

2017

# Changes in appetite, energy intake, body composition and circulating ghrelin constituents during an incremental trekking ascent to high altitude

Jamie Matu

John O'Hara

Neil Hill

Sarah B. Clarke

*Northern Michigan University*

Christopher Boos

*See next page for additional authors*

Follow this and additional works at: [https://commons.nmu.edu/facwork\\_journalarticles](https://commons.nmu.edu/facwork_journalarticles)



Part of the [Exercise Science Commons](#), and the [Sports Sciences Commons](#)

---

## Recommended Citation

Matu, Jamie; O'Hara, John; Hill, Neil; Clarke, Sarah B.; Boos, Christopher; Newman, Caroline; Holdsworth, David; Ispoglou, Theocharis; Duckworth, Lauren; Woods, David; Mellor, Adrian; and Deighton, Kevin, "Changes in appetite, energy intake, body composition and circulating ghrelin constituents during an incremental trekking ascent to high altitude" (2017). *Journal Articles*. 360. [https://commons.nmu.edu/facwork\\_journalarticles/360](https://commons.nmu.edu/facwork_journalarticles/360)

This Journal Article is brought to you for free and open access by the FacWorks at NMU Commons. It has been accepted for inclusion in Journal Articles by an authorized administrator of NMU Commons. For more information, please contact [kmcdonou@nmu.edu](mailto:kmcdonou@nmu.edu), [bsarjean@nmu.edu](mailto:bsarjean@nmu.edu).

---

**Author(s)**

Jamie Matu, John O'Hara, Neil Hill, Sarah B. Clarke, Christopher Boos, Caroline Newman, David Holdsworth, Theocharis Ispoglou, Lauren Duckworth, David Woods, Adrian Mellor, and Kevin Deighton

1 **TITLE:** Changes in appetite, energy intake, body composition and circulating ghrelin constituents during an  
2 incremental trekking ascent to high altitude

3 **AUTHOR NAMES:** Jamie Matu<sup>1</sup>, John O'Hara<sup>1</sup>, Neil Hill<sup>2,4</sup>, Sarah Clarke<sup>1</sup>, Christopher Boos<sup>1,3</sup>, Caroline  
4 Newman<sup>4</sup>, David Holdsworth<sup>4</sup>, Theocharis Ispoglou<sup>1</sup>, Lauren Duckworth<sup>1</sup>, David Woods<sup>1,4</sup>, Adrian Mellor<sup>1,4</sup>, and  
5 Kevin Deighton<sup>1</sup>

6 **DEPARTMENT AND INSTITUTION:**

7 <sup>1</sup>Institute for Sport Physical Activity & Leisure, Leeds Beckett University, Leeds, UK

8 <sup>2</sup>Section of Investigative Medicine, Imperial College London, London, UK

9 <sup>3</sup>Poole Hospital NHS Trust, Longfleet Rd, Poole, UK

10 <sup>4</sup>Royal Centre for Defence Medicine, ICT Building, Vincent Drive, Birmingham, UK

11 **CORRESPONDING AUTHOR:** Dr Kevin Deighton, Institute for Sport Physical Activity & Leisure, Leeds  
12 Beckett University, Leeds, LS6 3QS, United Kingdom (email: [K.Deighton@leedsbeckett.ac.uk](mailto:K.Deighton@leedsbeckett.ac.uk)). ORCID: 0000-  
13 0001-7994-2137.

14 **TELEPHONE NUMBER:** +44 (0)113 8123582

15

16

17

18

19

20

21

22 **ABSTRACT**

23 Purpose: Circulating acylated ghrelin concentrations are associated with altitude-induced anorexia in laboratory  
24 environments, but have never been measured at terrestrial altitude. This study examined time course changes in  
25 appetite, energy intake, body composition and ghrelin constituents during a high altitude trek. Methods: Twelve  
26 participants (age: 28(4) years, BMI: 23.0(2.1)kg.m<sup>-2</sup>) completed a 14-day trek in the Himalayas. Energy intake,  
27 appetite perceptions, body composition, and circulating acylated, des-acylated and total ghrelin concentrations  
28 were assessed at baseline (113m; 12 days prior to departure) and at three fixed research camps during the trek  
29 (3619m, day seven; 4600m, day 10; 5140m, day 12). Results: Relative to baseline, energy intake was lower at  
30 3619m (P=0.038) and 5140m (P=0.016) and tended to be lower at 4600m (P=0.056). Appetite perceptions were  
31 lower at 5140m (P=0.027) compared with baseline. Acylated ghrelin concentrations were lower at 3619m  
32 (P=0.046) and 4600m (P=0.038), and tended to be lower at 5140m (P=0.070), compared with baseline. Des-  
33 acylated ghrelin concentrations did not significantly change during the trek (P=0.177). Total ghrelin  
34 concentrations decreased from baseline to 4600m (P=0.045). Skinfold thickness was lower at all points during the  
35 trek compared with baseline (P≤0.001) and calf girth decreased incrementally during the trek (P=0.010).  
36 Conclusions: Changes in plasma acylated and total ghrelin concentrations may contribute to the suppression of  
37 appetite and energy intake at altitude but differences in the time course of these responses suggests that additional  
38 factors are also involved. Interventions are required to maintain appetite and energy balance during trekking at  
39 terrestrial altitudes.

40 **KEYWORDS:** ghrelin; hypoxia; altitude-induced anorexia; terrestrial altitude

41

42 **Abbreviations**

AG:DG	Acylated ghrelin to des-acylated ghrelin ratio
AMS	Acute mountain sickness
ANOVA	Analysis of variance
BMI	Body mass index
CAS	Composite appetite score
GOAT	Ghrelin-O-acyltransferase
ISAK	International Society for the Advancement of Kinanthropometry
LLS	Lake Louise Score
MoDREC	Ministry of Defence Research Ethics Committee
RPE	Rating of perceived exertion
SD	Standard deviation
SE	Standard error
SpO <sub>2</sub>	Arterial oxygen saturations
VAS	Visual analogue scales

43

44

## 45 INTRODUCTION

46 Acute exposure to hypoxic environments has been demonstrated to suppress appetite and energy intake (Armellini  
47 et al. 1997; Matu et al. 2017; Wasse et al. 2012; Westerterp et al. 2000). This effect appears to be maintained  
48 during prolonged sojourns to high altitude, which results in significant decreases in body mass, of which greater  
49 than 50 % is from fat-free mass (Rose et al. 1988; Sergi et al. 2010). These declines in lean mass will likely lead  
50 to a drop in physical capabilities at altitude (Sergi et al. 2010), which can have deleterious implications for  
51 individuals ascending to high altitude. A better understanding of the time course of these changes during a trek,  
52 as well as the mechanisms involved, is required to develop guidance for those travelling to high altitudes.

53 Over the past twenty years, changes in the circulating concentrations of gastrointestinal hormones and  
54 leptin have been implicated as potential mechanisms for the alterations in appetite and energy intake at altitude.  
55 However, although several hormones such as pancreatic polypeptide (Riepl et al. 2012), leptin (Sierra-Johnson et  
56 al. 2008), glucagon-like-peptide-1 (Snyder et al. 2008) and total ghrelin (Benso et al. 2007; Riepl et al. 2012;  
57 Shukla et al. 2005) have been measured in response to terrestrial altitude exposure, the findings remain equivocal.  
58 One major limitation of the current research is the measurement of total ghrelin concentrations at altitude, rather  
59 than the constituent components of acylated and des-acylated ghrelin which have opposing effects on appetite  
60 regulation (Fernandez et al. 2016). The differentiation of ghrelin constituents in response to terrestrial altitude is  
61 imperative as acylated ghrelin has been found to be particularly responsive to hypoxic exposure in a laboratory  
62 environment with decreases in this hormone correlated with a reduction in appetite (Bailey et al. 2015) and energy  
63 intake (Wasse et al. 2012). Furthermore, recent evidence suggests that des-acylated ghrelin may inhibit the  
64 orexigenic effects of acylated ghrelin (Fernandez et al. 2016), which further emphasises the need to measure both  
65 hormones as well as the ratio between the two (Al Massadi et al. 2014). It seems feasible that the measurement of  
66 total ghrelin in previous research (Benso et al. 2007; Debevec et al. 2014; Riepl et al. 2012) may have masked  
67 changes in acylated and des-acylated ghrelin, which may explain the lack of association between changes in  
68 appetite and circulating ghrelin concentrations at altitude.

