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1 APPETITE AND ENERGY INTAKE RESPONSES TO ACUTE ENERGY DEFICITS

2 IN FEMALES VERSUS MALES

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27 ABSTRACT

Purpose: To explore whether compensatory responses to acute energy deficits induced by 28 exercise or diet differ by sex. Methods: In experiment one, twelve healthy women completed 29 30 three 9 h trials (control, exercise-induced (Ex-Def) and food restriction induced energy deficit (Food-Def)) with identical energy deficits being imposed in the Ex-Def (90 min run, ~70% of 31 VO₂ max) and Food-Def trials. In experiment two, 10 men and 10 women completed two 7 h 32 trials (control and exercise). Sixty min of running (~70% of VO₂ max) was performed at the 33 beginning of the exercise trial. Participants rested throughout the remainder of the exercise 34 35 trial and during the control trial. Appetite ratings, plasma concentrations of gut hormones and ad libitum energy intake were assessed during main trials. Results: In experiment one, an 36 energy deficit of ~3500 kJ induced via food restriction increased appetite and food intake. 37 38 These changes corresponded with heightened concentrations of plasma acylated ghrelin and 39 lower peptide YY₃₋₃₆. None of these compensatory responses were apparent when an equivalent energy deficit was induced by exercise. In experiment two, appetite ratings and 40 41 plasma acylated ghrelin concentrations were lower in exercise than control but energy intake did not differ between trials. The appetite, acylated ghrelin and energy intake response to 42 exercise did not differ between men and women. Conclusions: Women exhibit compensatory 43 appetite, gut hormone and food intake responses to acute energy restriction but not in 44 45 response to an acute bout of exercise. Additionally, men and women appear to exhibit similar 46 acylated ghrelin and PYY₃₋₃₆ responses to exercise-induced energy deficits. These findings advance understanding regarding the interaction between exercise and energy homeostasis in 47 48 women.

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50 KEY WORDS: sex-based differences; gastrointestinal hormones; compensation; energy
51 balance; females

52 INTRODUCTION

The regulation of appetite control and energy balance is an area of scientific enquiry which 53 continues to receive widespread attention across disciplines. To date, as in many fields of 54 science, the foundation of our knowledge within appetite regulation has been gleaned from 55 studies conducted predominantly in men. Consequently, less in known specifically regarding 56 the regulation of appetite control and energy balance in women and the potential for sex-57 based differences has not been thoroughly investigated. Preliminary research has hinted that 58 appetite and appetite-regulatory hormones may display divergent responses to nutritional 59 60 interventions between men and women however this proposition continues to be debated (5). Specifically, compared to men, it has been suggested that women exhibit more potent 61 compensatory responses (appetite, appetite regulatory hormones, food intake) to energy 62 63 deficits in order to preserve energy balance and reproductive function (14). This viewpoint is 64 supported by studies demonstrating that men exhibit greater reductions in body fat and body mass than women in response to supervised exercise training (8,19,35). Conversely, other 65 66 research has suggested that differences in weight loss and adiposity responses to exercise are unrelated to sex (5,6). 67

68

Sex-based differences in the short-term regulation of appetite and energy balance were 69 previously investigated in a carefully designed experimental study using consecutive days of 70 71 exercise to induce an energy deficit in male and female participants (15). The researchers showed that this acute exercise-induced energy deficit triggered a compensatory increase in 72 circulating acylated ghrelin (appetite stimulating hormone) in women but not in men. These 73 74 changes corresponded with higher appetite ratings in women than men and suggest that sexbased differences may be apparent in the early appetite and gut hormone response to 75 76 exercise-induced energy deficits.

