Association between Body Water Status and Acute Mountain Sickness

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Abstract

Purpose: The present study determined the association between body fluid variation and the development of acute mountain sickness (AMS) in adults.

Methods: Forty-three healthy participants (26 males and 17 females, age: 26±6 yr, height: 174±9 cm, weight: 68±12 kg) were passively exposed at a FiO2 of 12.6% (simulated altitude hypoxia of 4500 m, PLO2 = 83.9 mmHg) for 12-h. AMS severity was assessed using the Lake Louise Score (LLS). Food and drink intakes were consumed ad libitum and measured; all urine was collected. Before and after the 12-h exposure, body weight and plasma osmolality were measured and whole-body bioimpedance analysis was performed.

Results: The overall AMS incidence was 43% (38% males, 50% females). Participants who developed AMS showed lower fluid losses (3.0±0.9 vs. 4.5±2.0 ml/kg/h, p = 0.002), a higher fluid retention (1.9±1.5 vs. 0.6±0.8 ml/kg/h, p = 0.022), greater plasma osmolality decreases (−7±7 vs. −2±5 mOsm/kg, p = 0.028) and a larger plasma volume expansion (11±10 vs. 1±15%, p = 0.041) compared to participants not developing AMS. Net water balance (fluid intake − fluid loss) and the amount of fluid loss were strong predictors whether getting sick or not (Nagelkerkes r² = 0.532). The LLS score was related to net water balance (r = 0.358, p = 0.018), changes in plasma osmolality (r = −0.325, p = 0.033) and sodium concentration (r = −0.305, p = 0.047). Changes in the impedance vector length were related to weight changes (r = −0.550, p < 0.001), fluid intake (r = −0.533, p < 0.001) and net water balance (r = −0.590, p < 0.001).

Conclusions: Participants developing AMS within 12 hours showed a positive net water balance due to low fluid loss. Thus measures to avoid excess fluid retention are likely to reduce AMS symptoms.

Introduction

Rapid ascents of non-acclimatized mountaineers to altitudes above 2,500 m are associated with the development of acute mountain sickness (AMS). This normally self-limiting syndrome is characterized by non-specific symptoms such as headache, dizziness, nausea, vomiting, loss of appetite, fatigue, and insomnia [1,2]. Symptoms of AMS typically appear 6–12 h after arrival at high altitude [3]. Risk factors associated with the development of AMS include the absolute altitude reached [4,5,6], the rate of ascent [4,7] and the individual susceptibility [1,3,7,8]. Controversy exists whether other factors such as low fluid intake and dehydration [6,9,10] or overhydration [11,12] promote AMS because of the lack of agreement in other studies [13,14,15]. These divergent findings may be explained by varied experimental designs (e.g., laboratory vs. field studies, resting vs. exercise conditions) and the lack of well controlled studies [15]. Less controversy exists on the impact of a decreased diuresis and fluid retention on AMS development [14,16,17,18]. The majority of studies point towards augmentation of AMS incidence by such disturbances even though prospective investigations have not always revealed a consistent difference in fluid balance between subjects developing AMS and those remaining well [19]. Controversy may arise due to the methodological divergences mentioned above. Additionally, changes in fluid balance have been reported to be different between hypobaric hypoxia vs. normobaric hypoxia conditions suggesting that both, reduced barometric pressure and oxygen partial pressure contribute to fluid retention [20]. However, the effect of different conditions on the AMS development has not been established yet. Thus, assessment of fluid homeostasis during passive normobaric hypoxia exposure (i.e., controlled normobaric hypoxic conditions, without the influence of exercise, hypobaria, cold) might help to clarify the pathophysiological relevance of hypoxia on the hydration status and the concomitant AMS development.

Citation: Gatterer H, Wille M, Faulhaber M, Lukaski H, Melmer A, et al. (2013) Association between Body Water Status and Acute Mountain Sickness. PLoS ONE 8(8): e73185. doi:10.1371/journal.pone.0073185

Editor: Jose A. L. Calbet, University of Las Palmas de Gran Canaria, Spain

Received April 25, 2013; Accepted July 17, 2013; Published August 27, 2013

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Funding: The study was funded by the Austrian National Bank (OENB). The funders had no role in study design, data collection and analysis, decision to publish, or preparation of the manuscript.

Competing Interests: The study was funded by the Austrian National Bank (OENB). This does not alter the authors’ adherence to all the PLOS ONE policies on sharing data and materials.

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PLOS ONE | www.plosone.org 1 August 2013 | Volume 8 | Issue 8 | e73185
A non-invasive method to monitor body fluid variation is the bioelectrical impedance analysis (BIA). BIA provides quantitative estimates of total, as well as intra- and extracellular water content based on regression equations. The R-Xc graph method on the other hand uses solely resistance (R) and reactance (Xc) normalized for standing height (bioimpedance vector analysis or BIVA) [21] and enables the classification of the hydration status independently of regression predictions which is considered the major weakness described for conventional BIA [22]. Piccoli et al. suggested it to be a useful tool in the planning of climber’s appropriate hydration and fluid intake during ascent to altitude [23].