69 Although circulating total ghrelin concentrations have been extensively investigated in response to  
70 hypoxic exposure (Benso et al. 2007; Debevec et al. 2014; Debevec et al. 2016; Mekjavic et al. 2016; Riepl et al.  
71 2012; Shukla et al. 2005), the investigation of acylated ghrelin is currently limited to four studies, all of which  
72 lasted for  $\leq 7$  h and were all conducted in normobaric environments (Bailey et al. 2015; Matu et al. 2017;  
73 Morishima and Goto 2016; Wasse et al. 2012). Although laboratory studies of this nature are valuable to gain

74 greater mechanistic understanding, further field studies are required to assess the combined effects of trekking,  
75 gradual ascent and other environmental stimuli such as cold exposure which occur during real life ascent to high  
76 altitude. The measurement of acylated and des-acylated ghrelin during ascent to terrestrial altitude is vital to  
77 understand the changes that occur during a real-world environment and the importance of these changes as a basis  
78 for the development of future interventions. The lack of investigation into the constituents of total ghrelin to date  
79 is likely due to the complexities of the necessary chemical preparation required to prevent the degradation of the  
80 analytes (Hosoda et al. 2004), which is particularly difficult to achieve in an extreme field environment.

81 The purpose of this study was to investigate the effects of a high altitude trek to 5300 m on appetite,  
82 energy intake and body composition responses in healthy men and women, with a further focus on circulating  
83 acylated and des-acylated ghrelin concentrations as mechanistic variables. These data provide novel insights into  
84 the time course of changes in appetite, energy intake and body composition during a real-life ascent to high  
85 altitude. This study also provides a better understanding of the mechanisms responsible for altitude-induced  
86 anorexia, representing the first investigation of acylated ghrelin and des-acylated ghrelin at terrestrial altitude. We  
87 hypothesised that exposure to increasingly high altitudes would suppress appetite, circulating acylated ghrelin  
88 concentrations and energy intake, which would be associated with a reduction in lean and total body mass.

89

## 90 METHODS

91 **Participants.** This study was conducted according to the guidelines laid down in the Declaration of  
92 Helsinki and all procedures were approved by the Ethics Advisory Committee at Leeds Beckett University and  
93 the Ministry of Defence Research Ethics Committee (MoDREC; protocol number 624). Twelve members (nine  
94 male, three female) of the British Military volunteered to participate in this study. Informed consent was obtained  
95 from all participants included in the study. All participants were non-smokers, had no known disease, allergies or  
96 intolerances, and had not been to an altitude over 1000 m for at least 3 months. All participants were physically  
97 fit and could run 2.4 km on a treadmill at a 2 % gradient in under 13 minutes 37 seconds in accordance with  
98 military requirements. The physical characteristics of participants (mean (SD)) were as follows: age 28 (4) years,  
99 body mass 71.3 (10.3) kg, body mass index (BMI) 23.0 (2.1) kg.m<sup>-2</sup>.

100 **Study design.** This study represents part of the 'British Services Dhaulagiri Medical Research  
101 Expedition' which took place in March – May 2016 (Mellor et al. 2017). In April 2016, participants in the present  
102 study travelled from the UK to Nepal and completed a 14-day trek around the Dhaulagiri circuit in the Himalayas.  
103 Travel from the UK to Nepal lasted for one day and participants were in Nepal for three days prior to starting the  
104 trek. The trek commenced from Darbang (~1100 m), peaked on day 11 at the French Pass (~5300 m) and ended  
105 on day 14 at Marpha (~2700 m). Pre-planned rest days were included at fixed camps at 3619 m (Camp 1; day  
106 seven), 4600 m (Camp 2; day 10) and 5140 m (Camp 3; day 12). Participants walked a mean distance of 8.2  
107 km.day<sup>-1</sup> with a mean elevation gain of 471 m.day<sup>-1</sup> whilst carrying a day pack weighing ~ 5 kg. Further  
108 information about the ascent profile and trek characteristics has been published elsewhere (Mellor et al. 2017).  
109 Data collection took place at baseline (113m; 12 days prior to departure from the UK) and at each fixed camp. On  
110 the day preceding data collection at camp 1, camp 2 and camp 3 participants walked 4.3 km, 4.3 km and 9.1 km  
111 and gained an elevation of 512 m, 528 m and 540 m, respectively. All trekking on these days was completed by  
112 5pm. All participants wore the same type of clothing throughout the trek and experienced the same degree of cold  
113 exposure. Baseline measurements were collected in the laboratories at Leeds Beckett University and the measures  
114 at each camp were collected in a designated research tent. At the time data was collected ambient temperatures in  
115 the laboratory and research tents were 19.8 °C, 4.9 °C, 1.2 °C and -6.4 °C at baseline, 3619 m, 4600 m and 5140  
116 m, respectively. All participants remained rested on the day of testing. Participants were staggered for the  
117 collection of all measurements between 7 am and 10 am, with each participant having their measures taken at a  
118 consistent time on all occasions and after an overnight fast of at least 10 h.



119           **Food Provision.** Throughout the trek all food and fluid was available *ad libitum*, and was provided by  
120 Nepalese cooks and staff who accompanied the trekking team. Typical foods offered at each meal were as follows:  
121 Breakfast – cereals, porridge, omelette, pancakes; Lunch – noodles, meats, soup, beans, vegetables, fruit; Dinner  
122 – curry, pasta, pizza, potatoes, dumplings, cheese, vegetables; Snacks – chocolate bars, biscuits, cake, fruit. At all  
123 meals participants were given more than one option and thus could decide what they wanted to eat. The mean  
124 (SD) macronutrient composition of the food consumed during the trek was 49.0 (6.6) % carbohydrate, 36.3 (6.2)  
125 % fat and 14.7 (2.6) % protein, respectively.

126           **Food intake.** Energy intake was assessed at baseline via a 24-hour dietary recall interview by an  
127 experienced researcher (Academic Associate of the Sport and Exercise Nutrition Register) using the multiple-pass  
128 approach (Guenther et al. 1997). In addition to collecting dietary intake information, this approach was used to  
129 demonstrate the level of detail required from the participants when completing a food diary during the trek. All  
130 participants completed a food diary on the day preceding each fixed camp and this process was monitored and  
131 verified by the same experienced researcher throughout the trek. Although there are acknowledged limitations of  
132 self-reported dietary intake methods (Hill and Davies 2001), the oversight of food diaries by the researcher present  
133 on the trek ensured accurate completion of all food diaries. Food intake was monitored during the day before each  
134 fixed camp to consider acute dietary changes when interpreting the data for fasted appetite ratings and blood  
135 samples at the fixed camps. The palatability of each meal consumed was measured using 100 mm visual analogue  
136 scales (VAS) with the anchors “the worst taste that I have ever experienced” and “the best taste that I have ever  
137 experienced” at each end of the scale. A mean palatability score was calculated for each day to control for any  
138 influence of palatability on food consumption.

