $\dot{V}E$ at 40 mmHg was retained by
267 38% at POST7 (P=0.004 vs. ALT1). Compared to ALT16, $\dot{V}E$ at 40 mmHg was lower at POST7 and

268 POST21 ($P=0.001$ & $P<0.001$, respectively), but these values remained higher when compared to
269 SL ($P<0.001$ & $P=0.001$, respectively).

270

271 *Modified rebreathing method (Table 1)*

272 Similar to the steady-state method, the rebreathing MCAv-CO₂ slope was elevated at ALT1
273 (by $137 \pm 117\%$, $P<0.001$), and further elevated at ALT16 (by $35 \pm 33\%$ vs. ALT1, $P=0.040$). The
274 rebreathing $\dot{V}E$ -CO₂ slope was elevated at ALT1 compared to SL (by 1.61 ± 1.14 L/min/mmHg,
275 $P=0.038$), and further elevated at ALT16 (by 2.86 ± 2.61 L/min/mmHg vs. ALT1, $P=0.004$). The
276 ventilatory recruitment threshold was lowered at ALT1 (by 4.4 ± 4.0 mmHg, $P<0.001$ vs. SL) and
277 further lowered at ALT16 (by 4.4 ± 3.2 mmHg vs. ALT1, $P<0.001$).

278

279 *Acid-base buffering capacity correlations (Figure 2)*

280 Based on previous findings (16), we performed correlations between the pooled steady-
281 state data with [HCO₃⁻] and found resting [HCO₃⁻] correlated with steady-state MCAv-CO₂ slope
282 ($R=-0.771$) and $\dot{V}E$ at 40 mmHg ($R=-0.723$, $P<0.001$ for both).

283

284 *Steady-state vs. modified rebreathing (Figure 3)*

285 We observed correlations between the steady-state and rebreathing MCAv-CO₂ slope at SL
286 ($R=0.609$, $P=0.003$), ALT1 ($R=0.817$, $P<0.001$) and ALT16 ($R=0.596$, $P=0.007$), while the pooled
287 MCAv-CO₂ slopes (combined SL, ALT1 and ALT16) between the two methods also correlated well
288 ($R=0.860$, $P<0.001$). Likewise, there were significant correlations between $\dot{V}E$ at 40 mmHg and the
289 rebreathing $\dot{V}E$ -CO₂ slope at SL ($R=0.476$, $P=0.029$), ALT1 ($R=0.506$, $P=0.038$) and ALT16 ($R=0.927$,
290 $P<0.001$), while the pooled ventilatory data across all time points were also correlated ($R=0.904$,
291 $P<0.001$).

292

293

294 **Discussion**

295 The present study is the first to assess the effect of altitude acclimatisation and re-
296 exposure on cerebrovascular CO₂ reactivity using both the steady-state and modified rebreathing
297 methods. We demonstrate that cerebrovascular CO₂ reactivity was elevated immediately upon
298 arrival to 5,260m and is further elevated following 16 days acclimatisation, regardless of the
299 method of assessment. In addition, we found that cerebrovascular and ventilatory responsiveness
300 to CO₂ remains elevated upon re-exposure to altitude, despite 7 or 21 days at low altitude. Since
301 these changes in cerebrovascular and ventilatory responsiveness to CO₂ correlated with the
302 changes in resting arterial [HCO₃⁻] across all time points, we speculate that these changes might be
303 partly due to an altered pH buffering capacity associated to exposure high altitude. Our data thus
304 demonstrate that the changes in cerebrovascular and ventilatory control gained due to altitude
305 acclimatisation over a period of 16 days are partially preserved upon subsequent exposure to
306 altitude, at least for up to a period of 3 weeks spent at low altitude.

307

308 *Effects of acclimatisation on cerebrovascular CO₂ reactivity*

309 Our findings extend those from Fan et al., (16, 17) by demonstrating that the MCAv-CO₂
310 slope is elevated upon arrival at 5,260 m and further elevated following 16 days of acclimatisation
311 (Fig. 1A). Importantly, previous studies by Fan et al., (16, 17) assessed MCAv-CO₂ slope in subjects
312 whom spent 8 days ascending to 5,050 m, while the subjects in the present study ascended rapidly
313 to altitude (•3 hours), thus making direct comparison difficult. Our findings contradict those of
314 Lucas et al., (30), who found that the MCAv-CO₂ slope was initially elevated at 5,050 m, but had
315 returned towards sea level values following two weeks at 5,050 m. However, because PETO₂ was
316 not controlled, the MCAv-CO₂ slopes reported by Lucas et al., (30) reflect MCAv changes from
317 *hypoxic hypocapnia* (room air breathing at 5,050 m; PETO₂ •48 mmHg & PETCO₂ 26-22 mmHg) to

318 *hypercapnic hyperoxia* ($PETO_2 > 310$ mmHg & $PETCO_2 \bullet 30$ mmHg), and thus do not represent
319 isolated reactivity to CO_2 . Rupp et al., (49) recently found the MCAv response to steady-state
320 *hypoxic hypercapnia* ($PETO_2 = 55$ mmHg) to be reduced following 5 days at 4,350 m. Therefore,
321 discrepancies between findings Rupp et al., (49) and those of the present study can be attributed
322 the differences in $PETO_2$ (55 mmHg vs. >200 mmHg), altitude (4,350 m vs. 5,260 m), and the
323 acclimatisation state of the subjects (5 days vs. 16 days). The results from the present study
324 demonstrate, for the first time, that cerebrovascular CO_2 reactivity *per se* is enhanced with
325 acclimatisation to high altitude when studied using a background level of hyperoxia. Furthermore,
326 discrepancy between studies highlights how methodological differences can yield vastly different
327 results. Thus future studies are warranted to clarify the effect of hypoxic and hyperoxic
328 background on assessing cerebrovascular functions at both sea-level and following ascent to high
329 altitude.

330

331 *Altered acid-base buffering capacity?*

332 During altitude acclimatisation, there is a progressive and parallel reduction in arterial and
333 cerebrospinal fluid (CSF) bicarbonate concentration which serves to compensate for the changes
334 in pH associated with hyperventilation-induced hypocapnia (12, 13, 20). These changes in acid-
335 base buffering capacity, in both the arterial and CSF compartments, would lead to a greater rise in
336 arterial and CSF $[H^+]$ for a given rise in $PaCO_2$. In support of this notion, lowering CSF bicarbonate
337 concentration elevates the cerebrovascular CO_2 reactivity in an anaesthetised dog model (27),
338 while bicarbonate infusion increases cerebral perfusion pressure in post-traumatic head injury
339 patients (9), elevates cerebral blood volume in preterm infants (57), and lowers ventilation in
340 healthy exercising humans at sea-level (44). As such, it has been suggested that the MCAv
341 responses to CO_2 at high altitude are linked to changes in arterial acid-base balance (16, 25). In the
342 present study, we observed concomitant increases in cerebrovascular and ventilatory

343 responsiveness to CO₂ with acclimatisation to high altitude and re-exposure (Fig. 1), which
344 occurred in parallel to the changes in [HCO₃⁻] (Fig. 2). While it should be acknowledged that such
345 correlations do not imply causality, the possible role for acid-base status changes on
346 cerebrovascular and ventilatory responsiveness to CO₂ at high altitude remains to be further
347 studied.