As mentioned before, only one study analysed the effects of passive normobaric hypoxia exposure on fluid balance. The authors suggested extracellular fluid shifts to be involved in AMS development, but did not establish the exact nature of its impact [20]. Therefore, the present study aimed to assess the relationship between body fluid shifts/variation and the development of AMS during the early hours of a standardized passive normobaric hypoxia exposure in a large sample of men and women. According to Burtscher et al. there is no clear evidence for sex differences in the response to hypoxia [24] but as AMS incidence was sometimes reported to differ between sexes [25] perhaps because of the effects of sex hormones on ventilatory control [26] a comparison between men and women was made. We hypothesized that net fluid retention and extracellular fluid shifts during acute exposure to simulated hypoxia are associated with development of AMS with no differences between sexes and that we would be able to demonstrate this by the use of BIVA in combination with measures of conventional hydration markers (e.g., plasma osmolality).

Materials and Methods

Participants

Participants were recruited by advertisements on the homepage of the Austrian Alpine Association and by information via the mailing list of the University of Innsbruck. Exclusion criteria were reported cardiovascular, respiratory, neurological and psychiatric diseases, migraine, chronic headache, smoking, permanent residence at altitudes exceeding 1,000 m, an overnight stay at altitudes >2,500 m in the previous month, or exposure above >2,500 m two wk prior to the 12-hr hypoxic exposure. Fifty-six adults (32 males and 24 females) aged 18 to 45 yr, who fulfilled the inclusion criteria, volunteered to participate in the study. In the course of the 12-hr session, 13 participants suffered an emesis and the volume could not have been determined. Therefore, these participants were excluded from analysis, except for the calculation of the overall incidence rate. The final data set for all analysis included 43 participants, 26 males and 17 females.

Participants were instructed to abstain from all anti-inflammatory medication and nutritional supplements for two weeks prior to the exposure and from alcohol starting the day before the experiment. Caffeine was not allowed on the day of the exposure. All participants gave their written informed consent prior to the participation in the study. The study was carried out in conformity with the ethical standards laid down in the 2008 declaration of Helsinki and was approved by the ethics committee of the Medical University of Innsbruck.

Procedures

Participants were passively exposed at a FiO₂ of 12.6% (corresponding to a simulated altitude hypoxia of 4,300 m, P'O₂ = 83.9 mmHg) for 12 hr. Room temperature and humidity were kept constant at 22–24°C and 23–27%, respectively. Prior to entering the hypoxic chamber participants were examined, including a medical routine check. Women were tested for pregnancy before the 12-hr hypoxic exposure. During the simulated hypoxia exposure, the same food (i.e., brown bread, cheese, boiled ham, cucumber, banana, apple, cookies and chocolate) and drinks (water and apple juice) were provided and consumed ad libitum. Most of the time the participants stayed but some activities (e.g., standing, walking and stretching) were also performed. Recumbent position or sleeping was not allowed. Measurements, described in detail in the following sections, were performed before, during or and after the session.

Measurements and instruments

Lake Louise Score (LLS) and AMS. The LLS was used to assess the incidence and severity of AMS [27]. It is a self-assessment questionnaire including five symptom complexes (headache, gastrointestinal symptoms like anorexia, nausea or vomiting, fatigue and/or weakness, dizziness and/or light-headedness, difficulty sleeping); scores range from 0 to 3. The subjects self-rated their status: 0 for no discomfort, 1 for mild, 2 for moderate and 3 for severe symptoms. Because participants did not stay overnight in the hypoxic chamber, the symptom complex difficulty sleeping was not taken into account. AMS was diagnosed when the symptom headache was present and at least one other symptom, with a total score of either ≥ 3 to recognize mild diseases or AMS at an early stage or ≥ 4 to recognize a more severe diseases state [28]. The LLS at the point when leaving the chamber was taken to distinguish between AMS+ and AMS–.

Diuresis, insensible water loss, fluid and food intake. Urine volume, fluid and food intake (amount and composition) was recorded during the 12-hr exposure. Measuring cups were used to determine urine and fluid intake volume. For every participant the same limited number of food products was provided and food was weighted with a food scale before ingestion. Fluid content of the food was calculated by using the DGE-PC Professional software (Version 3.2, Bonn, Germany). The solid food content was calculated as the amount of food (g) minus the fluid content (ml) and was used to correct the body weight after the exposure (body weight minus solid food content). Insensible water loss (respiratory and surface water loss) was estimated to be 0.5 mL/kg/h [29]. Net water balance was calculated as fluid intake (water intake + water content of the food) minus fluid loss (urinary volume + insensible water). All volumes were normalised for body weight and exposure hours (ml/kg/h).