139           **Appetite.** Appetite perceptions were measured after an overnight fast at baseline and upon waking at  
140 each fixed camp using validated 100 mm VAS for hunger, satisfaction, fullness and prospective food consumption  
141 (PFC) (Flint et al. 2000). Using these scales, a composite appetite score (CAS) was calculated using the following  
142 formula: composite appetite score = ([hunger + prospective food consumption + (100 – fullness) + (100 –  
143 satisfaction)] / 4) (Stubbs et al. 2000). A higher value is associated with a greater appetite sensation and  
144 subsequently a stronger motivation to eat. In addition, the extent to which participants desired sweet, salty,  
145 savoury, and fatty foods was assessed using VAS anchored at each end with “yes, very much” and “no, not at all”.

146           **Acute Mountain Sickness, oxygen saturation and rating of perceived exertion.** Acute mountain  
147 sickness (AMS) was assessed every morning and evening using the Lake Louise AMS (LLS) score (Roach et al.

148 1993); mild AMS was defined as LLS of  $\geq 3$  in the presence of a headache and severe AMS was defined as LLS  
149 of  $\geq 6$  in the presence of a headache. Arterial oxygen saturations ( $SpO_2$ ) were measured via a fingertip pulse  
150 oximeter (Nellcor™ PM10N; Medtronic, Minneapolis, MN) every morning and evening while the participants  
151 were resting in a seated position. Rating of perceived exertion (RPE) (Borg 1982) was recorded as the hardest  
152 exertion experienced during the day of trekking preceding each fixed camp, as used previously in similar  
153 environments (Mellor et al. 2014).

154 **Body composition.** Participants were weighed at baseline and each fixed camp in a fasted state whilst  
155 wearing minimal clothing and no footwear. A portable multicomponent force plate (Kistler, Switzerland) was  
156 used and was stabilised on the mountain using specialised levelling feet (JVD Design & Automation Ltd, Leeds,  
157 UK). Skinfolds of the triceps, subscapular, biceps, iliac crest, supraspinale, abdominal, front thigh and medial calf  
158 were measured using calibrated Harpenden callipers (John Bull, British Indicators, West Sussex, UK) to the  
159 nearest 0.1 mm. The sum of skinfolds was calculated by the addition of each of the eight skinfold values in mm.  
160 Girth measurements of the waist and calf, as well as the upper arm in a relaxed and flexed state, were obtained  
161 using a steel anthropometric tape (Lufkin W606PM, Cooper Hand Tools, Tyne & Wear, UK) to the nearest 1 mm.  
162 All anthropometric assessments were conducted by one researcher who was trained by an individual accredited  
163 by the International Society for the Advancement of Kinanthropometry (ISAK). All measures were conducted in  
164 duplicate and in accordance with ISAK guidelines on the right side of the body. The coefficient of variation for  
165 skinfolds and girths was 2.2% and 0.4%, respectively.

166 **Blood sampling.** Venous blood samples were obtained from an antecubital vein via venepuncture using  
167 a 21-gauge butterfly needle (Safety-Lok™; BD, Oxford, UK). Samples were collected at baseline and at all  
168 research camps with participants in a fasted state. One 4.9 mL pre-cooled EDTA monovette (Sarstedt, Leicester,  
169 UK) was used to obtain samples for the determination of plasma acylated and des-acylated ghrelin concentrations.  
170 Monovettes were pre-treated on the morning of testing, to prevent the degradation of acylated ghrelin, with 50 $\mu$ l  
171 of a solution containing p-hydroxymercuribenzoic acid, potassium phosphate buffer and sodium hydroxide  
172 (Hosoda et al. 2004). Immediately after filling, the tube was spun at 1500 x g for 10 minutes in a centrifuge  
173 (CompactStar CS4, VWR). Subsequently 1 mL of plasma was mixed with 100 $\mu$ l of 1M hydrochloric acid. This  
174 solution was then immediately frozen at either -20°C in a freezer (for baseline measurements) or within a dry  
175 shipper containing liquid nitrogen at <-80°C (at each fixed camp) before being transferred to a -80°C freezer at  
176 the university and stored until analysis. Plasma volume changes as a result of altitude exposure were not assessed  
177 during the trek because it is the absolute plasma hormonal concentrations that would determine the body's

178 response at that specific time (Kargotich et al. 1998). However, to prevent any extraneous influences from postural  
179 changes, all blood samples were collected after the participant had been seated for at least 5 minutes (Fawcett and  
180 Wynn 1960).

181 **Blood analyses.** Commercially available enzyme immunoassays were used to determine plasma  
182 concentrations of acylated and des-acylated ghrelin (SPI BIO, Montigny Le Bretonneux, France). To eliminate  
183 interassay variation, all samples from each participant were analysed on the same plate. In addition, all samples  
184 were analysed in duplicate and on the same day. The within batch coefficients of variation were 2.5% for acylated  
185 ghrelin and 2.4% for des-acylated ghrelin. Total ghrelin was computed via the addition of acylated and des-  
186 acylated ghrelin concentrations. The ratio between acylated ghrelin and des-acylated ghrelin concentrations  
187 (AG:DG) was calculated as acylated ghrelin divided by des-acylated ghrelin, as previously described (Delhanty  
188 et al. 2015).

189 **Statistical analysis.** Data are expressed as mean (SD) in text and tables and mean (SE) in figures to  
190 avoid distortion of the graphs. Diet records were inputted into Nutritics dietary analysis software (v1.8 for  
191 Windows; Nutritics, Dublin) to assess energy intake. All data were analysed using IBM SPSS statistics (v22.0 for  
192 Windows; SPSS, Chicago, IL). One way repeated measures analysis of variance (ANOVA) was used to assess  
193 altitude-based differences in SpO<sub>2</sub>, AMS scores, RPE scores, body composition measures, appetite perceptions,  
194 energy intake, fluid intake and plasma ghrelin concentrations. Significant effects were further explored using  
195 Student's paired *t* tests. Effect sizes are presented as Cohen's *d* and interpreted as  $\leq 0.2$  trivial,  $> 0.2$  small,  $> 0.6$   
196 moderate,  $> 1.2$  large,  $> 2$  very large and  $> 4$  extremely large (Hopkins 2004). The Pearson product moment  
197 correlation coefficient was used to investigate relationships between variables at each altitude. The exclusion of  
198 participants reporting AMS did not alter the interpretation of the findings; subsequently all participants were  
199 included in the data analysis. Based on evidence that males and females exhibit similar appetite, energy intake  
200 and gut hormone responses to exercise- and diet-induced energy deficits (Alajmi et al. 2016), data from both  
201 genders were combined for analyses. The sample size used within this study was deemed sufficient to detect a  
202 significant difference in energy intake between altitudes. The anticipated effect size for a difference in energy  
203 intake was based on a similar previous study which investigated energy intake in individuals climbing at  
204 approximately 4500 m for 16 days (Armellini et al. 1997). Based on the effect size and an alpha value of 5 %, a  
205 sample size of 12 participants would generate a power  $>95$  %. Calculations were performed using G\*power (v3  
206 for Windows; Düsseldorf) (Faul et al. 2007).

207 **RESULTS**

208 Measurements of SpO<sub>2</sub>, AMS, RPE, body composition, fluid intake and energy intake were successfully obtained  
209 from all 12 participants. One male participant withdrew consent for blood sampling at the research camps during  
210 the trek and one male participant did not complete the appetite perception measurements during the trek.  
211 Therefore, data is presented for 11 participants for plasma hormone concentrations and appetite perceptions.