348

349 *Interaction between cerebrovascular and ventilatory responsiveness to CO₂*

350 Interaction between cerebrovascular CO₂ reactivity and the central chemoreceptor
351 activation was first alluded to by Heyman et al., (22) and has been subsequently expanded upon
352 by others (10, 16-18, 38, 43, 60-62). It was postulated that changes in cerebrovascular CO₂
353 reactivity affect the stability of ventilatory response to CO₂ by modulating the degree of H⁺
354 washout at the level of the central chemoreceptor (38). Accordingly, a blunted cerebrovascular
355 CO₂ reactivity would lead to less central H⁺ washout and subsequently greater central
356 chemoreceptor activation. Conversely, an enhanced cerebrovascular CO₂ reactivity would result in
357 lower central [H⁺] and therefore lower ventilatory CO₂ sensitivity. In agreement with previous
358 altitude studies (16, 17), we observed concomitant increases cerebrovascular and ventilatory
359 responsiveness to CO₂ (Fig. 1). These findings seem to contradict the modulating role of
360 cerebrovascular CO₂ reactivity on central chemoreceptor activation, possibly due to other
361 overriding factors such as enhanced central chemosensitivity and changes in acid-base balance
362 associated with ascent to high altitude. Future work is necessary to further unravel the interaction
363 between the regulation of cerebral blood flow and ventilation.

364

365 *Going back up*

366 Despite the large body of literature regarding high altitude acclimatisation over the past
367 century, the effect of prior exposure on physiological parameters during subsequent exposures is

368 not well documented. Most attention focused on the effect of a recent altitude exposure on the
369 risk for AMS (7, 31, 45, 51), or the rate of ascent (56). However, the dose of previous altitude
370 exposure and acclimatisation were generally not controlled in these studies. Wu et al., (59) found
371 a progressive reduction in the incidence of AMS, lower HR and higher SpO₂ in lowland railroad
372 workers over the course of several seven-month exposures to high altitude interspersed with 5
373 months spent at low altitude. Similarly, MacNutt et al., (32) found faster rate of ascent, lower AMS
374 and higher SpO₂ in trekkers with a recent altitude exposure compared to altitude naive trekkers,
375 despite a 7-30 day de-acclimatisation period. In the present study, we compared the
376 cerebrovascular and ventilatory responsiveness to CO₂ with acclimatisation and upon re-exposure
377 to 5,260 m following a period of either 7 or 21 days at low altitude. We found that 38% of the gain
378 in ventilatory response to CO₂ over acclimatisation was retained at POST7 (Fig. 1C), while
379 essentially none of the gain in MCAv-CO₂ reactivity over acclimatisation was retained at POST7
380 (Fig. 1A). Regardless of the underpinning mechanism(s), our findings suggest that the effect of
381 previous altitude acclimatisation over 16 days on ventilatory response to CO₂ is partially retained
382 after 7 days at low altitude, while it is reversed in the cerebrovascular response to CO₂. Our data
383 extends those by Muza et al., (36) which showed that ventilatory acclimatisation gained at 4,300
384 m is retained following 8 days spent at low altitude. Since we found the CVCi-CO₂ slope to be
385 consistently elevated by 60-80% across all time points (Fig. 1D), while the changes MAP-CO₂ slope
386 closely follows the changes in MCAv-CO₂ slopes (Fig. 1B), we speculate that the changes in MCAv-
387 CO₂ slopes at high altitude can be primarily accounted for by an enhanced sensitivity of the
388 cerebral vessels to CO₂, whereas the remainder can be attributed to an enhanced perfusion
389 pressure response.

390

391 *Steady-state or modified rebreathing method?*

392 There has been much debate over the use of the steady-state or the modified rebreathing
393 method for the assessment of cerebrovascular and ventilatory control, and attempts at consensus
394 have produced no uniform agreement [(18, 40), also see (2, 14) for reviews]. The steady-state
395 ventilatory responses to CO₂ were found to be either similar (34, 37, 40-42, 47) or lower (6, 18, 23,
396 55) when compared to rebreathing estimates., while steady-state cerebrovascular CO₂ reactivity
397 has been shown to be consistently higher than rebreathing values (18, 40-42). The present study
398 demonstrates that the changes in cerebrovascular and ventilatory CO₂ responsiveness with
399 altitude acclimatisation were similar between the steady-state and the modified rebreathing
400 method (Table 1) – possibly due to tight control of arterial PCO₂ and PO₂ with our end-tidal
401 clamping setup. Moreover, we observed strong correlations in these parameters between the two
402 methods across all time points (Fig. 3). We therefore conclude that both methods can be used to
403 assess the changes in cerebrovascular and ventilatory responses to CO₂ with high altitude
404 exposure and acclimatisation, provided that the level of CO₂ is comparable across all the
405 conditions, under identical level of background O₂.

406

407 **Limitations**

408 Although the present study provided the opportunity to assess the effects of
409 acclimatisation and re-exposure to 5,260 m on the cerebrovascular CO₂ reactivity, an important
410 methodological consideration should be acknowledged when interpreting our findings. In the
411 present study, transcranial Doppler ultrasound (TCD) was used to measure the MCAv, as an index
412 of global CBF changes during initial exposure, acclimatisation and subsequent re-exposure to
413 5,260 m. This is based on the assumption that: i) the MCA carries approximately upwards of 80%
414 of the overall blood flow to the respective hemisphere (29); ii) changes in MCAv reflect changes in
415 global CBF (8, 52); iii) the changes in MCAv in response to PaCO₂ changes are comparable to the
416 changes of internal carotid blood flow (50); and iii) the diameter of the MCA does not change

417 during the observed changes in arterial blood gases (52). In support, MCAv has been shown to
418 reflect changes in CBF assessed with the direct Fick method, at least during initial exposure to high
419 altitude (33, 35, 48).

420 Recent findings by Wilson et al., (58) indicate that the diameter of MCA, as measured using
421 TCD, vary, depending on the altitude (e.g., 5.30 mm at 75 m, 5.51 mm at 3,500 m, 5.23 mm at
422 5,300 m and 9.34 mm at 7,950 m). Importantly, the results from Wilson et al., (58) demonstrate
423 that the MCA diameter remains relatively unchanged up to 5,300 m. It should be noted that the
424 MCA diameters measured with TCD in that study were 80-90% greater than the values obtained
425 using magnetic resonance imaging in the same subjects. Since our measurements were carried out
426 in background hyperoxia ($PETCO_2 > 300$ mmHg), it seems unlikely that our cerebral blood velocity
427 values would be confounded by any effect of hypoxia-induced vasodilation of the MCA. Further
428 studies are needed to evaluate MCAv responses to CO_2 while holding $PETO_2$ at consistent levels of
429 hypoxia.