BIA, BIVA and body weight. Directly before entering and after leaving the chamber body weight was determined after urinating, wearing the same clothes and without shoes to the nearest 0.1 kg using a calibrated digital scale (Axensse easy coach, Bosch, Stuttgart, Germany). Bioimpedance analysis (BIA) was performed before and immediately after leaving the chamber using an impedance plethysmograph that emits an 800-μA 50-kHz alternating current (BIA-101, RJL/Akern Systems, Clinton Township, MI, USA). Before performing the measurements participants were asked to empty their bladder. Measurements were taken with subjects being in a supine position for 5 minutes, according to the manufacturers guidelines. Surface, contact electrodes (Biatrodes, Akern, Florence, Italy) were placed on the right hand and foot. The position of the electrodes was marked with a permanent marker in case the electrodes went off; otherwise they were left throughout the hypoxic exposure. A comparison between newly placed electrodes and electrodes left for 12 hr was performed in 6 participants. R and Xc values did not differ between condition [R = 473±39Ω vs. 473±40Ω (ICC r = 0.996,
Arterial oxygen saturation ($SpO_2$) was measured by pulseoxymetry (Onyx II 9550, NONIN, Plymouth, MI, USA) before the capillary blood samples were taken.

**Statistical analysis**

This study tested the hypothesis that altered fluid balance affects the development of AMS during simulated exposure to hypoxia. Data analyses were performed with the use of the SPSS statistical-software package (PASW Statistic 18). The chi-square test was used to calculate difference in incidence of AMS between men and women. Analysis of variance (ANOVA) for repeated measurements with 2 between-subjects factors (AMS+ vs. AMS−, men vs. women) and one within-subject factor (measurement time before and after the 12-hr exposure) was performed. Unpaired student-t tests were used to determine differences between sex and between AMS+ and AMS− for parameters of fluid balance (urinary volume, fluid intake, net water balance). Paired sample Hotellings $T^2$ test were used to calculate significant vector displacements and two sampe Hotellings $T^2$ test were used to determine vector differences between AMS− and AMS+. Pearson correlation analysis was used to assess the relationship between measured or calculated values and difference variables ($\Delta$) [post minus pre]. A stepwise logistic regression analysis, adjusted for sex and age, with backward variable selection was applied to identify predictor hydration variables for AMS+ and AMS−. Variables included in the analysis were $Na^+$, $ApOsm$, $Delta$vector length, $A$ weight, fluid intake, fluid loss and net water balance. The mean of values measured after 6 hr and immediately before leaving the chamber (i.e. $SpO_2$, $Na^+$, $HCO_3^-$, $PCO_2$, $PO_2$ and $pH$) was used for statistical analysis. A p-value of less than 0.05 (two-tailed) was considered to indicate statistical significance. Results are expressed as mean±standard deviation (SD), 95% confidence interval (CI) and odds ratio (OR).

**Results**

**AMS incidence**

The AMS incidence when using a LLS cut off of $\geq 3$ was 46.5% with no differences between sexes (30% and 41% for males and females, respectively; $p=0.571$). When using a LLS cut off of $\geq 4$ the AMS incidence was 26% and was similar between sexes (23% and 29% for males and females, respectively; $p=0.642$).

**Table 1.** Baseline characteristics (mean±SD) for participants who developed AMS (AMS+) and participants who did not (AMS−) separated for sexes.

|                | all | male | female | all AMS+ | all AMS− | male AMS+ | male AMS− | female AMS+ | female AMS− |
|----------------|-----|------|--------|----------|----------|-----------|-----------|-------------|-------------|
| n              | 43  | 26   | 17     | 32       | 11       | 6         | 5         | 12          |             |
| age (years)    | 26.1±5.7 | 26.1±5.6 | 26.1±6.1 | 30±7    | 25±5*    | 30.7±7.1  | 24.7±4.4  | 29.0±7.6    | 24.8±5.2    |
| body weight [kg] | 67.9±11.7 | 73.4±9.9 | 59.4±9.1* | 70.0±13.2 | 67.2±11.3 | 73.6±13.9 | 72.5±8.6  | 62.3±7.8    | 58.2±9.7    |
| body height [m] | 1.74±0.09 | 1.79±0.07 | 1.66±0.06* | 1.73±0.09 | 1.75±0.09 | 1.80±0.06 | 1.80±0.07 | 1.64±0.06   | 1.67±0.06   |
| Posm [mOsm/kg] | 297±3 | 298±3 | 296±3 | 297±3 | 297±3 | 296±4 | 298±3 | 297±3 | 295±4 |
| Na [mmol/L]    | 140±2 | 140±2 | 140±2 | 140±2 | 140±2 | 140±2 | 140±2 | 140±2 | 141±2 |
| R/height [m]   | 315±49 | 284±25 | 364±35* | 312±44 | 317±51 | 280±28 | 285±24 | 350±19 | 369±40 |
| Xc/height [l/m] | 37±4 | 36±4 | 39±4* | 37±5 | 37±4 | 35±4 | 36±4 | 40±3 | 39±4 |
| vector length  | 318±49 | 286±25 | 366±38* | 314±44 | 319±51 | 282±29 | 287±24 | 353±18 | 371±40 |

