Live High-Train Low Altitude Training: Responders and Non-Responders

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Abstract

Objective: Investigate differences between athletes that responded (improved performance) compared to those that did not, after a 20-day "live high-train low" (LHTL) altitude training camp.

Methods: Ten elite triathletes completed 20 days of live high (1545-1650 m), train low (300 m) training. The athletes underwent (i), two 800-m swimming time trials at sea-level (1 week prior to and 1 week after the altitude camp) and (ii) two 10-min standardised submaximal cycling tests at altitude on day 1 and day 20 of the altitude camp. Acute mountain sickness (AMS) was also measured during the camp. Based on their 800-m swimming time trial performances, athletes were divided into responders (improved by 3.2 ± 2.2%, mean ± SD, n=6) and non-responders (decreased by 1.8 ± 1.2%, n=4).

Results: Compared to non-responders, the responders had lower exercise heart rates (-6.3 ± 7.8%, mean ± 90% CL, and 3.2 ± 2.2%, mean ± SD, n=6) and non-responders (decreased by 1.8 ± 1.2%, n=4).

Conclusion: Changes in SpO2, heart rate and some respiratory variables during exercise and resting AMS scores may help determine athletes that respond to LHTL altitude training camps from athletes that fail to respond to such training.

Keywords: Oxygen saturation; Acute mountain sickness; Submaximal exercise; Altitude training; Triathlon; Elite athletes

Introduction

The LHTL protocol (live at high altitude and train at or near sea level) has been shown to be the most effective altitude training method for improving sea level performance among athletes [1,2] and remains a very popular training method for elite athletes [3]. One of the key issues in the final outcome of LHTL altitude training is the problem of 'non-responders'. Some athletes thrive on altitude training and their performance improves, while others either fail to improve or suffer performance decline. It is currently not known why some athletes 'respond' to altitude and why others do not.

Many researchers have investigated the individual variability in athletes’ response to altitude training. Investigators have suggested that some athletes experience a better haematological response at altitude than others [4-6], which may help reduce stress by allowing more oxygen to be transported to the working muscle. Chapman et al. found that performance improvement was associated with increased Erythropoietin (Epo) which increased total red cell volume and subsequently VO2max after a 28-day altitude training camp in responders [5]. However, wide inter-individual variability has been found in erythropoietic response to altitude training which might not identify those elite athletes who respond to altitude training [7]. For example, Ge et al. reported individual responses in Epo ranged from -41 to 400% from baseline after 24 hours at 2800 m [8].

Recently researchers have searched for genetic determinants of the individual variability of the Epo response to altitude which may help explain why some athletes respond and some do not. While some researchers have identified specific genes associated with successful human existence at high altitude [9], others have reported little association between the proposed genes involved in Epo regulation and Epo response to hypoxia [10]. In addition, Hypoxia Inducible Factor (HIF-1) gene expression quantification after a 3-h hypoxia test performed before training was unable to predict poor and good responder athletes to the living high-training low model [11]. Others have suggested that individual difference in the tolerance to hypoxia may be explained by an increased Acute Mountain Sickness (AMS) incidence [12], which subsequently affects training and adaptation ultimately. In short, the differentiation between responders and non-responders is probably based on many factors including genetic predisposition [13], automatic nervous system adaptation [14], hypoxia-induced ventilator drive [15], underlying individual fitness levels, fatigue recovery and motivation [16]. It is clear that there is considerable individual variation in the physiological responses of athletes using altitude training, which makes the prediction of responders and non-responders very difficult. Some researchers have recently suggested screening individuals prior to altitude training may help identify those that may be negatively affected by such training [6], however pre-altitude testing is not always possible particularly on national squads that are commonly dispersed around the world.

The aim of the current project was to investigate different physiological (i.e., oxygen saturation, oxygen consumption, heart rate and performance responses) and subjective (acute mountain sickness (AMS), rate of perceived exertion (RPE)) responses of athletes undergoing training at a 20-day altitude camp in the hope of identifying possible parameters that may help identify responders from non-responders.