430

431 *Conclusion*

432 Findings from the present study clearly show that both cerebrovascular and ventilatory
433 responsiveness to CO_2 is elevated upon arrival at high altitude and further elevated with
434 acclimatisation. We demonstrate, for the first time, that this effect of high altitude acclimatisation
435 on the ventilatory response to CO_2 is partially retained after a period at low altitude, while prior
436 acclimatisation has no effect of the cerebrovascular response to CO_2 . Our data suggest that the
437 increased cerebrovascular CO_2 reactivity with acclimatisation may be accounted for by the
438 changes in acid-base balance in the blood and possibly the cerebrospinal fluid compartment.

439

440

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448

449 **Author contributions**

450 JF contributed to conception and design of the experiments, data collection, data analysis,
451 data interpretation, manuscript drafting and editorial process. OE contributed to the design of the
452 experiments, data collection, data analysis and manuscript revision. NB contributed to the data
453 collection and the manuscript revision. BK contributed to the interpretation of the data and the
454 revision of the manuscript. AL, AS and RR conceived, designed and executed the AltitudeOmics
455 study of which the present study was a part, and contributed to manuscript revision. AS also
456 contributed to data collection and data interpretation. All authors approved the final version of
457 the manuscript.

458

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- 641

642 **Table 1.** Cerebrovascular and ventilatory reactivities parameters during the steady-state and modified rebreathing (mean \pm SD).
 643

| | SL (n=21) | ALT1 (n=17) | ALT16 (n=20) | POST7 (n=14) | POST21 (n=5) |
|---------------------------------------------------|------------------|--------------------|---------------------|----------------------|--------------------|
| Steady-state | | | | | |
| MCAv-PaCO ₂ slope (cm/s/mmHg) | 1.19 \pm 0.42 | 2.16 \pm 1.05* | 3.39 \pm 0.89*† | 2.68 \pm 0.88*§ | 2.06 \pm 0.57*§ |
| CVCi-PaCO ₂ slope (%/mmHg) | 3.35 \pm 1.21 | 5.87 \pm 2.60* | 5.75 \pm 1.85* | 5.89 \pm 1.23* | 5.41 \pm 1.78* |
| MAP-PaCO ₂ slope (L/min) | 0.03 \pm 0.24 | 0.28 \pm 0.19* | 1.06 \pm 0.45*† | 0.56 \pm 0.29*§ | 0.32 \pm 0.18*§ |
| $\dot{V}E$ at 40 mmHg (L/min) | 19.15 \pm 4.89 | 34.06 \pm 12.23* | 80.05 \pm 32.32*† | 49.03 \pm 13.68*§† | 43.25 \pm 7.56*§ |
| Modified rebreathing | | | | | |
| MCAv-PETCO ₂ slope (cm/s/mmHg) | 1.34 \pm 0.60 | 2.95 \pm 1.11* | 3.67 \pm 0.87*† | - | - |
| $\dot{V}E$ -CO ₂ slope (L/min/mmHg) | 1.90 \pm 0.81 | 3.49 \pm 1.51* | 6.28 \pm 3.56*† | - | - |
| $\dot{V}E$ recruitment threshold (mmHg) | 38.7 \pm 3.4 | 33.7 \pm 3.7* | 29.2 \pm 2.1*† | - | - |

644 * different from SL (P<0.05); † different from ALT1 (P<0.05); § different from ALT16 (P<0.05).
 645
 646

647 **Figure legend**

648 **Figure 1** Changes in steady-state estimates of cerebrovascular, cardiovascular and ventilatory
649 responsiveness to CO₂ with acclimatisation and re-exposure to 5,260 m. Values expressed as mean
650 ± SD. * different from SL (P<0.05), † different from ALT1 (P<0.05), § different from ALT16 (P<0.05).

651

652 **Figure 2** Relationship between standard basic excess and steady-state cerebrovascular, ventilatory
653 and cardiovascular responsiveness to CO₂ with acclimatisation to altitude. * significant
654 correlations (P<0.05).

655

656 **Figure 3** Comparison of steady-state and rebreathing estimates of cerebrovascular and ventilatory
657 responsiveness of CO₂ with acclimatisation to 5,260 m. * significant correlations (P<0.05).