Plasma osmolarity (Posm), sodium (Na), resistance divided by body height (R/height), reactance divided by body height (Xc/height).

* indicates differences between sexes ($p<0.05$).

† indicates differences between AMS+ and AMS− ($p<0.05$).

‡ tendency to be different between AMS+ and AMS− ($0.05<p<0.1$).

doi:10.1371/journal.pone.0073185.t001
Body hydration, AMS and explanatory measures

When using the LLS cut off of ≥ 3 only a trend towards significant differences between AMS− and AMS+ were found with respect to net water balance (0.7±0.9 ml/kg/h vs. 1.3±1.3 ml/kg/h, p = 0.092, respectively). No differences were found for fluid intake and fluid loss (p > 0.05). Therefore in the following only the results for the more severe disease state (LLS ≥ 4) were shown. Baseline characteristics separated for sexes are shown in Table 1. AMS− and AMS+ differed in regard to fluid loss (4.5±2.0 ml/kg/h vs. 3.0±0.9 ml/kg/h for AMS− and AMS+, respectively; p = 0.002) and net water balance (0.6±0.8 ml/kg/h vs. 1.9±1.5 ml/kg/h for AMS− and AMS+, respectively; p = 0.022). No difference was found in regard to fluid intake (5.1±2.3 ml/kg/h vs. 4.9±1.4 ml/kg/h for AMS− and AMS+, respectively; p = 0.990). The regression analysis identified net water balance (1.09–8.68 CI, 3.1 OR) and fluid loss (0.29–1.05 CI, 0.55 OR) as predictor variables for the development of AMS (Nagelkerkes $\text{R}^2$ = 0.532 of the final model). Sodium intake did not differ between AMS− and AMS+ (1.7±0.8 g vs. 1.7±0.8 g, respectively; p = 0.861). Table 2 shows the outcomes for the hydration parameters. There were significant interactions (time x AMS+/−) for weight and Posm (p = 0.001 and p = 0.028). APV, obtained from the formula of Van Beaumont (1972) [32], showed significant differences between AMS+ and AMS− (11±10 vs. 11±15%, respectively, p = 0.041). Outcome of the correlation

### Table 2. Changes in hydration parameters (mean±SD) over the course of the 12 hour exposure (n=43).

| Parameter          | AMS+       | AMS−       | ANOVA main effect: time | ANOVA interaction: time x AMS+− |
|--------------------|------------|------------|-------------------------|---------------------------------|
| body weight (kg)   | pre        | post       | Δ%                      | pre                             | post       | Δ%                      |                                  |
|                    | 69.9±13.2  | 70.6±13.4  | +1.0                    | 67.2±11.3                       | 67.0±11.3  | −0.3                    | 0.037                            | 0.001                            |
| Posm [mOsm/kg]     | 297±3      | 290±9      | −2.4                    | 297±3                           | 295±5      | −0.7                    | <0.001                           | 0.028                            |
| Na [mmol/L]        | 140±2      | 137±4      | −2.1                    | 140±2                           | 140±2      | 0                       | 0.001                            | 0.050                            |
| Hct (%)            | 49.8±3.6   | 47.3±4.0   | −5.0                    | 49.2±4.0                        | 49.3±3.9   | +0.2                    | 0.050                            | 0.054                            |
| PV* [L]            | 2.7±0.5    | 3.0±0.5    | +11.1                   | 2.6±0.4                         | 2.6±0.5    | 0                       | 0.028                            | 0.052                            |
| TBW [L]            | 39.7±7.6   | 40.9±8.2   | +3.0                    | 39.4±6.7                        | 40.3±7.0   | +2.3                    | <0.001                           | 0.286                            |
| ECW [L]            | 16.6±3.0   | 17.5±3.2   | +5.4                    | 16.6±2.1                        | 17.2±2.4   | +3.6                    | <0.001                           | 0.151                            |
| ICW [L]            | 23.1±4.7   | 23.4±5.1   | +1.3                    | 22.8±4.9                        | 23.1±4.8   | +1.3                    | 0.005                            | 0.586                            |
| vector length      | 314±44     | 302±49     | −3.8                    | 319±51                          | 307±51     | −3.8                    | <0.001                           | 0.852                            |
| Δ values           |            |            | Unpaired t-test         |                                  |                                  |                                  |                                  |                                  |
| PV [%]             | 11±10      | 1.2±15     |                          |                                  |                                  |                                  |                                  | 0.041                            |
| net water balance  | 1.9±1.5    | 0.6±0.8    |                          |                                  |                                  |                                  |                                  | 0.022                            |
| fluid loss (ml/kg/h)| 3.0±0.9    | 4.5±2.0    |                          |                                  |                                  |                                  |                                  | 0.002                            |
| fluid intake (ml/kg/h)| 4.9±1.4  | 5.1±2.3    |                          |                                  |                                  |                                  |                                  | 0.999                            |