77 Over the past decade our laboratory has conducted many acute experimental trials seeking to 78 enhance understanding concerning the short-term regulation of appetite and energy balance (21,31). In a sample of male participants, we recently demonstrated that the induction of an 79 80 acute energy deficit by food restriction elicited a rapid and robust compensatory appetite, gut hormone (acylated ghrelin and PYY₃₋₃₆) and energy intake response whilst the same energy 81 82 deficit imposed by exercise had no effect (20). These findings suggest that the method by 83 which an energy deficit is imposed has a marked impact on the subsequent physiological and behavioural response. It is currently unknown whether women exhibit the same acute 84 85 responses to exercise and food restriction as men. This information has important implications regarding the utility of lifestyle therapies to assist weight control in women. 86

87

88 Within this report we describe the findings from two acute experimental studies which sought 89 to provide new information regarding the short-term appetite, food intake and appetite hormone responses to exercise and food-induced energy deficits in men and women. In 90 91 experiment one; we compared the appetite, energy intake, acylated ghrelin and PYY₃₋₃₆ responses to an equivalent energy deficit induced by exercise or energy restriction in women. 92 In experiment two, we directly compared appetite, food intake and circulating acylated 93 ghrelin responses to an exercise-induced energy deficit in men verses women. Our findings 94 95 identify a high degree of similarity in the acute response to energy deficits in men and 96 women.

97

98 METHODS

99 Experimental protocol

100 This investigation contained two experiments which were conducted according to the 101 guidelines laid down in the Declaration of Helsinki. All procedures were approved by the

Institutional Ethics Advisory Committee and written informed consent was obtained from all participants. Study participants were non-smokers, not taking medication, weight stable for at least six months before participation and were not dieting. Participants had no known history of cardiovascular/metabolic disease and with respect to female participants, were of reproductive age but were not pregnant. In each study, participants were recreationally active i.e. were familiar with exercise, but were not formally trained in endurance activities such as running or cycling.

109

110 Participants completed a weighed food diary in the 24 h before the first main trial of each experiment and replicated this before each subsequent trial. Alcohol, caffeine and strenuous 111 physical activity were not permitted during this period. All trials commenced between 8am 112 113 and 9am after an overnight fast of at least 10 h and participants exerted themselves minimally 114 when travelling to the laboratory, using motorised transport when possible. Verbal confirmation of dietary and exercise standardisation was obtained at the beginning of each 115 experimental trial. Female participants completed all main trials within the follicular phase of 116 the menstrual cycle (4). 117

118

119 **Preliminary trials**

In order to determine the running speed required to elicit 70% of maximum oxygen uptake (VO₂ max) for each individual, participants completed a preliminary trial before the main trials for each experiment. This consisted of a submaximal running test and a VO₂ max test on a motorised treadmill (34). Anthropometric measurements and study questionnaires e.g. Three Factor Eating Questionnaire (TFEQ) (33) was also taken/completed at this time. At this visit, participants also verbally confirmed acceptability of the test meals and ad *libitum* meals subsequently to be provided during main experimental trials. In experiment one a second preliminary trial was completed to determine the net energy costof exercise which was needed to calculate food provision in the main trials and to enable trial

randomisation in advance. During this session participants ran for 90 min at 70% of VO_2 max with expired air samples being collected into Douglas bags at 15 min intervals to calculate energy expenditure using the equations provided by Frayn (12).

133

134 Experiment One

135 Twelve female participants performed three 9 h experimental trials (control (Con)), exerciseinduced energy deficit (Ex-Def) and diet-induced energy deficit (Food-Def)) separated by 136 one-week in a randomised counterbalanced design. To ensure standardisation of menstrual 137 138 phase, participants' first main trial was undertaken at the beginning of their follicular phase with their second trial occurring one-week later. Participants' third main trial was 139 subsequently undertaken at the beginning of their next cycle approximately four weeks later. 140 Participants rested within the laboratory throughout all trials with participants being 141 permitted to read, work at a computer or watch DVDs which had been screened to ensure that 142 there was no overt emphasis on food and drink. The exception to this occurred at 0-1.5 h 143 during Ex-Def where participants performed 90 min of treadmill running at ~70% of VO₂ 144 max (identical to that performed during the preliminary trial). Resting expired air samples 145 146 were collected from 0 - 1.5 h during the Con and Food-Def trials to calculate the net energy expenditure of exercise (gross energy expenditure of exercise minus energy expenditure at 147 rest) (12). 148

149

150 Identical test meals were provided at 2 h (breakfast) and 4.75 h (lunch) and were each 151 consumed within 15 min. The meals consisted of a tuna and mayonnaise sandwich, salted