Methodology

Altitude

When deciding on an altitude to conduct a live-high train-low
model of training it is important to consider the effects of using an altitude that is high enough to produce a positive physiological response, but not so high as to reduce the recovery from training or cause unwanted clinical effects (nausea, dehydration etc.). A number of studies have suggested an altitude between 2000 and 2500 m is ideal for live-high train-low training, based mainly on the haematological response [8,17]. However recent research suggests athletes can improve performance after altitude training without positive haematological adaptation [18]. Therefore basing the altitude training threshold purely on haematological changes is probably unwise [19,20]. The altitude of 1545-1650 m used in this study was based mainly on practical reasons (a convenient ski lodge with all the necessary essentials was available at this altitude), but was also backed by contemporary research. Low to moderate altitude has previously been shown to produce significant improvements in sea level swim time trial performance (~1.9%) [21]. Indeed, in a recent study proclaiming the use of altitudes between 2000-2500 m the authors also found positive physiological (red cell mass volume ~7%, VO2 ~2%) and performance (~1%) effects for athletes on immediate return to sea level after living at 1754 m [17]; something that is not uncommon at these low altitudes (1200 m) [22]. Performance improvements of this magnitude would indicate meaningful effects for the very elite triathletes involved in this study [23].

Subjects

Ten elite triathletes were recruited from the New Zealand Academy of Sport development programme. All of the subjects (6 males and 4 females) who participated in the present study were international level athletes. The research was conducted over the summer period at Snow Farm (1545-1650 m), Wanaka, New Zealand. The study was approved by the Lincoln University Human Ethics Committee. Informed, voluntary, written consent was obtained from each subject prior to the start of the study. All subjects were free from injury, lived at sea level and had not been residents at altitude within the past 6 months. Subject characteristics and baseline measures of training are presented in Table 1.

Study design

The elite triathletes slept and stayed at the Snow Farm (1545-1650 m) and travelled to train at Wanaka (300 m) for approximately 2.7 ± 1.2 hour/day for 20 days during the camp. All subjects performed two exercise tests; swimming and submaximal cycling on day 1 and day 20 of the training camp. During the altitude camp, five subjects were randomly selected to receive iron supplementation of 1 capsule a day 20 of the training camp. During the altitude camp, five subjects performed two exercise tests; swimming and submaximal cycling on day 1 and day 20 of the training camp. During the altitude camp, five subjects were randomly selected to receive iron supplementation of 1 capsule a day.

Table 1: Characteristics and training workloads of athletes in the two training groups.

<table>
<thead>
<tr>
<th>Variables</th>
<th>Responders (n = 6)</th>
<th>Non-responders (n = 4)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Age (y)</td>
<td>23.5 ± 4.2</td>
<td>21.0 ± 2.0</td>
</tr>
<tr>
<td>Body mass (kg)</td>
<td>64.7 ± 7.4</td>
<td>66.1 ± 7.4</td>
</tr>
<tr>
<td>Height (cm)</td>
<td>173.2 ± 6.4</td>
<td>175.2 ± 5.1</td>
</tr>
<tr>
<td>Gender</td>
<td>Male 3; Female 3</td>
<td>Male 3; Female 1</td>
</tr>
<tr>
<td>BMI (kg/m²)</td>
<td>21.8 ± 1.3</td>
<td>21.8 ± 3.1</td>
</tr>
<tr>
<td>Swim Training (Trimp.d’)</td>
<td>163.7 ± 69.5</td>
<td>161.7 ± 68.3</td>
</tr>
<tr>
<td>Bike Training (Trimp.d’)</td>
<td>218.3 ± 123.7</td>
<td>184.3 ± 98.4</td>
</tr>
<tr>
<td>Run Training (Trimp.d’)</td>
<td>137.1 ± 96.1</td>
<td>113.9 ± 73.4</td>
</tr>
<tr>
<td>Total Training (Trimp.d’)</td>
<td>519.0 ± 41.4</td>
<td>459.8 ± 35.9</td>
</tr>
</tbody>
</table>

Note: Data are mean ± SD. No substantial difference between groups.

BP 325 mg (equivalent to 105 mg element iron), vitamin C 500 mg as sodium ascorbate (FERROGRAD® C, Abbott Laboratories (NZ) Ltd, Naenae)), while five subjects received a placebo. No contraindications for iron supplementation were found in any participants prior to the training camp, with all subjects showing normal iron, haemoglobin and ferritin levels (19.6 ± 4.4 umol.L-1, 145.0 g.L -1 ± 7.8, 82.9 ± 30.9 ug.L-1 mean ± SD for serum iron, haemoglobin and ferritin respectively). Retrospectively, based on their 800-m swim time trial, athletes were divided into responders (n=6) who had positive results (decreased their swimming time) and non-responders (n=4) who had negative results (increased their swimming time).