212

213 **Oxygen saturations, acute mountain sickness and rating of perceived exertion.** One way ANOVA  
214 revealed a main effect of altitude for SpO<sub>2</sub> ( $P < 0.001$ ). Post-hoc analysis demonstrated lower SpO<sub>2</sub> at each fixed  
215 camp compared with the previous location (baseline: 98.4 (0.9) %; 3619 m: 92.4 (2.5) %,  $P < 0.001$ ,  $d = 3.19$ ;  
216 4600 m: 83.5 (4.1) %,  $P < 0.001$ ,  $d = 2.62$ ; 5140 m: 79.8 (5.6) %,  $P = 0.007$ ,  $d = 0.75$ ). A positive diagnosis of  
217 mild AMS was reported in 50% of participants at some point during the trek. The first incidence of AMS occurred  
218 on the eighth day of the trek (the day after the first rest day; 4072 m). Incidence of AMS at the four fixed locations  
219 was as follows: baseline: zero participants; 3619 m: zero participants; 4600 m: two participants; and 5140 m: two  
220 participants. One way ANOVA revealed a significant effect of altitude for RPE ( $P < 0.001$ ). Relative to 3619 m  
221 (11.8 (1.5)), RPE was significantly higher at 4600 m (13.3 (1.5),  $P = 0.009$ ,  $d = 1.01$ ) and relative to 4600 m RPE  
222 was significantly higher at 5140 m (16.5 (2.5),  $P = 0.003$ ,  $d = 1.58$ ).

223

224 **Energy and fluid intake.** One way ANOVA revealed a significant effect of altitude for energy intake  
225 ( $P = 0.015$ ). Relative to baseline, energy intake was significantly lower at 3619 m ( $P = 0.038$ ,  $d = 1.05$ ) and 5140  
226 m ( $P = 0.016$ ,  $d = 1.00$ ) and tended to be lower at 4600 m ( $P = 0.056$ ,  $d = 0.82$ ). There were no differences observed  
227 between research camps during the trek (all  $P \geq 0.333$ ,  $d \leq 0.22$ ) (Figure 1a).

228 One way ANOVA revealed a main effect of altitude for fluid intake ( $P = 0.029$ ). Relative to baseline  
229 (2769 (1156) mL.day<sup>-1</sup>) fluid intake was significantly higher at 3619 m (4438 (1847) mL.day<sup>-1</sup>,  $P = 0.008$ ,  $d =$   
230 1.08) and 4600 m (4236 (2120) mL.day<sup>-1</sup>,  $P = 0.027$ ,  $d = 0.86$ ), but not significantly higher at 5140 m (3645 (2026)  
231 mL.day<sup>-1</sup>,  $P = 0.126$ ,  $d = 0.53$ ). There were no differences observed between camps (all  $P \geq 0.266$ ,  $d \leq 0.29$ ).

232 One way ANOVA revealed a main effect of altitude for the daily palatability of food consumed ( $P =$   
233 0.018). Relative to baseline, palatability was significantly higher at 3619 m ( $P = 0.030$ ,  $d = 1.17$ ). However,  
234 palatability was not different at 4600 m ( $P = 0.147$ ,  $d = 0.66$ ) or 5140 m ( $P = 0.509$ ,  $d = 0.32$ ) compared with

235 baseline. Palatability was significantly lower at 5140 m compared with 3619 m ( $P = 0.020$ ,  $d = 0.97$ ) and 4600 m  
236 ( $P = 0.013$ ,  $d = 0.71$ ) (Figure 1b).

237 One way ANOVA revealed a main effect of altitude on the desire to eat salty ( $P = 0.025$ ) and savoury ( $P$   
238  $< 0.001$ ) foods, but not sweet ( $P = 0.604$ ) or fatty ( $P = 0.354$ ) foods. Relative to baseline (67 (12) mm), the desire  
239 to eat salty foods was significantly lower at 3619 m (41 (23) mm,  $P = 0.018$ ,  $d = 1.42$ ) and 5140 m (39 (27) mm,  
240  $P = 0.024$ ,  $d = 1.38$ ), and also tended to be lower at 4600 m (45 (27) mm,  $P = 0.066$ ,  $d = 1.09$ ). There were no  
241 differences observed between camps (all  $P \geq 0.159$ ,  $d \leq 0.22$ ). The desire to eat savoury foods was significantly  
242 increased at 3619 m (70 (13) mm,  $P < 0.001$ ,  $d = 2.41$ ), 4600 m (67 (15) mm,  $P < 0.001$ ,  $d = 2.03$ ) and 5140 m  
243 (53 (19) mm,  $P = 0.011$ ,  $d = 1.08$ ) compared with baseline (33 (18) mm). In addition, the desire to eat savoury  
244 foods reduced significantly from 4600 m to 5140 m ( $P = 0.026$ ,  $d = 0.79$ ) with no difference observed between  
245 3619 m and 4600 m ( $P = 0.516$ ,  $d = 0.23$ ).

246

247 **Appetite perceptions.** One way ANOVA revealed a main effect of altitude for CAS ( $P = 0.005$ ). Post-  
248 hoc analysis revealed that CAS was significantly lower at 5140 m compared with baseline ( $P = 0.027$ ,  $d = 1.07$ ),  
249 3619 m ( $P = 0.005$ ,  $d = 1.19$ ) and 4600 m ( $P = 0.05$ ,  $d = 0.69$ ). No other differences were observed between  
250 altitudes (all  $P \geq 0.116$ ,  $d \leq 0.48$ ) (Figure 1c).

251

252 **Plasma acylated and des-acylated ghrelin concentrations.** One way ANOVA revealed a significant  
253 effect of altitude for plasma acylated ghrelin concentrations ( $P = 0.048$ ; Figure 2a), plasma total ghrelin  
254 concentrations ( $P = 0.047$ ; Figure 2d) and the AG:DG ratio ( $P = 0.046$ ; Figure 2c). A main effect of altitude was  
255 not detected for plasma des-acylated ghrelin concentrations ( $P = 0.177$ ; Figure 2b).

256 Relative to baseline, plasma acylated ghrelin concentrations were significantly lower at 3619 m ( $P =$   
257  $0.046$ ,  $d = 0.25$ ) and 4600 m ( $P = 0.038$ ,  $d = 0.29$ ), and tended to be lower at 5140 m ( $P = 0.070$ ,  $d = 0.28$ ). There  
258 were no differences observed between camps (all  $P \geq 0.512$ ,  $d \leq 0.04$ ; Figure 2a). Plasma AG:DG ratio decreased  
259 significantly from baseline to 3619 m ( $P = 0.034$ ,  $d = 0.37$ ), and tended to be lower than baseline at 4600 m ( $P =$   
260  $0.069$ ,  $d = 0.23$ ) and 5140 m ( $P = 0.070$ ,  $d = 0.25$ ). There were no differences observed between camps (all  $P \geq$   
261  $0.362$ ,  $d \leq 0.15$ ) (Figure 2c). Plasma total ghrelin concentrations decreased significantly from baseline to 4600 m

262 (P = 0.045, *d* = 0.36), however no other significant differences were observed between altitudes (all P ≥ 0.111, *d*  
263 ≤ 0.31) (Figure 2d).

264

265 **Body composition.** One way ANOVA revealed a significant effect of altitude for body mass (P < 0.001),  
266 sum of skinfolds (P < 0.001), calf girth (P = 0.010), waist girth (P = 0.016) and relaxed arm girth (P = 0.029), with  
267 no significant differences observed for flexed arm girth (P = 0.173) (Table 1).

268 Body mass increased from baseline to 3619 m (P = 0.002, *d* = 0.18), decreased between 3619 m and  
269 4600 m (P < 0.001, *d* = 0.22) and did not change between 4600 m and 5140 m (P = 0.415, *d* = 0.03). Sum of  
270 skinfolds was lower at 3619 m (P = 0.001, *d* = 0.30), 4600 m (P < 0.001, *d* = 0.34) and 5140 m (P = 0.001, *d* =  
271 0.24) compared with baseline. There were no significant differences observed between each of the camps during  
272 the trek (all P ≥ 0.116, *d* ≤ 0.09).