Figure 1

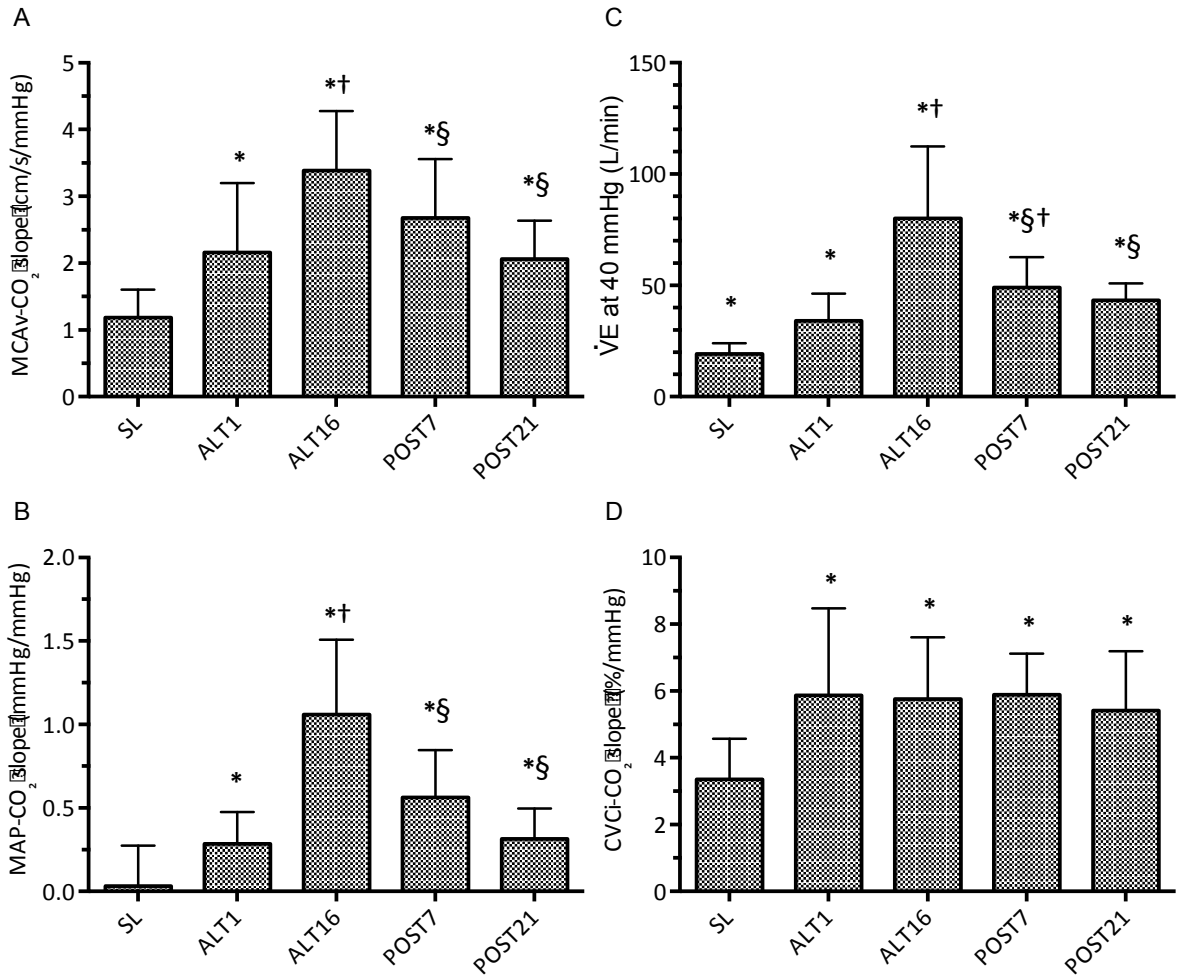


Figure 2

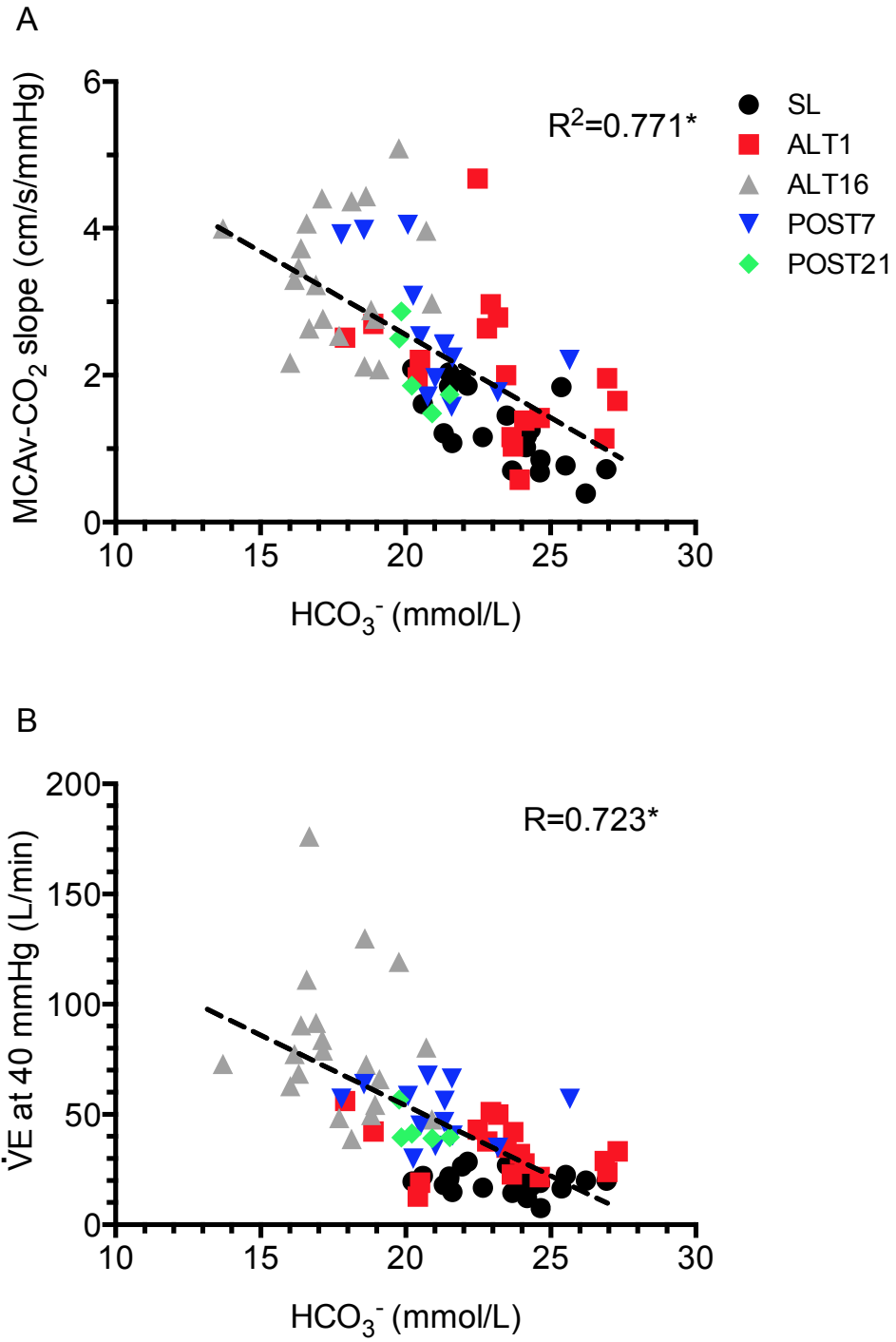
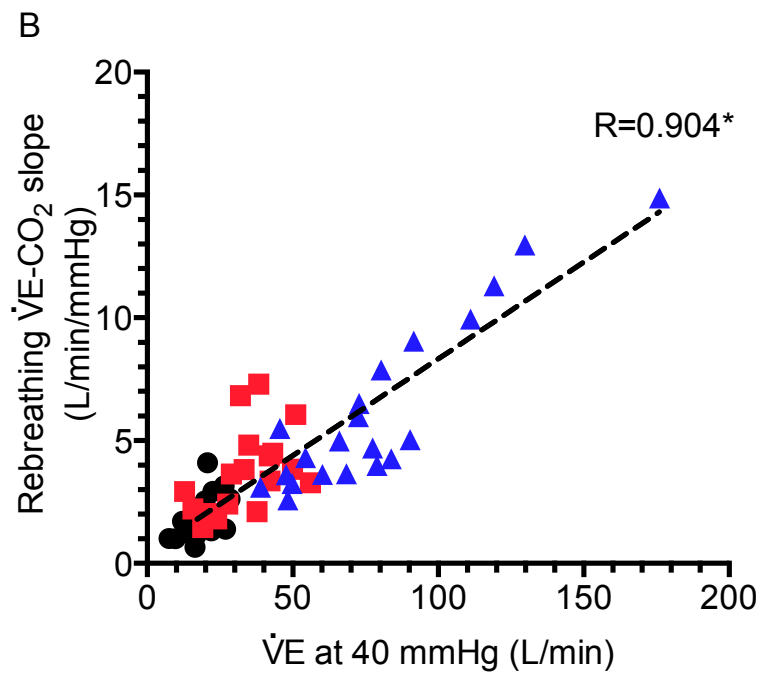
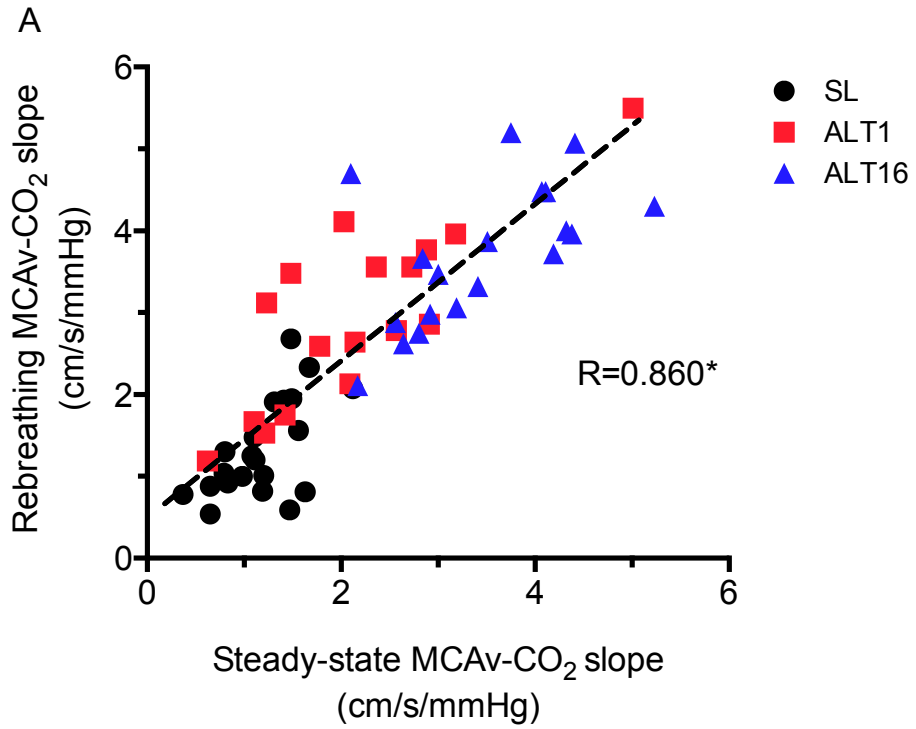