Plasma osmolarity (Posm), sodium (Na), hematocrit (Hct), total body water (TBW), resistance divided by body height (R/height), reactance divided by body height (Xc/height), plasma volume (PV).

*calculated from Hct assuming 77 ml/kg of blood volume.
#calculated from the formula of Van Beaumont W (1972) [32].

doi:10.1371/journal.pone.0073185.t002

Body hydration, AMS and explanatory measures

When using the LLS cut off of ≥ 3 and 6.7±2.2 and 1.6±1.1 when using the cut off of ≥ 4. When participants with emesis were included the overall incidence increased to 59% or 43% for cut off ≥ 3 or 4, respectively (males 59% or 38% and females 58% or 50%; p = 0.993 or 0.350). Due to very severe symptoms 3 males and 3 females out of 43 participants left the chamber before the end of the 12 hr (mean stay of these participants: males = 8.6±2.4 hr, females = 7.2±2.0 hr). No sex differences were found for the fluid intake, the fluid loss and for the net water balance (p > 0.05). No sex effects (interaction: time x sex) for changes in weight, Na**, Hct, R, Xc and vector length (p > 0.05) were found, except for Posm (p < 0.05). No interaction (time x sex x AMS+/−) was found for any of the analyzed variables. Because only small sex effects (see section above) were detected combined results of male and females were depicted in the following sections.

### Table 3. Correlation analysis for the LLS and the explanatory variables (n=43).

| Parameter          | Δ Posm | Δ Na+ | Δ weight | Δ Hct | SpO2 | HCO3 | PCO2 |
|--------------------|--------|-------|----------|-------|------|------|------|
| LLS                | 0.358* | −0.325* | −0.305* | 0.296* | −0.26* | −0.376* | 0.388* | 0.386* |

Lake Louise Score (LLS), plasma osmolarity (Posm), sodium (Na), hematocrit (Hct), arterial oxygen saturation (SpO2), bicarbonate concentration (HCO3−), partial pressure of carbon dioxide (PCO2).

*p < 0.05; **p < 0.1.
doi:10.1371/journal.pone.0073185.t003
Discussion

The main outcomes of the present investigation were that a net gain in water balance due to a low fluid loss (water retention) but not due to the amount of fluid intake discriminated participants who during the early hours of a passive normobaric hypoxia exposure developed a severe disease state (LLS ≥ 4). Furthermore, outcomes show that each 1 ml/kg/h of water gain leads to a 3.1 times higher, while every 1 ml/kg/h increase of fluid loss to a 45% reduced probability to develop AMS. Moreover, estimates of TBW and impedance vector displacements (Figure 2) show whole body fluid gain for both, AMS− and AMS+, whereas solely the AMS+ group showed fluid shifts into the intravascular system (indicated by Δ Posm and estimated Δ PV differences).

Only a limited number of studies investigated the influence of the body hydration status on the development of AMS with inconsistent findings. When considering solely fluid intake, Basnyat et al. and Mairer et al. found that low fluid intake was associated with AMS [6,9]. In contrast to this finding but in agreement with others [14,15], the present results do not support that low fluid intake was linked with AMS. Examination of net balance of fluid intake and fluid loss as well as the amount of fluid loss reveals their importance in the development of AMS compared to fluid intake alone. Accordingly, both, net water balance and the amount of fluid loss are strong predictors whether getting sick or not [Nagelkerkes r² = 0.532]. Even though Aoki et al. did not find body hydration to be a factor for the development of AMS [13], most studies found water retention to be linked to the onset of AMS symptoms [14,18,33,34]. This water retention was shown to occur already within the first 3 hours of hypoxic exposure [18] indicating that reduced fluid loss occurs ahead of AMS symptoms. The mechanisms explaining the effects of fluid retention on AMS development are not established. Some proposed effects include acute hypoxia-induced elevations of circulating arginine vasopressin levels [12,14,18], increased aldosterone, and atrial natriuretic peptide levels [14] that might cause anti-diuresis and/or water redistribution [14,12]. Importantly, bioimpedance and the R-Xc graph demonstrated a significant shortening of the vector that indicates fluid retention (Figure 2). This shortening was related to Δ body mass and parameters of fluid intake and output, but not with AMS symptoms or Δ Posm. As BIVA reflects whole body hydration status the finding of a significant correlation between Δ Posm and Δ Na⁺ with AMS and the larger PV expansion in AMS+ could mean that fluid shifts into the intravascular system were present.