Testing

Maximal exercise test: Performance was assessed by individual 800-m timed swim trials in a standard 25- m pool near sea level. Trial times were performed at the same time each day with similar water temperature and brands of swimming costumes. Subjects completed a standardised 10-15 min warm-up with stretching prior to the swim. Subjects were instructed to try to achieve the best time possible during the swim, but were not provided with feedback on time or pace.

Submaximal exercise test: The submaximal cycling test was performed at altitude on each athlete’s own bicycle set up on a stationary ergometer (Cycle Ops Fluid 2, Madison, WI, USA) at the same time of day, on the first and last day of the altitude camp. Subjects performed the 10-min submaximal test at 250 W (for males) or 200 W (for females) during which heart rate (S610; Polar, Kempele, Finland), oxygen saturation (Sport-Stat, Nonin Medical, Minneapolis, MN) and rate of perceived exertion were recorded every minute. In addition, ventilation and expired gases were measured breath-by-breath using a portable gas exchange system (MetaMax® 3B; Cortex Biophysik, Leipzig, Germany). Before testing, the gas analyser was calibrated for volume (Hans Rudolph 5530 3 L syringe; Kansas City, MO, USA) and gas composition (15% O2 and 5% CO2). Oxygen consumption (VO2), minute ventilation (VE), and respiratory exchange ratio (RER) were measured continuously during the submaximal test, however only the mean of the final 5 minutes of the test was used for analysis to ensure steady state measures. Face masks (Hans Rudolph, Kansas City, MO, USA) with small dead spaces (approximately 70 mL) were fitted to participants allowing simultaneous breathing at the mouth and nose. Subjects performed a 10-minute, self-selected warm-up before the test followed by a 5-minute self-selected warm-down.

Ongoing monitoring

All of the subjects were monitored daily for morning heart rate, BP, body weight, oxygen saturation and subjective perception of fatigue, muscle soreness, sleep, stress and training performance. The Lake Louise Acute Mountain Sickness Score and training intensity and volume were recorded throughout the camp.

Training

Training loads were calculated via the training impulse (TRIMP) method [24], which was expressed as product of stress (duration of activity) and strain (a 5-point Likert-type scale based on exercise heart rate; easy=1, steady=2, moderately hard=3, hard=4, very hard=5). The heart rate training zones were identified initially from laboratory based cycling and running lactate tests, and confirmed or adjusted by training sessions performed in the field prior to the altitude training camp. The following training zones were established; easy, corresponded to heart rates where lactate was ≤ resting lactate...
Data analysis

All data are means and standard deviations (SD) within and between groups and subjects. We used the baseline data as a covariate in the analysis to adjust for any differences between groups at baseline. We used a contemporary statistical approach because small performance changes can be beneficial for elite athletes [25], whereas conventional statistics can be less sensitive to such small but worthwhile changes. Specifically, we used magnitude-based inferences about effect sizes, and then to make inferences about true (population) values of the effect, the uncertainty in the effect was expressed as 90% confidence limits (CL). The probability that the true value of the effect was practically negative, trivial, or positive accounted for the observed difference, and typical error of measurement [26]. The natural logarithm of each measure was analysed to reduce any effects from non-uniformity errors and then back-transformed. For the physiological variables, the value was determined by multiplying the baseline between-subject standard deviation by Cohen’s value of the smallest worthwhile effect of 0.2 [27]. The unequal variances t statistic was used to analyse differences in the mean change between groups. The difference in the proportion of responders taking iron supplementation and having Acute Mountain Sickness were analysed using the general linear modelling procedure (Proc Genmod version 8.2, SAS Institute Inc., Cary, N.C.) and given as relative risk. The smallest worthwhile change in 800-m performance was assumed to be a reduction or increase in performance time of more than 1.0%, based on previous research into elite athletes competitive performance [23].