152 crisps, chocolate muffin and green apple. The macronutrient composition of the meal was 47% carbohydrate, 18% protein and 35% fat. The energy content of the test meals was 153 identical in Con and Ex-Def (2778 (109) kJ) with each meal providing 35% of participants' 154 estimated daily energy needs for a sedentary day. This calculation was based upon an 155 estimation of each participant's daily energy needs which was determined using a validated 156 equation for resting metabolic rate (28) that was multiplied by an activity factor (1.4) deemed 157 appropriate for a sedentary day (10). In Food-Def, the energy content of the test meals was 158 reduced (1025 (159) kJ) by deducting the net energy expenditure of exercise from the energy 159 160 provided at the test meals during Con and Ex-Def. This energy deficit was individually prescribed based on the exercise energy expenditure data derived from the preliminary trials 161 and the total amount of energy deducted was divided equally between breakfast and lunch. 162 163 Therefore, equivalent energy deficits were induced in Ex-Def and Food-Def relative to Con. The macronutrient percentage of the test meals was identical across main trials i.e. only the 164 meal energy content was altered in the Food-Def trial. 165

166

167 Experiment Two

Ten female and 10 male participants performed two 7 h experimental trials (exercise and 168 control) separated by one week in a randomised counterbalanced design. Female participants 169 completed both main trials during the follicular phase (days 1 - 11) of their menstrual cycle. 170 171 Participants rested within the laboratory throughout each trial, except from 0 - 1 h during the exercise trial where participants performed 60 min of treadmill running at \sim 70% of VO₂ max. 172 Expired air samples were collected as described earlier to calculate the net energy 173 174 expenditure of exercise. A test meal was provided at 2 h, consisting of a ham sandwich, banana, salted crisps and chocolate bar. The macronutrient composition of the meals was 175

176 63% carbohydrate, 9% protein and 28% fat. The energy content was 42 kJ per kg body mass
177 (men 3167 (395) kJ; women 2599 (305) kJ).

178

179 Appetite perceptions and *ad libitum* buffet meals

Appetite perceptions (hunger, satisfaction, fullness and prospective food consumption) were 180 assessed at baseline and every 30 min during both experiments using 100 mm visual analogue 181 scales (11). An overall appetite rating was calculated for each time-point as the mean value of 182 the four appetite perceptions after inverting the values for satisfaction and fullness (32). At 8 183 184 h during experiment one and 5 h during experiment two, participants were given 30 min access to a buffet meal from which they were free to select and consume food *ad libitum*. The 185 buffet was set up identically before each meal with food being presented in excess of 186 187 expected consumption. The items available were milk, three varieties of cereal, cereal bars, white bread, brown bread, ham, cheese, tuna, mayonnaise, butter, margarine, cookies, 188 chocolate rolls, apples, oranges and bananas. Participants were told to eat until satisfied and 189 190 that additional food was available if desired. Participants were not overtly aware that their food intake was being monitored with actual intake being deduced by experimenters covertly 191 re-weighing leftover foods after ad libitum meals. Energy and macronutrient intake was 192 determined using values provided by the food manufacturers. All meals were consumed in 193 194 isolation so that social influence did not affect food selection. Water was available ad libitum 195 throughout each trial.

196

197 Blood sampling and analysis

During the experimental trials, venous blood samples were collected via a cannula (Venflon,
Becton Dickinson, Helsinborg, Sweden) inserted into an antecubital vein. Blood samples
were collected at baseline, 2, 3, 4.75, 6, 7, 8 and 9 h in experiment one and baseline, 0.5, 1, 2,

201 2.5, 3, 4, 4.5, 5, 5.5, 6, and 7 h in experiment two. Plasma acylated ghrelin concentrations 202 were measured from blood samples in both experiments and PYY_{3-36} was additionally 203 measured in experiment one. Details on acylated ghrelin and PYY_{3-36} sample collection and 204 processing have been described in-depth previously (7).