Figure 3



1 **AltitudeOmics: Cerebral autoregulation during ascent, acclimatization, and re-**
2 **exposure to high altitude and its relation with acute mountain sickness**

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4
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21 Running Head: Cerebral autoregulation at altitude

22
23 Key Words: transcranial Doppler, cerebral blood flow, cerebral oxygenation,
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46 **Abstract**

47 Cerebral autoregulation (CA) acts to maintain brain blood flow despite fluctuations
48 in perfusion pressure. Acute hypoxia is thought to impair CA, but it is unclear if CA is
49 affected by acclimatization or related to the development of acute mountain
50 sickness (AMS). We assessed changes in CA using transfer function analysis of
51 spontaneous fluctuations in radial artery blood pressure (indwelling catheter) and
52 resulting changes in middle cerebral artery blood flow velocity (transcranial
53 Doppler) in 21 active individuals at sea level (SL), upon arrival at 5,260 m (ALT1),
54 after 16 days of acclimatization (ALT16), and upon re-exposure to 5,260m after 7
55 days at 1,525 m (POST7). The Lake Louise Questionnaire (LLQ) was used to evaluate
56 AMS symptom severity. CA was impaired upon arrival at ALT1 ($P < 0.001$) and did
57 not change with acclimatization at ALT16 or upon re-exposure at POST7. CA was not
58 associated with AMS symptoms (all $R < 0.50$, $P > 0.05$). These findings suggest that
59 alterations in CA are an intrinsic consequence of hypoxia and are not directly
60 related to the occurrence or severity of AMS.

61

62 Introduction

63 Cerebral autoregulation (CA) is a general term used to describe dynamic myogenic,
64 neurologic and metabolic responses that adjust cerebrovascular resistance to
65 maintain relatively constant cerebral blood flow across a wide range of perfusion
66 pressures (25). Dynamic CA is said to be impaired if fluctuations in mean arterial
67 blood pressure lead to concurrent fluctuations in mean cerebral blood flow.
68 Impairments in CA are associated with cerebrovascular disorders (3, 24, 31), yet the
69 relative importance of CA in the development and course of certain pathologies is
70 unclear.

71

72 Our initial interest in CA stemmed from the hypothesis that impaired CA may be
73 involved in the development of acute mountain sickness (AMS), high-altitude
74 headache and cerebral edema (5, 7, 9, 16, 37). Conversely, we showed that
75 impairments in CA upon acute exposure to hypobaric hypoxia preceded, but were
76 not associated with, the development of AMS (2, 33, 35). Furthermore, since several
77 cross-sectional studies demonstrated that impairments in CA persist from 1 to 30
78 days of high-altitude exposure (1, 2, 11, 12, 17) - when AMS is not present - and are
79 evident in healthy, permanent high-altitude residents (12, 13), it seems reasonable
80 to suggest that a shift in CA may be an inherent and relatively benign consequence
81 of hypoxemia.

82

83 To date, no longitudinal studies have characterized CA and tested its relation with
84 AMS during acute and chronic high-altitude exposures. Previous studies have either

85 omitted CA measurements upon arrival at high altitude (7, 11, 17), or followed slow
86 ascent profiles that allow for partial acclimatization prior to initial measurements
87 (1, 12, 39). In this study, we present novel data from sea-level residents who rapidly
88 ascended to high altitude (5,260 m), acclimatized for 16 days, and were
89 subsequently re-exposed to high altitude after spending 7 days at low altitude
90 (1,525 m). Specifically, we tested the hypotheses that CA would be: 1) impaired
91 upon rapid ascent to high altitude, 2) unaffected by 16 days of acclimatization, 3)
92 unaffected upon re-exposure to the same altitude, and 4) unrelated to the
93 occurrence or severity of AMS.
94

95 **Methods**

96 **Study overview**

97 This study was conducted as part of the AltitudeOmics project. Briefly, institutional
98 ethics approval was obtained from the Universities of Colorado and Oregon and the
99 US Department of Defense Human Research Protection Office. Young, healthy sea-
100 level residents were recruited from the greater Eugene, Oregon area (elevation 128
101 m) and screened to exclude anyone who was born or had lived at altitudes >1,500 m
102 for more than one year or had traveled to altitudes > 1,000 m in the past 3 months.
103 After obtaining written consent, physical exams and the Army Physical Fitness Test
104 (push ups, sit ups and 3.2 km run) were performed to verify health and fitness
105 status. Approximately 4 weeks following sea-level (SL) measurements in Eugene,
106 Oregon, subjects were flown to La Paz, Bolivia. They spent two nights at low altitude
107 (Coroico, Bolivia, 1,525 m) before being driven to the Chacaltaya research station at
108 5,260 m while breathing supplemental oxygen. Acute responses to high altitude
109 were assessed ~4 hours after arrival and cessation of supplemental oxygen (ALT1).
110 Subjects acclimatized to altitudes ranging from 3,800 to 5,260 m over the next 15
111 days, with most of the time (75%) spent at 5,250 m. On the 16th day (ALT16),
112 measurements were repeated at 5,260 m before subjects were driven down to
113 Coroico for either 7 or 21 days. Subjects were driven back to the laboratory at 5,260
114 m for POST 7 or POST 21 re-exposure measurements.

115

116 This report focuses on novel data regarding resting CA evaluated immediately prior
117 to a series of cerebrovascular, respiratory and exercise interventions, as outlined
118 elsewhere (32). We have carefully avoided replication of data between reports,
119 except where common variables were necessary to describe subjects' basic
120 physiologic status at the time points of interest (e.g. heart rate, blood pressure,
121 arterial blood gases).