Results

Based on any improvement (change) in the sea level 800-m, swim time trial, all of the subjects were retrospectively grouped as responders and non-responders (Figure 1). Responders’ performance improved by 3.2 ± 2.2%, mean ± SD and non-responders performance decreased by 1.8 ± 1.2%, therefore compared to non-responders, the responders improved performance on average by 5.0% (90% CL ± 2.2%) as a result of the 20-day altitude training camp.

Physiological variables

Heart rate and SpO2: Compared to non-responders, responders heart rate at the end of the 5 and 10 minute periods of the submaximal cycle test as a result of the altitude camp were lower (-3.8 ± 4.4% and -6.3 ± 7.8%, mean ± 90% CL, respectively). Relative to the non-responders, the responders also showed a substantial increase in SpO2 at the start (0 minute 2.1 ± 1.7, mean ± 90% CL) and at the end of the 10 minute submaximal cycle test (1.2 ± 1.3%). Responders SpO2 at the end of the 5th minute of the cycle was elevated compared to non-responders but was not substantially different (Table 2). Compared to the responders, the non-responders had substantially higher V E and V E/ V O2 (ventilatory equivalent) measures during the submaximal test on day 20 of the camp (Table 3). As a result of the 20-day altitude training camp, the economy (i.e. power (Watts) per litre of oxygen consumed) of the non-responders compared to the responders deteriorated (i.e. non-responders required more oxygen per watt of work completed). Compared to non-responders, responders showed overall improvement SpO2 per wattage during the 10 minute submaximal cycle test (responders 3.6 ± 1.5% and non-responders 1.4 ± 1.2% mean ± SD).

Iron supplementation: Interestingly, 4 of the 6 responders who improved their swimming performance were taking iron tablets during the altitude camp while 1 of the 4 non-responders were taking iron. In other words, those participants that underwent iron supplementation were 4.0 times (90% CL=1.3-12.8) more likely to improve performance than participants who did not take the supplement.

Subjective variables

Compared to non-responders, responders were more likely to have a lower RPE score before (0 min) and after (end of 5 min recovery) the 10-min submaximal cycle tests (Table 3). Changes to RPE score during the 10-min cycle test were unclear.

There were substantial differences in morning SpO2 at day 1, 2 and 3 between responders and non-responders (1.1 ± 2.0 %, 1.0 ± 1.1 % and 0.8 ± 0.8 %, mean ± 90% CL, respectively). The morning heart rates were not substantially different between groups. The body weights in non-responders tended to decline from day 1 to day 6 (no substantial difference from day 1) of altitude camp, and then plateau. Compared to responders, non-responders body weights were
Chances that true differences are substantial\(a\)

An unexpected but potentially clinically useful finding was that non-responders had consistently higher submaximal exercise RPE scores compared to responders (Table 4).

Substantially higher at day 1, 2 and 3 \((5.8 \pm 7.6\) kg, \(6.0 \pm 7.5\) kg, \(6.0 \pm 7.6\), mean \(\pm 90\%\) CL) of camp (Figure 2).

Subjective scores for fatigue, muscle soreness, the Lake Louise Acute Mountain Sickness (LL-AMS) score and training performance are presented in Figure 3. The scores for fatigue and muscle soreness in non-responders showed mostly higher scores from day 1 to day 20 compared to responders during the camp. For the LL-AMS, non-responders were 3.0 times (90% CL=0.5-16.6) more likely to suffer symptoms of AMS during first 5 days of altitude compared to responders. Specifically, at day 3 and 5, the mean AMS score in non-responders showed mostly higher scores from day 1 to day 20 of training, especially different between responders and non-responders over the course of the training (day 20 minus day 1). Based on the smallest substantial change of 0.2 for all measures.

Discussion

This study had two main findings. First, in response to the 20-day camp utilizing a LHTL protocol, responders had a substantially higher exercise \(\mathrm{SpO}_2\) and lower exercise HR during a submaximal cycle test at altitude. Second, based on the Lake Louise Acute Mountain Sickness Score and the subjective perception scores of fatigue and muscle soreness during altitude training, it seems that responders coped with the LHTL protocol better than non-responders. An unexpected but potentially clinically useful finding was that non-responders had consistently higher submaximal exercise RPE scores compared to responders (Table 4).