273 Calf girth did not differ significantly between baseline and 3619 m (P = 0.127, *d* = 0.30), however was  
274 significantly decreased at 4600 m (P = 0.039, *d* = 0.44) and 5140 m (P = 0.008, *d* = 0.60) compared with baseline.  
275 Calf girth was also significantly lower at 4600 m compared with 3619 m (P = 0.031, *d* = 0.14), and tended to be  
276 lower at 5140 m compared with 4600 m (P = 0.069, *d* = 0.14). Waist girth did not differ between baseline and any  
277 of the three camps (all P ≥ 0.122, *d* ≤ 0.15), however was significantly lower at 5140 m than 3619 m (P < 0.001,  
278 *d* = 0.29) and 4600 m (P = 0.04, *d* = 0.13). Relaxed arm girth was significantly lower at 3619 m (P = 0.022, *d* =  
279 0.30) and 4600 m (P = 0.047, *d* = 0.23) and tended to be lower at 5140 m (P = 0.073, *d* = 0.20) compared with  
280 baseline. There was a significant increase in relaxed arm girth between 3619 m and 5140 m (P = 0.047, *d* = 0.10),  
281 with no other differences observed between camps (all P ≥ 0.191, *d* ≤ 0.07).

282

283 **Correlations.** There were no correlations observed at any altitude between energy intake and CAS (all *r*  
284 ≤ 0.311, P ≥ 0.352). At 3619 m CAS tended to be associated with plasma acylated ghrelin (*r* = 0.603, P = 0.065)  
285 and total ghrelin (*r* = 0.626, P = 0.053) concentrations. Additionally, at 3619 m energy intake was significantly  
286 correlated with des-acylated ghrelin concentrations (*r* = 0.686, P = 0.029). At 4600 m CAS was significantly  
287 correlated with acylated ghrelin concentrations (*r* = 0.633, P = 0.049) and the AG:DG ratio (*r* = 0.667, P = 0.035).  
288 There were no other significant correlations observed between any variable, at any altitude (all *r* ≤ 0.511, P ≥  
289 0.108).

290

291 **DISCUSSION**

292 This study presents an assessment of the changes in appetite perceptions, energy intake, body composition, and  
293 ghrelin constituents throughout a trek to high terrestrial altitude. The findings demonstrate a reduction in energy  
294 intake and skinfold thickness during the trek, with a progressive reduction in appetite at increasing altitudes. This  
295 study provides the first investigation of acylated ghrelin and des-acylated ghrelin concentrations at terrestrial  
296 altitude and demonstrates a suppression of acylated- but not des-acylated ghrelin during the trek. These findings  
297 highlight the importance of measuring ghrelin constituents in addition to total ghrelin concentrations as small  
298 fluctuations in des-acylated ghrelin may mask changes in acylated ghrelin if only total ghrelin were to be  
299 measured. This phenomenon would have occurred at 3619 m in the present study as observed by a significant  
300 decrease in acylated ghrelin levels in the absence of any significant changes in des-acylated and total ghrelin  
301 concentrations. The findings from this study also demonstrate the need for interventions to maintain appetite  
302 during exposure to terrestrial altitudes, particularly above 4600 m.

303 In the present study energy intake was reduced by 27 % (9326 kJ) at 3619 m, 22 % (9886 kJ) at 4600 m,  
304 and 27 % (9238 kJ) at 5140 m compared with baseline, which substantiates previous findings at similar altitudes.  
305 Armellini et al. (1997) observed a 29 % decrease in energy intake in individuals climbing at approximately 4500  
306 m for 16 days, whilst Aeberli et al. (2013) demonstrated a 32 % reduction in energy intake two days after rapid  
307 ascent to 4559 m. One study however, found that energy intake, as well as fat and muscle mass, could be  
308 maintained up to an altitude of 5050 m when a wide choice of palatable foods were available in a comfortable  
309 setting (Kayser et al. 1993). It may therefore be argued that the reduction in energy intake in the present study  
310 was caused by a lack of food availability and reduced palatability of food whilst trekking in a foreign country.  
311 However, at 3619 m, energy intake was significantly suppressed compared with baseline while food was widely  
312 available and the mean palatability of the food consumed was significantly higher than baseline. These findings  
313 agree with those of Rose et al. (1988) who found a significant reduction in *ad-libitum* energy intake during a  
314 simulated ascent of Mount Everest, despite a variety of palatable foods being available. Despite the reduction in  
315 energy intake, fasting appetite perceptions were similar between baseline and 3619 m, which suggests a greater  
316 satiating effect of the energy consumed. This response was maintained at 4600 m but appetite perceptions  
317 decreased significantly at 5140 m despite consistent food intake. Observations during the trek suggested that  
318 participants were consciously trying to maintain energy intakes throughout the trek in an attempt to maintain

319 physical performance. This would support the observation that food intake was similar between the three camps  
320 but that appetite perceptions and the desire for foods decreased with increasing altitude. This mismatch between  
321 appetite perceptions and energy intake is further supported by the lack of correlation between the two variables at  
322 each altitude. The reduced palatability of the foods consumed at 5140 m also supports a reduction in appetite, and  
323 occurred despite the same ad libitum food provision throughout the trek. This effect accords with the findings  
324 from previous animal studies which suggest that hypoxia degrades the taste of food (Ettinger and Staddon 1982).  
325 Considering the significant suppression of appetite at 5140 m, it is unclear whether food intakes could continue  
326 to be maintained over a more prolonged period and targeted interventions to better maintain appetite above 4600  
327 m may be beneficial.

328 From a mechanistic perspective, trekking to high altitude induced a suppression of acylated but not des-  
329 acylated ghrelin concentrations, which resulted in a suppression of total ghrelin levels and the AG:DG ratio.  
330 However at 5140 m, the only altitude in which CAS was significantly suppressed, no correlations were observed  
331 between any blood marker and CAS or energy intake. These findings suggest that appetite regulation during high-  
332 altitude trekking may be influenced by other hormonal (e.g. leptin, glucagon-like peptide-1 and peptide YY  
333 (Debevec 2017)) and non-hormonal (e.g. taste degradation (Ettinger and Staddon 1982) potentially altering food  
334 reward (Berthoud 2006)) factors. Appetite regulation is a complex multifaceted system which involves the  
335 integration of a wide range of neuroendocrine and psychological factors (Murphy and Bloom 2006). Subsequently,  
336 appetite suppression at altitude is unlikely to be solely explained by the measurement of a single hormone.  
337 However, a better understanding of the neuroendocrine responses to high altitude trekking could be beneficial in  
338 the design of interventions to minimise appetite suppression at altitude.

339 The reductions in fasted acylated and total ghrelin concentrations, the day after significantly reduced  
340 energy intakes compared with baseline measurements, are particularly interesting considering the evidence that  
341 ghrelin levels and appetite perceptions increase in response to reduced food intake at sea level (Alajmi et al. 2016).  
342 Furthermore, acylated ghrelin levels remained depressed during the trek despite reductions in body mass between  
343 3619 m and 4600 m and the established inverse relationship between body mass and ghrelin concentrations at sea  
344 level (Chen et al. 2009; Shiiya et al. 2002). These observations suggest that the reductions in acylated and total  
345 ghrelin during this study were genuine effects of high altitude exposure rather than being secondary to any changes  
346 in food intake or body composition. Although these changes in ghrelin were small, they appear to be  
347 physiologically relevant as changes of this magnitude have previously been associated with reductions in appetite  
348 and energy intake in a laboratory environment (Bailey et al. 2015; Wasse et al. 2012). It would be beneficial for



349 future research to attempt to increase circulating plasma acylated ghrelin concentrations at altitude, in order to  
350 quantify these effects on appetite responses. Potential methods of accomplishing this include ghrelin infusion  
351 (Druce et al. 2005) or dietary interventions to manipulate ghrelin constituents (e.g. increased medium chain  
352 triglyceride intake as a substrate for ghrelin acylation (Kawai et al. 2017; Nishi et al. 2005)).