The findings of an estimated 0.3 ± 0.3 L increase of plasma volume in the AMS+ group (Table 2) along with the finding that TBW and ECW changes only differed by this volume (i.e., 0.3 L, Table 2) between AMS+ and AMS− supports this finding. Accordingly,

### Table 4. Changes in HCO$_3^-$, SpO$_2$, pH and the blood gases (mean±SD) over the course of the 12 hour exposure.

|                  | AMS+                | AMS−                | ANOVA main effect: time | ANOVA interaction: time x AMS+/− |
|------------------|---------------------|---------------------|-------------------------|----------------------------------|
|                  | pre     | post | Δ%  | pre     | post | Δ%  |                  |                     |
| SpO$_2$ [%]      | 98±2    | 81±5 | −17 | 98±1    | 85±4 | −13 | <0.001            | 0.008               |
| HCO$_3^-$ [mmol/L]| 24.3±1.9 | 23.8±1.9 | −0.6 | 23.8±1.6 | 22.4±1.4 | −5.9 | <0.001            | 0.038               |
| pH               | 7.40±0.02 | 7.46±0.02 | +0.8 | 7.41±0.03 | 7.47±0.03 | +0.8 | <0.001            | 0.673               |
| PO$_2$ [mmHg]    | 73.1±5.6 | 35.9±3.8 | −50.9 | 77.0±6.9 | 38.2±3.8 | −50.4 | <0.001            | 0.413               |
| PCO$_2$ [mmHg]   | 39.6±2.7 | 34.2±3.2 | −13.6 | 37.9±3.8 | 31.5±3.2 | −16.9 | <0.001            | 0.393               |

Arterial oxygen saturation (SpO$_2$), bicarbonate concentration (HCO$_3$−), partial pressure of oxygen and carbon dioxide (PO$_2$ and PCO$_2$).

doi:10.1371/journal.pone.0073185.t004
and AMS group from before to after the sample size was large. AMS findings to the early development of AMS. Even though overall effects of hypoxia alone on fluid balance and connects these retention [20]. Nonetheless this study adds information on the pressure and reduced oxygen availability contribute to fluid outcomes of the study at hand. Present findings only hold true which could have influenced statistical outcomes. Not excluding parameters of body hydration, however, could not be detected.

Figure 2. Changes in the vector length and the 95% confidence ellipses of the AMS+ and AMS group from before to after the hypoxia exposure. In both groups a significant shortening of the vector was present (p<0.001, paired one sample Hotelling’s T2 test) indicating fluid gain [22]. There were no differences between groups (two sample Hotelling’s T2 test).

doi:10.1371/journal.pone.0073185.g002

Westerterp et al. found that subjects developing AMS showed the largest extracellular water shifts, even though not always in the same direction [34]. Furthermore, Loepky et al. reported increased peripheral blood flow and vasodilation in subjects with AMS [35].

The link of the peripheral responses (i.e. fluid retention and/or fluid shifts) to the development of AMS remains unexplained but increased extracellular water levels, beside other factors (e.g. changed endothelial permeability, up-regulation of inducible nitric oxide synthase), could influence BBB permeability and might increase intracranial pressure that causes symptoms of AMS [12]. Furthermore, inadequate diuresis might lessen excretion of bicarbonate, which otherwise would compensate for the respiratory alkalosis [36]. This results in a decrease in ventilation and as a consequence lower SpO2 which impacts AMS development [37]. Accordingly we found lower HCO3− and higher SpO2 values in AMS−, an association between HCO3−, PCO2 and SpO2 and an association between SpO2, HCO3− and PCO2 with the AMS score. A direct association between these parameters and parameters of body hydration, however, could not be detected.