It is obvious from our results that the LHTL strategy is not effective for all athletes trying to improve sea-level performance. The mechanisms that contribute to individual variability in responders and non-responders are difficult to elucidate. Large individual variations in physiological responses to hypoxia have been reported, including, erythropoietin release [7] and maximum oxygen consumption [5]. Previous research has suggested a possible relationship between performance variation and altitude training, as differences in gene expression [11,13] and may also relate to different stress thresholds within athletes [14]. Such variation may create an environment where some athletes might find the demand of coping with altitude in addition to normal training too stressful and maladaptation may ensue, while others may thrive, giving rise to the performance variation observed.

Physiological variables

Our study found responders showed a substantial improvement in \(\mathrm{SpO}_2\) at 0 min (or just prior to the test after 5 min standardised warm up) and in the last minute of submaximal cycle test compared to non-responders. In fact, during the submaximal cycle test, all athletes decreased their \(\mathrm{SpO}_2\) to varying degrees which underscores the individual variability in physiological variables among elite athletes. Some researchers have reported significant differences in arterial oxyhaemoglobin saturation which contributed to significant improvement in exercise performance with LHTL (1800-1900 m) via supplemental oxygen [28]. Similar findings have been noted among climbers after a 21-day expedition to 6,194 m [29]. However, these two studies did not separate responders from non-responders. Chapman et al. reviewed the evidence for individual responses to LHTL training and found that athletes that desaturate more during

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**Table 2:** Mean changes in physiological measures during a 10-min submaximal cycle test (day 20 minus day 1) and chances that the differences represent real changes.

<table>
<thead>
<tr>
<th>Physiological Measure</th>
<th>Responders (mean ± SD)</th>
<th>Non-Responders (mean ± SD)</th>
<th>Difference (mean ± 90% CL)</th>
<th>Chances that true differences are substantial (Qualitative inference)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Body weight</td>
<td>1.3 ± 0.6</td>
<td>0.3 ± 2.9</td>
<td>1.9 ± 6.4</td>
<td>42 Unclear</td>
</tr>
<tr>
<td>HR 0 min</td>
<td>4.8 ± 6.6</td>
<td>10.3 ± 14.7</td>
<td>-6.9 ± 18.4</td>
<td>68 Unclear</td>
</tr>
<tr>
<td>HR 5 min</td>
<td>-4.7 ± 3.9</td>
<td>-1.0 ± 10.6</td>
<td>-3.8 ± 4.4</td>
<td>86 Likely</td>
</tr>
<tr>
<td>HR 10 min</td>
<td>-2.5 ± 8.5</td>
<td>3.3 ± 12.6</td>
<td>-6.3 ± 7.8</td>
<td>87 Likely</td>
</tr>
<tr>
<td>(\mathrm{SpO}_2) 0 min</td>
<td>1.5 ± 2.8</td>
<td>-0.8 ± 2.0</td>
<td>2.1 ± 1.7</td>
<td>95 Very likely</td>
</tr>
<tr>
<td>(\mathrm{SpO}_2) 5 min</td>
<td>4.0 ± 3.8</td>
<td>2.3 ± 0.9</td>
<td>1.4 ± 2.4</td>
<td>73 Unclear</td>
</tr>
<tr>
<td>(\mathrm{SpO}_2) 10 min</td>
<td>3.3 ± 1.5</td>
<td>2.3 ± 0.5</td>
<td>1.2 ± 1.3</td>
<td>83 Likely</td>
</tr>
</tbody>
</table>

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**Table 3:** Mean changes in respiratory variables over the last 5 minutes of a 10-min submaximal cycle test conducted on day 1 and day 20 of altitude training.