353 The reasons for the observed suppression of acylated ghrelin at altitude are unclear. However,  
354 considering the lack of change in total ghrelin levels at 3619 m, it seems plausible that the post-translational  
355 acylation of ghrelin may have been inhibited during the early stages of the trek due to inhibited ghrelin-O-  
356 acyltransferase (GOAT) activity or reduced availability of medium chain fatty acids as the substrate for acylation  
357 (Nishi et al. 2005). Alternatively, the reduction in both acylated ghrelin and total ghrelin at 4600 m suggests  
358 inhibited secretion of ghrelin from the P/D1 cells of the stomach (Kojima et al. 1999). A reduction in gut blood  
359 flow at altitude (Loshbaugh et al. 2006) has also been proposed to reduce ghrelin concentrations (Wasse et al.  
360 2012), however this concept has been disputed (Kalson et al. 2010; Mekjavic et al. 2016). The depression of  
361 acylated ghrelin levels prior to the depression of total ghrelin levels at the subsequent research camp suggests that  
362 acylated ghrelin may be a more sensitive measure of altered appetite signalling at altitude. Furthermore, the  
363 AG:DG ratio of  $>1$  in the present study supports recent data that acylated ghrelin constitutes a much larger  
364 proportion of total ghrelin than previously thought (Delhanty et al. 2015) and demonstrates that preservation of  
365 this peptide can be achieved during field research in extreme environments.

366 In accordance with previous research, considerable changes in body composition were observed during  
367 the trek. This includes a mean reduction in body mass of 2.3 kg in the three days between 3619 m and 4600 m,  
368 which was associated with significant decreases in calf girth despite very high levels of physical activity (mean  
369 distance walked:  $8.8 \text{ km}\cdot\text{day}^{-1}$ , mean elevation gain:  $491 \text{ m}\cdot\text{day}^{-1}$ ). It seems likely that the reductions in body mass  
370 were not only caused by decreases in energy intake, but also by increases in energy expenditure due to the high  
371 altitude environment and high physical activity levels. Such decreases in body and muscle mass have been  
372 observed in previous altitude research (Benso et al. 2007; Rose et al. 1988; Shukla et al. 2005; Westerterp et al.  
373 2000) and it is likely that these losses would impair physical performance in these environments (Sergi et al.  
374 2010). The further reductions in calf girth between 3619 m and 5140 m also aligns with previous research (Rose  
375 et al. 1988) but the reasons for this response are unknown. We speculate that an increase in protein degradation  
376 may have occurred due to increasing altitude exposure (Holm et al. 2010). However, contrary to these changes in  
377 calf girth, an increase in relaxed arm girth was observed between 3619 m and 5140 m. Although this increase was  
378 statistically significant, the absolute increase of 0.3 mm seems trivial. This may represent a compensatory response

379 to the significant reduction in arm girth between baseline and 3619 m which is most likely due to atrophy caused  
380 by reduced activity of the arms during final preparations and low altitude trekking. Despite these insights into  
381 changes in body composition, it must be acknowledged that the baseline measures were collected 12 days before  
382 departure from the UK, which may have confounded comparisons between baseline values and those obtained  
383 during the trek due to changes in body composition during the final preparations for the expedition. Potential  
384 increases in lean body mass from final preparations and reductions in body fat would support the observed increase  
385 in body mass at the first fixed camp and reduced skinfold values at all camps relative to baseline. At sea level, an  
386 increase in body mass would usually result in a reduction in acylated ghrelin concentrations (Chen et al. 2009),  
387 which accords with our findings at the first fixed camp. However, in the present study body mass then significantly  
388 reduced at the second and third camps, without a subsequent increase in ghrelin. This suggests that changes in  
389 ghrelin were unlikely to be the result of fluctuations in body mass during the trek and were more strongly mediated  
390 by high altitude exposure.

391 Although the findings of the present study provide novel information regarding the appetite and  
392 metabolic responses during an incremental trekking ascent to high altitude, some notable limitations must be  
393 acknowledged. First, the current study design did not include a control group to separate the effects of trekking  
394 and high altitude exposure. Therefore it is not possible to conclude that the obtained results are a consequence of  
395 hypobaric hypoxia *per se*, but are a result of high altitude trekking which combines various environmental and  
396 psychological factors as well as demanding physical exercise. An example of this being that cold exposure may  
397 interfere with energy balance, potentially by increasing non-shivering thermogenesis (van der Lans et al. 2013).  
398 Although this limits interpretation of the influence of each of these factors individually, the study design allowed  
399 us to investigate the effects of a real world gradual ascent to high altitude in order to understand the practical  
400 implications for high altitude trekking. Second, it was not possible to standardise the trekking distance on the day  
401 preceding each fixed camp due to the extreme terrain and environment. The greater trekking distance performed  
402 on the day prior to the final research camp in combination with higher altitude exposure resulted in markedly  
403 higher RPE scores at 5140 m. However, although exercise-induced anorexia occurs acutely in response to  
404 strenuous exercise, tightly controlled laboratory studies suggest that this does not affect appetite perceptions and  
405 ghrelin concentrations during the next day (King et al. 2015). Therefore the suppression of appetite at 5140 m is  
406 unlikely to be due to greater exertion during trekking on the previous day. Furthermore, increased energy  
407 expenditure from the greater trekking distance and the continued suppression of energy intake suggests that  
408 participants would have been in their greatest energy deficit at the final research camp. This would be expected

409 to increase appetite under sea level conditions (Alajmi et al. 2016) and provides further support for a genuine  
410 altitude-mediated suppression of appetite. Third, the current study did not assess hydration status during the trek,  
411 therefore it is possible that the observed reduction in body mass could be partly attributed to dehydration.  
412 However, it should be noted that reductions in skinfold and girth measurements were observed during the trek  
413 which suggests that the changes in body mass were at least partly due to genuine reductions in fat mass and muscle  
414 mass. Although the estimation of body composition using skinfold and girth measurements contains limitations,  
415 this was deemed to be the most practical and achievable method of assessment considering the extreme research  
416 environment encountered in the present study. Additionally, mean fluid intake on the day before each fixed camp  
417 was  $>3.6 \text{ L}\cdot\text{day}^{-1}$  which makes it unlikely that the participants experienced severe dehydration. It is not expected  
418 that the higher fluid intake during the trek, compared with baseline, would have influenced the measured ghrelin  
419 constituents given that gastric distension from water ingestion does not appear to influence plasma ghrelin  
420 concentrations (Tschop et al. 2000).

421 In conclusion, this study represents the first investigation of circulating ghrelin constituents in response  
422 to terrestrial altitude exposure and provides a time course of the changes in appetite, energy intake and body  
423 composition during gradual trekking ascent to high altitude. These findings demonstrate consistently reduced  
424 energy intake during high altitude exposure and an incremental reduction in appetite perceptions with increasing  
425 altitude. These changes were associated with reductions in circulating concentrations of acylated and total ghrelin  
426 during the trek but differences in the time course of these responses suggests that additional factors are also  
427 involved.. A negative energy balance during the trek caused reductions in body mass and lower body muscle mass  
428 which may have negative consequences for physical performance. Future investigations are required to develop  
429 nutritional and/or physiological interventions to maintain appetite, energy intake and muscle mass at altitude.

430

431

432 **CONFLICT OF INTEREST**

433 No conflicts of interest, financial or otherwise, are declared by the author(s). The content of this manuscript is  
434 solely the responsibility of the authors and does not necessarily represent the official views of the Defence Medical  
435 Services.

436

437 **ETHICAL APPROVAL**

438 All procedures performed in this study were in accordance with the ethical standards of the institutional and  
439 Ministry of Defence research ethics committees and with the 1964 Helsinki declaration and its later amendments  
440 or comparable ethical standards.

441

442 **ACKNOWLEDGEMENTS**

443 This work was supported by the Surgeon General (Defence Medical Services) Research Fund and Leeds Beckett  
444 University. The authors would like to acknowledge the dedication of the participants to comply with the research  
445 protocols in such a severe environment. The authors would also like to acknowledge: Joe Sails, Becky Price,  
446 Georgia Head, Timothy Keitaro, Rebekah Brady, Milton Holt and Rachael Bradley for their help with the study.