122

123 **Physiology Protocol**

124 All subjects were familiarized with study procedures during a practice session at
125 least 48 hours before experimental testing at SL. Subjects followed standardized
126 exercise and dietary regimens for 24 hours prior to each measurement period. At
127 each time point, a 22-gauge catheter was inserted into a radial artery at least 1 hour
128 prior to instrumentation. Subjects were seated in an upright position for 15 min
129 while sensors were placed to measure physiologic variables of interest. Limb lead
130 electrodes were used to measure ECG (BioAmp, ADInstruments, Colorado Springs,
131 CO, USA). Arterial blood pressure (ABP) was monitored via a fluid filled pressure
132 transducer (Deltran II, Utah Medical, Salt Lake City, UT, USA) attached to the radial
133 artery catheter. Core temperature was telemetrically recorded from an ingested pill
134 (CorTemp, HQInc, Palmetto, FL, USA). Cerebral blood flow velocity (CBFv) in the left
135 middle cerebral artery was measured by transcranial Doppler (2MHz Spencer
136 Technologies, Seattle USA) at depths ranging from 43 to 54 mm. Signal quality was
137 optimized and an M-mode screen shot was recorded to facilitate subsequent probe
138 placements and insonation angles.

139

140 After verification of signal quality, resting data were recorded for 6 min while
141 subjects breathed room air to assess CA at each altitude. Continuous analog data
142 (ABP, CBFv, ECG, O₂ and CO₂) were recorded at 200 Hz (ADInstruments Powerlab
143 16/30, Colorado Springs, CO, USA) for offline analysis. Core temperature and
144 arterial blood samples (2 ml) were taken during the last 30 s of measurement
145 periods. Blood samples were taken from the radial artery catheter and blood gases
146 were analyzed for PaCO₂ and PaO₂ in triplicate (RAPIDLab 248, Siemens, Erlangen,
147 Germany) and corrected for body temperature (15, 29).

148 **Acute Mountain Sickness**

149 Self reported sections of the Lake Louise Questionnaire (LLQ) were used to assess
150 AMS on ALT1 and POST7 (~12 hours after arrival). Moderate and severe AMS were
151 defined as LLQ ≥ 3 and ≥ 6 , including headache, respectively (27).

152 **Data Analysis**

153 Transfer function analyses were used to assess dynamic CA, based on spontaneous
154 fluctuations in the raw ABP and CBFv signals, as previously described (33, 34).
155 Briefly, 6-min recordings of instantaneous ABP and CBFv were reduced to beat-by-
156 beat averages, resampled at 5 Hz and transformed from the time to frequency
157 domain using fast Fourier transformations (512 points per segment with 40%
158 overlap). The transfer function from mean ABP to CBFv was expressed in terms of
159 coherence, gain, and phase shift in the very low frequency range (0.02 - 0.07 Hz),
160 where dynamic CA is most active (21, 22), as well as in low (0.07 to 0.20 Hz) and

161 high (0.20 to 0.35 Hz) frequency ranges. All data were used in subsequent statistical
162 analyses. Reduction in phase shift was considered the primary criterion for
163 impaired CA because it signifies shorter delay in transmission of pressure (ABP)
164 into flow (CBFv), or a reduction in the ability of the cerebrovascular system to buffer
165 changes in ABP and maintain consistent blood flow. Yet, since increases in gain
166 (increase in CBFv relative to a change in ABP) and coherence (linear correlation
167 between ABP and CBFv) may also suggest CA impairment (8, 24, 41), all three
168 transfer function metrics are reported. To address difficulties in interpreting
169 possible permutations of these three variables, the inverse transfer function of the
170 resulting gain and phase shift was used to express results in the time domain as a
171 step function that could be fitted to one of 10 curves representing a single
172 autoregulation index (ARI) score (36). An ARI score of 0 indicates complete lack of
173 autoregulation and 9 indicates perfect autoregulation.

174 **Statistics**

175 After calculating descriptive statistics (mean \pm SD) and verifying normality
176 (D'Agostino and Pearson Test), variables were analyzed by repeated measures
177 ANOVAs to evaluate the effect of time on CA metrics with Fisher's LSD *post hoc* tests
178 and the Holm procedure to correct for multiple comparisons ($\alpha = 0.05$).

179

180 Spearman rho correlations were run to evaluate relations between CA metrics and
181 the severity of LLQ symptom scores. Specifically, we tested the ability of CA
182 assessments, measured at SL and upon arrival at ALT1, to predict ensuing
183 symptoms of AMS (7). Also, because AMS classification is dichotomous (i.e. positive

184 vs. negative), we used receiver operating characteristic (ROC) analyses (14, 18) to
185 evaluate the sensitivity (true positive rate) and specificity (true negative rate) of
186 ARI scores' ability to detect mild and severe AMS. The ROC area under the curve
187 (AUC) statistic was used as an indicator of test accuracy. An AUC of 1.0 signifies a
188 perfect test, with no chance of false positive or false negative results, while an AUC
189 of 0.5 signifies a meaningless test, where the probability of identifying a true
190 positive result is only 50%.

191 **Results**

192 **Subjects**

193
194 We studied 21 subjects at SL (12 males and 9 females; 21 ± 1 years old). Because of
195 logistical problems upon arrival in Bolivia, complete data sets were not obtained on
196 the first 7 subjects upon arrival at ALT1. Since the first 7 subjects comprised the
197 cohort studied at POST21, longitudinal assessments of CA were limited to the
198 remaining 14 subjects who completed the study at POST7.

199

200 **Effect of Rapid Ascent to High Altitude**

201

202 At SL, resting cardiovascular (HR, ABP, CBFv) and CA measurements (coherence,
203 gain, phase shift and ARI scores) were characteristic of young, healthy individuals
204 with intact CA (Table 1, Figure 1). From SL to ALT1, PaO₂ and PaCO₂ decreased (65
205 and 26%, respectively, $P < 0.001$, Table 1). This degree of hypoxia increased HR ($P <$
206 0.001), but did not affect mean ABP or CBFv. Very low frequency power spectral

207 density (PSD) of ABP and CBFv were unaltered, but increases in transfer function
208 coherence ($P < 0.001$) and decreases in phase shift ($P < 0.050$) and ARI score ($P <$
209 0.001) were consistent (in 13 of 14 subjects) with the definition of impaired CA at
210 ALT1.

211

212 **Effect of Acclimatization to High Altitude**

213 Acclimatization increased resting PaO_2 (27%) and decreased PaCO_2 (22%) from
214 ALT1 to ALT16 (both $P < 0.001$), without affecting HR, ABP or CBFv. Measures of CA
215 at ALT16 were unchanged from ALT1 and remained impaired relative to SL in the
216 very low frequency range (all $P < 0.010$, Table 1, Figure 1).