Some issues have to be addressed when interpreting the outcomes of the study at hand. Present findings only hold true for normobaric hypoxia conditions, as both, reduced barometric pressure and reduced oxygen availability contribute to fluid retention [20]. Nonetheless this study adds information on the effects of hypoxia alone on fluid balance and connects these findings to the early development of AMS. Even though overall sample size was large AMS+ group involved only 11 participants, which could have influenced statistical outcomes. Not excluding subjects with emesis would have increased AMS+ sample size but as we were not able to determine volume of emesis, this procedure seemed adequate as emesis would have influenced hydration markers. As we did not perform multiple measurements prior to the development of AMS symptoms it is not possible to state whether water retention was cause or effect of AMS. Yet Loepky et.al. showed that water retention starts within the first 3 hours of hypoxic exposure in subjects that afterwards develop AMS suggesting that water retention occurs ahead of AMS symptoms [18]. Moreover, TBW and ECW were estimated from BIA values by using equations. It has to be mentioned that such formulas are depending on hydration status which obviously was changed and therefore might have influenced outcomes. Nonetheless the R-Xc graph method, shown to be a valid method for detection of body fluid volume changes independent of equations [21,22], confirmed the water gain. Of course isotopic measurements might have been more precisely. Furthermore, we did not measure urine specific gravity as another widely used hydration parameter. Nevertheless, according to Cheuvront et al., Posm is the only useful marker for static dehydration assessment and as accurate as urine specific gravity in the setting of dynamic dehydration assessment [38]. Furthermore, increased ventilation in hypoxia might have enhanced insensible water loss. This partly might account for the differences between body mass changes and net water balance volumes (approximately 600 ml). Additionally, the obviously elevated Hct levels are due to a systematic measurement error of the device (approximately 10%) but reliability of measurements was high and the observed changes should be addressed correctly. In this context it should be mentioned that plasma volume changes calculated solely from Hct values might to some extent be inaccurate [32] as is the assumption that all participants have a blood volume of 77 ml/kg. Finally it has to be recognized that blood gases were obtained from capillary blood. Even though ear lobe was arterialized the sample might have contained some peripheral venous blood.

Conclusion

In conclusion, the present findings confirm that a positive net water balance due to low fluid loss at least partly is involved in the early development of AMS under normobaric hypoxia conditions. Furthermore data suggest that most of the water gain differences between AMS+ and AMS− occurred within the vascular system. These results argue against “forced” or “over”-hydration for the prevention of AMS. From a clinical point of view measures to avoid excess fluid retention are likely to reduce AMS symptoms. However, future studies that manipulate fluid intake are needed to rigorously establish water intake recommendations at altitude.

Acknowledgments

We would like to thank all the volunteers who reserved a great amount of time to participate in the study.

Author Contributions

Conceived and designed the experiments: HG MW MB. Performed the experiments: HG MW MF. Analyzed the data: HG HL AM CE MB. Contributed reagents/materials/analysis tools: AM CE MB. Wrote the paper: HG MW MF HL AM CE MB.

References

1. Burtscher M, Mairer K, Wille M, Broessner G (2011) Risk factors for high-altitude headache in mountaineers. Cephalalgia 31: 706–711.

2. Hackett PH, Roach RC (2001) High-altitude illness. N Engl J Med 345: 107–114.
Dr. Baertsch P, Shaw S, Weidmann P, Oelz O (1993) Schützt Trinken vor der acute mountain sickness? Schweiz Z Sportmed 41: 7–13.

Dr. Akoi V S, Robinson S M (1971) Body hydration and the incidence and severity of acute mountain sickness: influence of susceptibility, provocation, and ascent rate. Med Sci Sports Exerc 34: 1886–1891.

Dr. Savourey G, Laumay J C, Bernard Y, Guinet-Lebreton A, Alonso A, et al. (2007) Normo or hypobaric hypoxic tests: propositions for the determination of the individual susceptibility to altitude illnesses. Eur J Appl Physiol 100: 193–205.

Dr. Baunyat B, Lemaster J, Litch J A (1999) Everest or bust: a cross sectional, epidemiological study of acute mountain sickness at 4243 meters in the Himalayas. Aviat Space Environ Med 70: 867–873.

Dr. Cumbo T A, Baunyat B, Graham J, Lescano A G, Gambert S (2002) Acute mountain sickness, dehydration, and bicarbonate clearance: preliminary field data from the Nepal Himalaya. Aviat Space Environ Med 73: 890–901.

Dr. Richardson A, Watt P, Maxwell N (2009) Hydration and the physiological responses to acute normobaric hypoxia. Wilderness Environ Med 20: 212–220.

Dr. Roach R C, Hackett P H (2001) Frontiers of hypoxia research: acute mountain sickness. J Exp Biol 204: 3161–3170.

Dr. Aoki V S, Robinson S M (1971) Body hydration and the incidence and severity of acute mountain sickness. J Appl Physiol 31: 363–367.

Dr. Bartsch P, Shaw S, Weidmann P, Oelz O (1993) Schützt Trinken vor der Bergkrankheit? Schweiz Z Sportmed 41: 7–13.

Dr. Castellani J W, Muza S R, Cheuvront S N, Sils I V, Fulco C S, et al. (2010) Effect of hypohydration and altitude exposure on aerobic exercise performance and acute mountain sickness. J Appl Physiol 109: 1792–1800.

Dr. Bartels P, Shaw S, Francioli M, Gnaedinger M P, Weidmann P (1988) Arrithmatriaque peptide in acute mountain sickness. J Appl Physiol 65: 1929–1937.