447

448

449 **REFERENCES**

- 450 Aeberli I et al. (2013) Disturbed eating at high altitude: influence of food preferences, acute mountain sickness  
 451 and satiety hormones. *European journal of nutrition* 52:625-635. doi:10.1007/s00394-012-0366-9  
 452
- 453 Al Massadi O, Lear PV, Muller TD, Lopez M, Dieguez C, Tschop MH, Nogueiras R (2014) Review of novel  
 454 aspects of the regulation of ghrelin secretion. *Current drug metabolism* 15:398-413.  
 455
- 456 Alajmi N et al. (2016) Appetite and Energy Intake Responses to Acute Energy Deficits in Females versus  
 457 Males. *Medicine and science in sports and exercise* 48:412-420. doi:10.1249/mss.0000000000000793  
 458
- 459 Armellini F et al. (1997) The effects of high altitude trekking on body composition and resting metabolic rate.  
 460 *Hormone and Metabolic Research* 29:458-461. doi:10.1055/s-2007-979077  
 461
- 462 Bailey DP et al. (2015) Appetite and gut hormone responses to moderate-intensity continuous exercise versus  
 463 high-intensity interval exercise, in normoxic and hypoxic conditions. *Appetite* 89:237-245.  
 464 doi:10.1016/j.appet.2015.02.019  
 465
- 466 Benso A, Broglio F, Aimaretti G, Lucatello B, Lanfranco F, Ghigo E, Grotoli S (2007) Endocrine and  
 467 metabolic responses to extreme altitude and physical exercise in climbers. *European journal of*  
 468 *endocrinology* 157:733-740. doi:10.1530/eje-07-0355  
 469
- 470 Berthoud HR (2006) Homeostatic and non-homeostatic pathways involved in the control of food intake and  
 471 energy balance. *Obesity* 14:197-200. doi:10.1038/oby.2006.308  
 472
- 473 Borg GA (1982) Psychophysical bases of perceived exertion. *Medicine and science in sports and exercise*  
 474 14:377-381.  
 475
- 476 Chen C-Y, Asakawa A, Fujimiya M, Lee S-D, Inui A (2009) Ghrelin Gene Products and the Regulation of Food  
 477 Intake and Gut Motility. *Pharmacological Reviews* 61:430-481. doi:10.1124/pr.109.001958  
 478
- 479 Debevec T (2017) Hypoxia-Related Hormonal Appetite Modulation in Humans during Rest and Exercise: Mini  
 480 Review. *Frontiers in Physiology* 8 doi:10.3389/fphys.2017.00366  
 481
- 482 Debevec T, Simpson EJ, Macdonald IA, Eiken O, Mekjavic IB (2014) Exercise training during normobaric  
 483 hypoxic confinement does not alter hormonal appetite regulation. *PloS one* 9:e98874.  
 484 doi:10.1371/journal.pone.0098874  
 485
- 486 Debevec T, Simpson EJ, Mekjavic IB, Eiken O, Macdonald IA (2016) Effects of prolonged hypoxia and bed  
 487 rest on appetite and appetite-related hormones. *Appetite* 107:28-37. doi:10.1016/j.appet.2016.07.005  
 488
- 489 Delhanty PJ et al. (2015) The acylated (AG) to unacylated (UAG) ghrelin ratio in esterase inhibitor-treated  
 490 blood is higher than previously described. *Clinical endocrinology* 82:142-146. doi:10.1111/cen.12489  
 491
- 492 Druce MR et al. (2005) Ghrelin increases food intake in obese as well as lean subjects. *International journal of*  
 493 *obesity* (2005) 29:1130-1136. doi:10.1038/sj.jco.0803001  
 494
- 495 Ettinger RH, Staddon JE (1982) Decreased feeding associated with acute hypoxia in rats. *Physiology &*  
 496 *behavior* 29:455-458.  
 497
- 498 Fawcett JK, Wynn V (1960) Effects of posture on plasma volume and some blood constituents. *Journal of*  
 499 *clinical pathology* 13:304-310.  
 500
- 501 Fernandez G, Cabral A, Cornejo MP, De Francesco PN, Garcia-Romero G, Reynaldo M, Perello M (2016) Des-  
 502 Acyl Ghrelin Directly Targets the Arcuate Nucleus in a Ghrelin-Receptor Independent Manner and  
 503 Impairs the Orexigenic Effect of Ghrelin. *Journal of neuroendocrinology* 28:12349.  
 504 doi:10.1111/jne.12349  
 505

506 Flint A, Raben A, Blundell JE, Astrup A (2000) Reproducibility, power and validity of visual analogue scales in  
507 assessment of appetite sensations in single test meal studies. *International journal of obesity and related*  
508 *metabolic disorders* 24:38-48.  
509

510 Guenther PM, Kott PS, Carriquiry AL (1997) Development of an approach for estimating usual nutrient intake  
511 distributions at the population level. *The Journal of nutrition* 127:1106-1112.  
512

513 Hill RJ, Davies PS (2001) The validity of self-reported energy intake as determined using the doubly labelled  
514 water technique. *The British journal of nutrition* 85:415-430.  
515

516 Holm L, Haslund ML, Robach P, van Hall G, Calbet JAL, Saltin B, Lundby C (2010) Skeletal Muscle  
517 Myofibrillar and Sarcoplasmic Protein Synthesis Rates Are Affected Differently by Altitude-Induced  
518 Hypoxia in Native Lowlanders. *PloS one* 5:e15606. doi:10.1371/journal.pone.0015606  
519

520 Hopkins WG (2004) How to interpret changes in an athletic performance test. *Sportscience* 8:1-7.  
521

522 Hosoda H et al. (2004) Optimum collection and storage conditions for ghrelin measurements: octanoyl  
523 modification of ghrelin is rapidly hydrolyzed to desacyl ghrelin in blood samples. *Clinical chemistry*  
524 50:1077-1080. doi:10.1373/clinchem.2003.025841  
525

526 Kalson NS, Hext F, Davies AJ, Chan CW, Wright AD, Imray CH (2010) Do changes in gastro-intestinal blood  
527 flow explain high-altitude anorexia? *European journal of clinical investigation* 40:735-741.  
528 doi:10.1111/j.1365-2362.2010.02324.x  
529

530 Kargotich S, Goodman C, Keast D, Morton AR (1998) The influence of exercise-induced plasma volume  
531 changes on the interpretation of biochemical parameters used for monitoring exercise, training and  
532 sport. *Sports medicine* 26:101-117.  
533

534 Kawai K et al. (2017) Ghrelin activation and neuropeptide Y elevation in response to medium chain triglyceride  
535 administration in anorexia nervosa patients. *Clinical nutrition ESPEN* 17:100-104.  
536 doi:10.1016/j.clnesp.2016.10.001  
537

538 Kayser B, Narici M, Milesi S, Grassi B, Cerretelli P (1993) Body Composition and Maximum Alactic  
539 Anaerobic Performance During a One Month Stay at High Altitude. *International journal of sports*  
540 *medicine* 14:244-247. doi:10.1055/s-2007-1021171  
541

542 King JA, Garnham JO, Jackson AP, Kelly BM, Xenophontos S, Nimmo MA (2015) Appetite-regulatory  
543 hormone responses on the day following a prolonged bout of moderate-intensity exercise. *Physiology*  
544 *& behavior* 141:23-31. doi:10.1016/j.physbeh.2014.12.050  
545

546 Kojima M, Hosoda H, Date Y, Nakazato M, Matsuo H, Kangawa K (1999) Ghrelin is a growth-hormone-  
547 releasing acylated peptide from stomach. *Nature* 402:656-660.  
548