217 **Effect of Re-exposure to High Altitude**

218 Resting PaO_2 and PaCO_2 at POST7 fell between ALT1 and ALT16 values (all $P < 0.050$
219 vs. ALT1 and vs. ALT16), indicating that the degree of acclimatization achieved at
220 ALT16 was partially maintained at POST7. Assessments of CA at POST7 were similar
221 to those at ALT1 and ALT16 and remained impaired relative to SL in the very low
222 frequency range ($P < 0.050$, Table 1, Figure 1).

223

224 **Association between CA and AMS**

225

226 Of the 21 subjects, 17 reported symptoms of at least moderate AMS at ALT1 (LLQ =
227 6.4 ± 2.2), 10 of who met the criteria for severe AMS (LLQ = 7.8 ± 1.7). Correlations
228 between CA metrics preceding the development of AMS symptom were weak (all
229 $r < 0.50$, $P > 0.050$, Figure 2). The ROC analysis revealed that ARI scores measured at

230 SL were not sensitive or specific predictors of moderate (AUC=0.54, P=0.788) or
231 severe AMS (AUC=0.69, P=0.139). Additionally, the degree of impairment in CA
232 (measured as the change in ARI from SL to ALT1) was not a sensitive or specific
233 predictor of moderate (AUC=0.53, P=0.881) or severe AMS (AUC=0.72, P=0.124).
234 None of the 14 subjects studied at POST7 reported symptoms of AMS, thus
235 associations with CA could not be tested.

236 **Discussion**

237 The key findings of this study were that cerebral autoregulation, as assessed by
238 transfer function analysis, is 1) impaired upon rapid ascent to high altitude, 2)
239 unaffected by acclimatization, or 3) subsequent re-exposure to the same altitude,
240 and 4) not a sensitive or specific predictor of AMS. Based on our results we question
241 whether the so-called impairment in CA that persists at high altitude is
242 characteristic of pathological insufficiency in cerebrovascular regulation (16), or
243 alternatively reflects a relatively benign relaxation in autoregulation.

244

245 **Effect of high altitude on CA**

246 This is the first longitudinal study of CA at high altitude, from rapid ascent through
247 acclimatization and upon re-exposure after a short period at low altitude. We show
248 that impairment of CA was a consistent characteristic across this high-altitude
249 exposure profile.

250

251 Increased transfer function coherence and gain along with reduced phase shift and
252 ARI score upon rapid ascent were all consistent with the classic definition of
253 impaired CA (Table 1) and outside the normal range of expected variability (6),
254 implying that changes in ABP were more readily transmitted into the cerebral
255 circulation as changes in CBFv at high altitude. Our finding of impaired CA after less
256 than one day of travel from low to high elevation is consistent with our previous
257 findings after 4 hours in a hypobaric chamber (35) and fills an important gap in the
258 literature between studies conducted in laboratories with hypoxic gas mixtures,
259 where normobaric hypoxia was achieved in a matter of minutes (5, 10, 26, 34), and
260 studies of trekkers, where several days of progressive ascent preceded initial high-
261 altitude measurements (1, 2, 12, 37). Impaired CA at rest in acute hypoxia is a
262 consistent finding among all but one study (26), suggesting that neither the mode
263 nor rate of ascent appears to affect the general assessment.

264

265 By evaluating CA upon initial exposure and after 16 days at high altitude, we were
266 able to determine if changes in CA occur with acclimatization, as might be expected
267 with increased PaO₂ (2, 35), decreased PaCO₂ (19, 23, 26), and further
268 sympathoexcitation (1). On the contrary, we found no change in CA over the course
269 of acclimatization (Table 1). Our longitudinal findings are consistent with other
270 cross-sectional studies demonstrating impaired CA at various time points after
271 arrival at high altitude (1, 2, 7, 11, 12, 37) and in permanent high-altitude residents
272 (12, 13). These results may indicate that assessments of CA are less sensitive to
273 changes in PaO₂ and PaCO₂ near their respective extremes. Alternatively, a slight

274 improvement in CA due to increased PaO₂ (2, 35) may have been masked if the
275 opposing effects of PaCO₂ (19, 23, 26) and/or sympathoexcitation (1) on CA were
276 heightened over time at altitude. Further testing with manipulation of arterial gases
277 and sympathetic activity is necessary to determine the relative influence of arterial
278 gases and neural stimulation on CA at high altitude, yet impaired CA remains a
279 consistent functional consequence across time at high altitude.

280

281 As an additional test of the hypothesis that impaired CA is a consistent response to
282 hypoxemia, we sent subjects down to low altitude for 7 days and re-evaluated their
283 CA response after a second rapid ascent back to high altitude. Upon re-exposure, the
284 measured impairment in CA was similar to that observed upon the first ascent
285 (ALT1) and after acclimatization (ALT16). Together, these results demonstrate that
286 impaired autoregulation was a consistent characteristic of hypoxemia across our
287 study and imply that slow fluctuations in arterial pressure were less effectively
288 dampened by the cerebral vasculature regardless of the state of acclimatization.
289 What remains to be determined is if such a tenuous pressure-flow relation may be
290 potentially harmful.

291

292 **Relation of CA to AMS**

293 Impairment of CA has been suggested to play a role in the development of AMS by
294 either permitting cerebral overperfusion and mechanical disruption of the blood
295 brain barrier (i.e. vasogenic cerebral edema) when mean ABP is elevated, or by
296 cerebral under-perfusion and exacerbation of cerebral hypoxia/ischemia when

297 mean ABP is lowered (9, 16). In the present study, we found no correlation between
298 measures of CA and subsequent AMS symptom scores (Figure 2), which opposes the
299 notion that lower CA predisposes people to AMS, or conversely, that higher CA
300 confers protection from AMS. Our additional ROC analyses of AMS status, confirmed
301 that ARI scores were neither sensitive nor specific indicators for the development of
302 moderate or severe AMS upon arrival at high altitude. These findings are congruent
303 with our previous report following the time course of changes in CA and AMS
304 symptoms over the first 10 hours of exposure to hypobaric hypoxia (35), where we
305 found similar levels of CA impairments in subjects who eventually developed AMS
306 or stayed healthy, but are at odds with other studies showing some association
307 between CA and AMS symptoms (5, 37). Our data also counter a recent finding that
308 sea-level assessments of CA predict ensuing severity of AMS (7).

309

310 Discrepancies between studies may be explained by the various methods used to
311 assess CA (transfer function vs. leg cuff – see Limitations), the questionnaires used
312 to assess AMS (LLQ vs. Environmental Symptoms Questionnaire), and the statistical
313 approach used to evaluate the relation between CA and AMS (correlation vs. ROC).
314 We acknowledge that caution should be exercised when interpreting correlations
315 with an ordinal level variable, such as the LLQ score, because by definition the scale
316 has limited mathematical meaning. For example, an LLQ score of 6 does not imply
317 symptom severity is exactly twice that of a score of 3. Due to the intrinsic level of
318 measurement, we believe that LLQ scores are best restricted to dichotomous
319 classification of positive or negative AMS status, and thus place more emphasis of

320 the negative results of our ROC analysis. We encourage others to consider this
321 method of analysis for future AMS studies.