205

A commercially available enzyme immunoassay was used to determine plasma 206 concentrations of acylated ghrelin (SPI BIO, Montigny le Bretonneux, France). Plasma 207 concentrations of PYY₃₋₃₆ were determined using a commercially 208 available 209 radioimmunoassay (Millipore, Watford, UK). To eliminate interassay variation, samples from each participant were analysed in the same run. The within batch coefficient of variation for 210 211 the assays were 6.9 and 6.8% for acylated ghrelin and PYY₃₋₃₆, respectively.

212

213 Statistical analysis

Data was analysed using IBM SPSS statistics version 19 for Windows. Time-averaged area 214 under the curve (AUC) values were calculated using the trapezoidal method. For experiment 215 one, one-way repeated measures ANOVA was used to assess trial-based differences in 216 energy intake at the *ad libitum* meal as well as AUC values for appetite, acylated ghrelin and 217 PYY₃₋₃₆. For experiment two, independent samples t-tests were used to assess baseline 218 219 differences between male and female participants. Mixed measures, two-way ANOVA (sex x 220 trial) was used to assess differences in energy intake and AUC values for appetite and acylated ghrelin. Where significant main effects were found, post-hoc analysis was 221 performed using Holm-Bonferonni correction for multiple comparisons. Statistical 222 significance for this study was accepted as $P \le 0.05$. Results in text and tables are presented 223 as mean (SD). Graphical representations of results are presented as mean (SEM) to avoid 224 225 distortion of the graphs.

226 Sample size calculations

The sample sizes employed within this study were deemed sufficient to detect a significant 227 difference in energy intake between trials in experiment one and a significant difference in 228 229 relative energy intake between sexes in experiment two. These variables were selected as the 230 primary outcome measure for each experiment. The anticipated effect size for a difference in energy intake between trials for experiment one was based on previous findings from our 231 laboratory using an identical experimental protocol in men (20). The anticipated effect size 232 for a difference in relative energy intake between sexes for experiment two was based on the 233 234 findings from previous research that employed similar methods to the present experiment (16). Based on these effect sizes and an alpha value of 5%, a sample size of 12 participants in 235 experiment one would have > 95 % power to detect a difference in energy intake and 20 236 237 participants (10 men and 10 women) in experiment two would have > 87 % power to detect a difference in relative energy intake between sexes. All calculations were performed using 238 G*power (9). 239

240

241 **RESULTS**

242 Experiment One

243 Participant characteristics and exercise responses

The physical characteristics of participants are described in Table 1. Participants rated 'low' for each trait within the TFEQ (cognitive restraint 7.8 (3.3); disinhibition 7.9 (3.2); hunger 6.9 (3.1). Participants completed the 90 min run at 8.6 (1.0) km.h⁻¹. This elicited an oxygen consumption equivalent to 70.2 (1.5) % of VO₂ max and a net energy expenditure of 3560 (382) kJ. The non-protein respiratory exchange ratio was 0.86 (0.04) which reflected a proportional contribution to energy provision of 54 (13) % carbohydrate and 46 (13) % fat. Heart rate and rating of perceived exertion (RPE) were 175 (3) beats.min⁻¹ and 13 (1),
respectively.

252

253 Appetite and energy intake

Overall appetite ratings did not differ between trials at baseline (Ex-Def 71 (23); Food-Def 77 (12); Con 75 (16); P = 0.536). One-way ANOVA revealed higher appetite AUC in Food-Def than Ex-Def and Con across the 9 h trial (P < 0.0005; Figure 1 and 2). At the *ad libitum* buffet meal, total energy intake was significantly higher in Food-Def than Ex-Def and Control (Ex-Def 2774 (1682); Food-Def 3965 (1409); Control 2560 (1112) kJ; P < 0.0005). Similarly, energy intake from fat, protein and carbohydrate was significantly higher in Food-Def than Ex-Def and Control (all P < 0.004; data not presented).