549 Loshbaugh JE, Loeppky JA, Greene ER (2006) Effects of acute hypobaric hypoxia on resting and postprandial  
550 superior mesenteric artery blood flow. *High altitude medicine & biology* 7:47-53.  
551 doi:10.1089/ham.2006.7.47  
552

553 Matu J, Deighton K, Ispoglou T, Duckworth L (2017) The effect of moderate versus severe simulated altitude  
554 on appetite, gut hormones, energy intake and substrate oxidation in men. *Appetite* 113:284-292.  
555

556 Mekjavic IB et al. (2016) The effect of normobaric hypoxic confinement on metabolism, gut hormones and  
557 body composition. *Frontiers in Physiology* 7 doi:10.3389/fphys.2016.00202  
558

559 Mellor A et al. (2017) British Services Dhaulagiri Medical Research Expedition: A unique military/civilian  
560 research collaboration.  
561

562 Mellor AJ, Woods DR, O'Hara J, Howley M, Watchorn J, Boos C (2014) Rating of perceived exertion and acute  
563 mountain sickness during a high-altitude trek. *Aviation, space, and environmental medicine* 85:1214-  
564 1216. doi:10.3357/asem.4083.2014  
565

566 Morishima T, Goto K (2016) Ghrelin, GLP-1, and leptin responses during exposure to moderate hypoxia.  
567 Applied physiology, nutrition, and metabolism 41:375-381. doi:10.1139/apnm-2015-0311  
568

569 Murphy KG, Bloom SR (2006) Gut hormones and the regulation of energy homeostasis. Nature 444:854-859.  
570 doi:10.1038/nature05484  
571

572 Nishi Y et al. (2005) Ingested medium-chain fatty acids are directly utilized for the acyl modification of ghrelin.  
573 Endocrinology 146:2255-2264. doi:10.1210/en.2004-0695  
574

575 Riepl RL et al. (2012) Influence of acute exposure to high altitude on basal and postprandial plasma levels of  
576 gastroenteropancreatic peptides. PloS one 7:e44445. doi:10.1371/journal.pone.0044445  
577

578 Roach RC, Bärtzsch P, Oelz O, Hackett PH (1993) The Lake Louise acute mountain sickness scoring system. In:  
579 Sutton JR, Houston CS, Goates G (eds) Hypoxia and Molecular Medicine. Queens City Burlington,  
580 VT, pp 272-274  
581

582 Rose MS, Houston CS, Fulco CS, Coates G, Sutton JR, Cymerman A (1988) Operation Everest. II: Nutrition  
583 and body composition. Journal of applied physiology 65:2545-2551.  
584

585 Sergi G et al. (2010) Changes in total body and limb composition and muscle strength after a 6-8 weeks sojourn  
586 at extreme altitude (5000-8000 m). The Journal of sports medicine and physical fitness 50:450-455.  
587

588 Shiiya T et al. (2002) Plasma ghrelin levels in lean and obese humans and the effect of glucose on ghrelin  
589 secretion. The Journal of clinical endocrinology and metabolism 87 doi:10.1210/jcem.87.1.8129  
590

591 Shukla V, Singh SN, Vats P, Singh VK, Singh SB, Banerjee PK (2005) Ghrelin and leptin levels of sojourners  
592 and acclimatized lowlanders at high altitude. Nutritional neuroscience 8:161-165.  
593 doi:10.1080/10284150500132823  
594

595 Sierra-Johnson J, Romero-Corral A, Somers VK, Johnson BD (2008) Last word on viewpoint: effect of altitude  
596 on leptin levels, does it go up or down? Journal of applied physiology 105:1691.  
597 doi:10.1152/jappphysiol.90679.2008  
598

599 Snyder EM, Carr RD, Deacon CF, Johnson BD (2008) Overnight hypoxic exposure and glucagon-like peptide-1  
600 and leptin levels in humans. Applied physiology, nutrition, and metabolism 33:929-935.  
601 doi:10.1139/h08-079  
602

603 Stubbs RJ et al. (2000) The use of visual analogue scales to assess motivation to eat in human subjects: a review  
604 of their reliability and validity with an evaluation of new hand-held computerized systems for temporal  
605 tracking of appetite ratings. The British journal of nutrition 84:405-415.  
606

607 Tschop M, Smiley DL, Heiman ML (2000) Ghrelin induces adiposity in rodents. Nature 407:908-913.  
608 doi:10.1038/35038090  
609

610 van der Lans AA et al. (2013) Cold acclimation recruits human brown fat and increases nonshivering  
611 thermogenesis. The Journal of clinical investigation 123:3395-3403. doi:10.1172/jci68993  
612

613 Vissing K, McGee S, Farup J, Kjolhede T, Vendelbo M, Jessen N (2013) Differentiated mTOR but not AMPK  
614 signaling after strength vs endurance exercise in training-acclimated individuals. Scandinavian journal  
615 of medicine & science in sports 23:355-366. doi:10.1111/j.1600-0838.2011.01395.x  
616

617 Wasse LK, Sunderland C, King JA, Batterham RL, Stensel DJ (2012) Influence of rest and exercise at a  
618 simulated altitude of 4,000 m on appetite, energy intake, and plasma concentrations of acylated ghrelin  
619 and peptide YY. Journal of applied physiology 112:552-559. doi:10.1152/jappphysiol.00090.2011  
620

621 Westerterp KR, Meijer EP, Rubbens M, Robach P, Richalet JP (2000) Operation Everest III: energy and water  
622 balance. European journal of physiology 439:483-488. doi:10.1007/s004249900203  
623

ACCEPTED VERSION



625 **Figure 1** Energy intake (a), palatability score (b) and composite appetite score (c) at baseline, 3619 m, 4600 m  
626 and 5140 m. \*significant difference from baseline. ‡significant difference between 4600 m and 5140 m.  
627 §significant difference between 3619 m and 5140 m (One way ANOVA;  $P < 0.05$  after post-hoc analyses). Values  
628 are mean (SE),  $N = 12$  for energy intake and palatability,  $N = 11$  for composite appetite score

629

630

631 **Figure 2** Acylated ghrelin (a), des-acylated ghrelin (b), AG:DG ratio (c) and total ghrelin (d) concentrations at  
632 baseline, 3619 m, 4600 m and 5140 m. \*significant difference from baseline (One way ANOVA;  $P < 0.05$  after  
633 post-hoc analyses). Values are mean (SE),  $N = 11$

634

ACCEPTED VERSION

635 **Table 1** Body composition measurements at baseline, 3619 m, 4600 m and 5140 m.

	Baseline	3619 m	4600 m	5140 m
Body mass (kg)	71.3 (10.3)	73.1 (10.2)*	70.8 (10.7) <sup>†</sup>	71.1 (10.0) <sup>§</sup>
Sum of 8 Skinfolts (mm)	81.2 (23.7)	74.2 (22.3)*	73.5 (21.7)*	75.5 (23.9)*
Calf girth (cm)	38.1 (1.9)	37.5 (2.1)	37.2 (2.2)* <sup>†</sup>	36.9 (2.1)* <sup>§</sup>
Waist girth (cm)	77.5 (6.6)	78.2 (5.5)	77.3 (5.5)	76.6 (5.4) <sup>‡§</sup>
Relaxed arm girth (cm)	29.5 (3.0)	28.6 (3.0)*	28.8 (3.1)*	28.9 (2.9) <sup>§</sup>
Flexed arm girth (cm)	30.6 (3.2)	30.2 (3.2)	30.1 (3.2)	30.1 (3.1)

636 Values are mean (SD), N = 12. \*significant difference from baseline. <sup>†</sup>significant difference between 3619 m and  
 637 4600 m. <sup>‡</sup>significant difference between 4600 m and 5140 m. <sup>§</sup>significant difference between 3619 m and 5140 m  
 638 (One way ANOVA; P < 0.05 after post-hoc analyses)

639