<table>
<thead>
<tr>
<th>Respiratory Measure</th>
<th>Responders (mean ± SD)</th>
<th>Non-Responders (mean ± SD)</th>
<th>Difference (mean ± 90% CL)</th>
<th>Chances that true differences are substantial (Qualitative inference)</th>
</tr>
</thead>
<tbody>
<tr>
<td>(\dot{V}E) ((L/min^-1))</td>
<td>68.5 ± 19.9</td>
<td>79.6 ± 24.3</td>
<td>93.1 ± 16.6</td>
<td>95.5 ± 30.2</td>
</tr>
<tr>
<td>(\dot{V}O_2) ((L/min^-1))</td>
<td>1.7 ± 0.5</td>
<td>2.0 ± 0.7</td>
<td>1.3 ± 0.6</td>
<td>2.6 ± 0.7</td>
</tr>
<tr>
<td>RER</td>
<td>0.94 ± 0.02</td>
<td>0.84 ± 0.03</td>
<td>0.95 ± 0.02</td>
<td>0.87 ± 0.04</td>
</tr>
<tr>
<td>(\dot{V}E/\dot{V}O_2) ((L/L^-1))</td>
<td>40.2 ± 14.1</td>
<td>40.8 ± 9.9</td>
<td>76.8 ± 38.8</td>
<td>52.5 ± 13.3</td>
</tr>
<tr>
<td>Economy ((Watt/L^-1))</td>
<td>132.2 ± 42.2</td>
<td>127.3 ± 40.7</td>
<td>209.6 ± 86.3</td>
<td>94.2 ± 17.2</td>
</tr>
</tbody>
</table>
exercise (at sea level or altitude) tend to have difficulty in adapting to altitude training while those that maintain adequate oxygen saturation respond better [6]. The relative increase (compared to non-responders) in the current study in SpO2 in responders (which corroborates the theory of Chapman et al. could be explained in part by increased haemoglobin concentration [30], enhanced erythropoietic response [7,31], which may result in increased red cell volume [4] and subsequently increased arterial oxygen content. Weil et al. noted a graded response such that an increase in red cell mass was proportional to oxyhaemoglobin saturation at sea level and altitude [32]. This may suggest a haematological aspect is involved in the beneficial adaptation to altitude. The chronic hypoxic stress facilitates increased red cell production (when adequate iron level is available) which leads to an increase haemoglobin available to carry oxygen to muscle cells (and subsequently increase SpO2).

Relative to non-responders, the heart rate of responders at 5 min and 10 min during the submaximal cycling test were substantially
lower. In addition, responders had lower $\dot{V}E$, RER, $\dot{V}E/\dot{V}O_2$, but better economy (post training camp) and higher relative $\dot{V}O_2$ to wattage ratio than non-responders. This may imply that responders have a greater ability to maintain training workload during altitude camps. These findings are consistent with the results of the Chapman et al. study, where individual variability in the response to altitude training was accounted for by maintenance of training intensity and oxygen uptake values [5]. Previous work has also found similar improvements in exercise economy after altitude training based at higher (2500-3500 m) and lower (1200 m) altitudes [22]. We suggest that the higher exercise $\dot{V}O_2$ in responders reflects higher arterial partial pressure of oxygen. Responders are therefore able to train at higher intensities without delving into anaerobic energy systems and subsequently fatigue and therefore acquire performance improvement. The non-responders on the other hand, cannot maintain adequate oxygen levels in the blood resulting in lower $\dot{V}O_2$ causing an increase in heart rate and $\dot{V}E$. In addition, the lower levels of oxygen require a higher reliance on anaerobic metabolic processes, thereby elevating RER. The other possible cause for lower heart rates in responders during submaximal exercise may be due to increased vagal dominance [14], or to the elevation of red cell volume, stroke volume, and ultimately cardiac output, which subsequently decreases heart rate. A recent study demonstrated that red cell volume remained elevated in responders after 2 weeks of altitude training, while non-responders red cell volume returned to their original levels [4]. However, such linkages between heart rate changes and response to altitude require further research to confirm findings.

Subjective variables

We observed that non-responders found the submaximal exercise test subjectively harder (high RPE score) than responders. Svedenhag et al. found that 2 weeks of altitude (2000 m) training improved RPE during exercise compared to sea-level training [33] but there are currently no studies comparing the perceived exertion of responders with non-responders. Bailey et al. also reported that 4 weeks of altitude training (1500-2000 m) improved RPE when performing a submaximal exercise test at sea-level in elite distance runners [34]. A high correlation between RPE and HR was reported by Borg et al. in a variety of work tasks (cycling and treadmill) and under varying exercise conditions (moderate to heavy intensity) [35]. It seems a major influence on RPE scores is the build-up of blood lactate and hydrogen ions due to insufficient aerobic processes to meet the demands of heavy exercise [36]. We postulate that non-responders are required to meet the energy demands of exercise at altitude through greater anaerobic means thus pushing up RPE, heart rate and RER, however, this hypothesis is speculation as these measures were not taken in this study and further studies are required to substantiate this argument.