261

262 Plasma acylated ghrelin and PYY₃₋₃₆ concentrations

Due to problems with venous cannulation acylated ghrelin and PYY₃₋₃₆ data is only available 263 264 for 11 participants. Fasting plasma acylated ghrelin concentrations did not differ significantly between trials at baseline (Con 148 (100); Ex-Def 140 (86); Food-Def 148 (96) $pg.mL^{-1}$; P = 265 0.422). Acylated ghrelin concentrations were significantly higher in Food-Def and 266 significantly lowest in Ex-Def across the 9 h trial (P < 0.0005; Figure 1 and 2). Fasting PYY₃-267 ₃₆ concentrations did not differ significantly between trials at baseline (Con 77 (39); Ex-Def 268 76 (34); Food-Def 77 (36) pg.mL⁻¹; P = 0.989). Time-averaged AUC for PYY₃₋₃₆ was 269 significantly highest in Ex-Def and significantly lowest in Food-Def across the 9 h trial (P < 270 0.0005; Figure 1 and 2). 271

272

273

275 Experiment Two

276 Participant characteristics and exercise responses

The physical characteristics of the participants are described and contrasted (men versus 277 women) in Table 1. There were no differences between men and women in their TFEQ scores 278 for cognitive restraint (men: 6 (1); women: 8 (2)), disinhibition (men: 4 (1); women: 6 (1)) or 279 hunger (men: 6 (1); women: 7 (1)). The 60 min run was completed at a significantly higher 280 speed in men than women (men: 10.7 (0.7) km.h⁻¹; women: 8.4 (0.3) km.h⁻¹; P = 0.006). The 281 run also generated a greater net energy expenditure in men than women (men: 3971 (200) kJ; 282 283 women: 2536 (126) kJ; P < 0.0005). However, there was no difference in relative exercise intensity (70.9 (1.4) % and 73.3 (0.6) % of VO₂ max in men and women respectively; P =284 0.130). There was a tendency for a lower heart rate in men than women (men: 163 (4) 285 beats.min⁻¹; women: 174 (4) beats.min⁻¹; P = 0.068). Ratings of perceived exertion did not 286 differ between sexes (13 (1) and 12 (0) in men and women respectively; P = 0.797). 287

288

289 Appetite and energy intake

Appetite did not differ by trial (exercise vs. Con) or sex at baseline (Female-Ex 61 (22); Female Con 65 (11); Male Ex 70 (12); Male Con 74 (11); all P > 0.05). Two-way ANOVA revealed main effects of trial (P = 0.05) and sex (P = 0.01) for AUC appetite ratings across the 7 h trial, with higher appetite ratings in men than women and in control compared with exercise (Figure 3).

295

Two-factor ANOVA revealed a main effect of sex for energy intake (P = 0.023) and carbohydrate intake (P = 0.013) during the *ad libitum* buffet meal, indicating greater consumption by men than women. Differences between sexes no longer remained after intakes were adjusted for lean body mass (both P \ge 0.289). There was no effect of trial for energy or macronutrient intake and no differences between sexes for fat and protein intake (both P > 0.05; Table 2).

302

Two-factor ANOVA revealed a main effect of trial for relative energy intake (energy intake minus net energy expenditure of exercise) indicating lower relative energy intake in the exercise trial compared with control (Female Ex 442 (1711); Female Con 2916 (1510); Male Ex 1414 (2510); Male Con 4971 (2648) kJ; P < 0.0005). This resulted in a similar energy deficit for men and women in the exercise trial relative to control (men: 3557 (598); women: 2474 (406) kJ; P = 0.152).

309

310 Acylated ghrelin

Due to problems with venous cannulation, acylated ghrelin data is only available for 8 men and 8 women. Baseline values were not different between control and exercise trials (P > 0.05) but were significantly higher in women than men (Female Ex 155 (101); Female Con 178 (61); Male Ex 71 (31); Male Con 100 (56); P = 0.018). Two-way ANOVA revealed main effects of trial (P = 0.004) and sex (P = 0.034) for AUC acylated ghrelin concentrations across the 7 h trial, with higher concentrations in women than men and in control compared with exercise (Figure 4).