322

323 Overall, given the similarity in CA responses among individuals with a wide range of
324 AMS scores, we do not believe that changes in CA cause AMS. This assertion is
325 further supported by the complete lack of association between impaired CA at
326 POST7 when no symptoms of AMS were reported and previous reports
327 documenting impaired CA in healthy high-altitude natives (12, 13). Nonetheless, we
328 must acknowledge that the alteration in CA upon acute altitude exposure may set up
329 a tenuous pressure-flow relation which could permit AMS to develop, if other, yet
330 unidentified, factors are present at the same time.

331

332 Since impairment of CA appears to be a consistent physiological response in hypoxic
333 environments and unrelated to AMS status, it is tempting to speculate that the
334 underlying change in the cerebral pressure-flow relation may actually promote
335 successful acclimatization or adaptation to chronic states of hypoxemia (4). It is
336 possible that impairment of CA could promote cerebral oxygen delivery in a time of
337 need since it allows greater cerebral perfusion for a given increase in ABP. This
338 potentially beneficial consequence of impaired CA during hypoxemic stress might
339 outweigh the relative risk of reduced cerebral perfusion if ABP were to drop. We
340 therefore raise the possibility that the term impaired CA may be a misnomer
341 because it implies an association with pathology that has yet to be substantiated in
342 acute or chronic hypoxemia. We suggest that relaxation of CA might be a more

343 accurate term to describe changes in the cerebral pressure-flow relation from
344 normoxia to hypoxia in the absence of pathology.

345

346 **Limitations**

347 One major limitation affecting the field is the lack of a gold standard method to
348 assess CA. We have chosen to evaluate rhythmical fluctuations in CA via transfer
349 function analysis, primarily because we believe it captures the natural cerebral
350 pressure-flow relation over time and thus has greater practical relevance over
351 methods which induce larger, more abrupt changes in ABP, as with leg cuff
352 inflation/deflation, rapid tilting, or more sustained changes in ABP, as with
353 pharmaceutical interventions. Still, we acknowledge that transfer function analysis
354 of resting data monitors relatively subtle fluctuations in ABP and CBFv, which, if
355 amplified, may not show impairment in CA (39). These factors may limit the
356 generalizability of resting CA assessments and lead to overstatement of the clinical
357 relevance of the findings. Additionally, there are no universal standards for the
358 parameter settings used in transfer function analysis or interpretation of
359 subsequent results, which makes comparisons between studies problematic. Future
360 work is needed to clarify differences in methods used to assess CA in hypoxemic
361 states and evaluate if these changes are generalizable to clinical settings.

362

363 Most CA studies rely on transcranial Doppler measurements of flow velocity and
364 assume that vessel diameter is unchanged, yet there is evidence to suggest that this
365 assumption may be invalid at extreme altitudes (39, 40). Dilation of the MCA at

366 ALT1 may explain why MCAv did not follow the expected increase in CBF upon
367 acute exposure to high altitude (30). We do not believe potential MCA dilation
368 affected our interpretation because the phase shift - our primary criterion for
369 assessing changes in CA – measures the relative timing of oscillations in ABP and
370 CBFv and thus is largely independent of absolute flow. However, since small changes
371 in diameter can have profound affects on flow (flow \sim radius⁴), future studies must
372 consider the use of continuous flow measurements, instead of velocity
373 measurements, to accurately assess CA in hypoxia.

374

375 Finally, our measurements of CA were limited to the MCA and relied on pressure
376 measurements taken in the radial artery. Since regional differences in
377 cerebrovascular regulation have recently been reported (20, 28, 38), more specific
378 measurements of regional pressure and flow are needed to fully characterize CA.

379 **Conclusions**

380 Our data demonstrate that the initial impairment of CA upon acute exposure to high
381 altitude is invariant with acclimatization and re-exposure, suggesting that relaxation
382 in the regulation of the cerebral pressure-flow relation is a characteristic response
383 to hypoxia that is unaffected by the degree of acclimatization. Since changes in CA do
384 not follow the progression and resolution of AMS, we question the clinical relevance
385 of impaired CA at high altitude.

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389 hypoxia and its subsequent retention. Many people and organizations invested
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530 **Figure Captions**

531

532 Figure 1. Arterial blood pressure to cerebral blood flow velocity transfer function

533 metrics (mean \pm SD from 0 to 0.5 Hz) at sea level (SL), upon arrival at 5,260 m

534 (ALT1), after 16 days of acclimatization (ALT16), and upon re-exposure to 5,260 m

535 after 7 days at low altitude (POST7). Similar impairments in cerebral

536 autoregulation (increased coherence and gain and decreased phase shift) from SL

537 were seen in the very low frequency (0.02 to 0.07 Hz – shaded area) at ALT1, ALT16,

538 and POST7 ($P < 0.05$). * Different from SL. & Different from SL.

539

540 Figure 2. Scatter plots showing no relation ($P > 0.05$) between autoregulation

541 indices (ARI), measured at sea level (SL, top) and as the change from SL to arrival at

542 high altitude (ALT1, bottom), and AMS symptoms scores from the Lake Louise

543 Questionnaire at ALT1.

Table 1. Resting Data (n=14, mean \pm SD)

| Variable | | SL | ALT1 | ALT16 | POST7 |
|-------------------|-------------------------|-----------------|------------------|------------------|-------------------|
| PaO ₂ | mmHg | 103 \pm 5 | 36 \pm 3* | 45 \pm 4*# | 42 \pm 4*#& |
| PaCO ₂ | mmHg | 37 \pm 4 | 28 \pm 2* | 21 \pm 3*# | 24 \pm 3*#& |
| HR | bpm | 73 \pm 9 | 90 \pm 18* | 95 \pm 12* | 85 \pm 15*& |
| ABP | mmHg | 77 \pm 6 | 76 \pm 14 | 81 \pm 10 | 76 \pm 8 |
| CBFv | cm/s | 62 \pm 9 | 63 \pm 14 | 59 \pm 7 | 57 \pm 9 |
| PSD ABP | mmHg ² /Hz | 11 \pm 13 | 9 \pm 4 | 9 \pm 5 | 6 \pm 4 |
| PSD CBFv | (cm/s) ² /Hz | 13 \pm 19 | 14 \pm 16 | 10 \pm 6 | 11 \pm 8 |
| Coherence | | 0.42 \pm 0.12 | 0.64 \pm 0.15* | 0.70 \pm 0.16* | 0.55 \pm 0.12*& |
| Gain | %/% | 0.64 \pm 0.24 | 0.88 \pm 0.35* | 0.85 \pm 0.25* | 0.97 \pm 0.33* |
| Phase Shift | radians | 0.48 \pm 0.28 | 0.17 \pm 0.21* | 0.27 \pm 0.09* | 0.25 \pm 0.19* |
| ARI | | 4.4 \pm 1.0 | 2.8 \pm 0.9* | 2.8 \pm 1.0* | 3.3 \pm 1.6* |

* Different from SL

Different from ALT1

& Different from ALT16