Acute mountain sickness (AMS) and the subjective perception score on fatigue, muscle soreness and training performance demonstrated that non-responders were not acclimatizing to the extra stress of training and hypoxia to the same extent as responders. Fatigue and muscle soreness scores indicated more stress after 24-h exposure to altitude (day 1) in non-responders compared to responders. Overall non-responders had a higher daily score on most days of the altitude training camp in all of the subjective parameters. Previous research suggested that AMS sufferers have lower resting pain thresholds and subsequently increased exercise-induced muscle soreness which was caused by free-radical-mediated skeletal muscle damage [37]. In short, low AMS and subjective perception scores of fatigue and muscle soreness could be a sign of those who adapt well to moderate altitude. It seems the responders adapt to extra stress of training at altitude whereas the non-responders find the extra stress of training and altitude too much and subsequently maladapt. Such changes in these subjective measures of participants may be useful for coaches as a detection mechanism for those athletes not coping with the altitude and training.

Reasons behind the varied performance response to LIHTL training are probably multi-factorial however genetics may play a large part [38]. Studies have shown that a transcriptional factor, called a hypoxia-inducible factor (HIF), is essential in promoting cellular adaptation to changes in oxygen availability and regulating the hypoxic gene expression [39]. Under hypoxic conditions, the HIF-1α complex is stable, allowing for transcriptional activation and ultimate stimulation of proteins such as EPO [40] and vascular endothelial growth factor [41]. HIF-1α also moderates other physiological responses at altitude including glucose transport and glycolytic enzyme activity [42]. Conversely, the studies of Mason et al. (2004) and (2007) showed that under low-oxygen conditions glycolysis is the central source of anaerobic energy which is regulated by HIF-1α. It can be seen in HIF-1α knockout mice that loss of HIF-1α in skeletal muscle causes an adaptive response by shifting from glycolytic metabolism toward fatty acid oxidation leading to an increased capacity for endurance exercise [43,44]. We speculate that the non-responders in this study probably sensitive to the hypoxic condition consequently produced more HIF-1α compared to responders which may have increased the reliance on glycolysis and subsequently increased RER during exercise. However, more research is required in this area since other researchers have been unable to identify
genes that can be overwhelmingly linked to the individual variability observed in responders versus non-responders [10] and HIF-1α gene expression quantification is unable to predict poor and good responder-athletes to the LHTL model [11].

From this research it seems clear that athletes who respond to altitude are less stressed in terms of lower heart rate, \( V \) \( E \) and higher levels of arterial oxygen saturation during submaximal exercise. Athletes that are not responding seem to perceive themselves as being more fatigued, become less efficient (use more oxygen per watt), have more muscle soreness, and higher AMS scores, particularly over the first few days of altitude. It seems the non-responders are unable to adapt to the increased demand for oxygen required by the muscles during exercise at altitude, resulting in fatigue and poor performance adaptation. Possible confounding factors behind athletes who respond to altitude training and those that do not may be their iron stores. Clearly, there is the need for further studies to investigate possible mechanisms.

A limitation of this study is lack of a sea-level control group which does not allow us to distinguish whether the observed effects are due to the altitude or to the effect of training. Indeed, altitudes that are more common for such training (e.g. 2000-2500 m) may have been better at separating out clear differences between responders and non-responders, but such altitude training bases are unavailable in New Zealand. In addition, maximal testing (run or cycle) may have provided greater evidence for differences between the groups. However, we consider our observations are worthwhile given the lack of information on such groups (elite triathletes all ranked within the world top 100). Nevertheless, these results should remain speculative until further research is conducted in a more controlled study.

Conclusion

This investigation demonstrated that changes in some simple measures taken during submaximal exercise (SpO2, HR, \( V \) \( E \), economy and RPE) or at rest (perception of fatigue, muscle soreness, training performance and the acute mountain sickness score) during an altitude sojourn may be useful indicators for classifying athletes that are coping and are likely to improve subsequent sea-level performance from those that are suffering increased stress and are likely to decrease subsequent sea-level performance. These physiological indicators and subjective variables provide non-invasive screening for athletes who could profit from such hypoxic exposure.

Acknowledgements

The authors acknowledge the Lincoln University Research Fund for financial support, the New Zealand Triathlon athletes and their coaches, and the Snow Farm in Wanaka for their assistance.

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