Dr. Krasney J A A (1994) A neurogenic basis for acute altitude illness. Med Sci Sports Exerc 26: 193–208.

Dr. Loeppky J A, Icenogle M V, Maes D, Riboni K, Hinghofer-Szlakay H, et al. (2005) Early fluid retention and severe acute mountain sickness. J Appl Physiol 98: 591–597.

Dr. Bartsch P, Swenson E R, Paul A, Julg B, Hohenhaus E (2002) Hypoxic ventilatory response, ventilation, gas exchange, and fluid balance in acute mountain sickness. High Alt Med Biol 3: 361–376.

Dr. Loeppky J A, Roach R C, Maes D, Hinghofer-Szlakay H, Reesler A, et al. (2005) Role of hypobaria in fluid balance response to hypoxia. High Alt Med Biol 6: 60–71.

Dr. Piccoli A, Rossi B, Pillon L, Bucchiante G (1994) A new method for monitoring body fluid variation by bioimpedance analysis: The RXc graph. Kidney Int 46: 534–539.

Dr. Lukaski H C, Piccoli A (2012) Bioelectrical impedance vector analysis for assessment of hydration in physiological states and clinical conditions. In: Preedy V R, editor. Handbook of Anthropometry: Physical Measures of Human Form in Health and Disease. London: Springer. pp. 287–306.

Dr. Piccoli A, Piazza P, Noventa D, Pillon L, Zaccaria M (1996) A new method for monitoring hydration at high altitude by bioimpedance analysis. Med Sci Sports Exerc 28: 1517–1522.

Dr. Bartsch M, Maier K, Wille M, Gatterer H, Ruedd G, et al. (2012) Short-term exposure to hypoxia for work and leisure activities in health and disease: which level of hypoxia is safe? Sleep Breath 16: 435–442.

Dr. Houignan B, Thetis M K, Koetzcl-McLain J, Roach R, Yip R, et al. (1993) Acute mountain sickness in a general tourist population at moderate altitudes. Ann Intern Med 118: 587–592.

Dr. Behan M, Werninger J M (2008) Sex steroid hormones and respiratory control. Respir Physiol Neurobiol 164: 213–221.

Dr. Roach R C, Bartsch P, Oelz O, Hackett P H (1993) The Lake Louise acute mountain sickness scoring system. In: Sutton J R, Houston C S, Coates G, eds. Hypoxia and Molecular Medicine. Burlington, VT: Queen City Printers. pp. 272–274.

Dr. Bartsch P, Bailey D M, Berger M M, Knauth M, Baumgartner R W (2004) Acute mountain sickness: controversies and advances. High Alt Med Biol 5: 110–124.

Dr. Jacob M, Chappell D, Hofmann-Kieser K, Conzen P, Peter K, et al. (2007) Determinants of insensible fluid loss. Perspiration, protein shift and endothelial glycocalyx. Anaesthesist 56: 747–764.

Dr. Sun S S, Chumlea W C, Heysenfield S B, Lukaski H C, Schoeller D, et al. (2003) Development of bioelectrical impedance analysis prediction equations for body composition with the use of a multicomponent model for use in epidemiologic surveys. Am J Clin Nutr 77: 331–340.

Dr. Lukaski H C, Hall C B, Siders W A (2007) Assessment of changes in hydration in women during pregnancy and postpartum with bioelectrical impedance vectors. Nutrition 23: 535–550.

Dr. Van Beumont W (1972) Evaluation of hematocrit from hematocrit measurements. J Appl Physiol 32: 712–713.

Dr. Bartsch P, Pfhuiger N, Auerfart M, Shaw S, Weidmann P, et al. (1991) Effects of slow ascent to 6559 M on fluid homeostasis. Aviat Space Environ Med 62: 105–110.

Dr. Westerterp K R, Robach P, Wouters L, Richalet J P (1996) Water balance and acute mountain sickness before and after arrival at high altitude of 4,550 m. J Appl Physiol 80: 1968–1972.

Dr. Loeppky J A, Icenogle M V, Maes D, Riboni K, Scorto P, et al. (2003) Body temperature, autonomic responses, and acute mountain sickness. High Alt Med Biol 4: 367–373.

Dr. Cerretelli P, Samaja M (2003) Acid–base balance at exercise in normoxia and in chronic hypoxia. Revisiting the “lactate paradox”. Eur J Appl Physiol 90: 431–440.

Dr. Bartsch M, Flatz M, Faulhaber M (2004) Prediction of susceptibility to acute mountain sickness by SaO2 values during short-term exposure to hypoxia. High Alt Med Biol 5: 335–340.

Dr. Cheuvront S N, Ely B R, Kenefick R W, Sawka M N (2010) Biological variation and diagnostic accuracy of dehydration assessment markers. Am J Clin Nutr 92: 565–573.