318

319 **DISCUSSION**

In recent years there has been an explosion of research examining the interaction between exercise and energy homeostasis. One area which has received widespread attention is the influence of exercise and associated changes in energy balance on gut hormones which have been identified as key regulators of appetite, energy intake and adiposity (21,30,31). To date, the majority of research within these areas has been conducted using male particpants 325 meaning that much less is known regarding the interaction between acute exercise and food intake regulation in women. The findings of the present experiments demonstrate that women 326 respond similarly to men with regards to short-term responses to energy deficits induced by 327 328 exercise and food restriction. Specifically, in accordance with our previous results in male participants (20), in experiment one, our female sample demonstrated rapid and robust 329 compensatory appetite, energy intake and appetite hormone responses (acylated ghrelin and 330 PYY₃₋₃₆) to energy deficits induced by food restriction but not exercise. Additionally, in 331 experiment two, both male and female participants exhibited suppressed appetite and 332 333 circulating acylated ghrelin in response to exercise without any change in *ad libitum* energy intake being apparent. These data provide new information regarding short-term 334 physiological and behavioural responses to energy deficits in women. 335

336

Experiment one showed that in women an acute energy deficit of ~3500 kJ robustly 337 stimulated appetite and energy intake when induced via energy restriction but such 338 compensatory responses did not occur when an equivalent deficit was induced by exercise. 339 These outcomes are consistent with the findings from an identical previous study in men (20) 340 and highlight the importance of oro-gastric mechanisms e.g. stomach distention and/or 341 passage of nutrients through the gastrointestinal tract, for short-term appetite control in men 342 343 and women (2,36). Such regulatory mechanisms are complemented by a network of appetite 344 regulatory hormones, and the identification of higher circulating concentrations of acylated ghrelin, and lower PYY_{3-36} in response to energy restriction, is consistent with the known 345 acute regulatory actions of these hormones (24,25). In contrast, within experiment one, 346 347 exercise elicited reductions in circulating acylated ghrelin and elevations in PYY₃₋₃₆ across the 9 h trial in our female sample. These responses are consistent with previous studies in 348 men which have identified a potent capacity of exercise to perturb the circulating 349

350 concentrations of these hormones in directions associated with a reduction in appetite (30). The mechanisms promoting such changes are unclear and were not investigated in the present 351 experiments. It has been suggested that exercise-induced changes in sympathetic nervous 352 353 system activity (3,38) and splanchnic blood flow (29,37) may be important, however additional work is needed to investigate this issue. As per our previous findings in males (20), 354 the results from experiment one demonstrate the usefulness of exercise for weight 355 356 management in women to minimise compensatory responses associated with energy deficits produced solely by dietary restriction. Additional research is now needed to determine the 357 358 more prolonged impact of exercise and diet-related energy deficits on appetite and energy intake in men and women; research that will provide more tangible information for 359 individuals concerned with weight management. 360

361

The second experiment of this paper demonstrated that an acute bout of exercise, performed 362 at the same relative exercise intensity, decreased appetite ratings in men and women. 363 364 Furthermore, this response was consistent with lower acylated ghrelin concentrations in both sexes and the absence of any compensatory increase in *ad libitum* energy intake. These 365 findings are consistent with the suggestion that men and women do not differ in their 366 physiological and behavioural responses to exercise (5) and this notion is supported by 367 368 previous data, albeit with a very brief period of observation after exercise (16). Our findings 369 therefore add to the literature by demonstrating that acute responses to exercise do not differ 370 between men and women over a prolonged duration within the laboratory.

371

In contrast to the present results, previous research has shown that appetite is not suppressed in women during exercise (18,22,23). Furthermore, Larson-Meyer et al. (23) observed an increase in circulating acylated ghrelin in response to acute exercise; contrasting the

suppression reported in the present paper. The discrepant findings with regards to appetite 375 may be related to exercise intensity with the intensity in the present studies being much 376 greater (70% of VO₂ max) than that employed by Hopkins et al (18) (~50% of VO₂ max). 377 378 Training status and familiarity with exercise also moderate exercise-related appetite responses (26,27) and the lack of influence of exercise on appetite in the studies of King et al. 379 (22) and Larson-Meyer et al. (23) may be because their participants were regularly active and 380 particularly familiar with the mode of exercise employed. An increase in circulating acylated 381 ghrelin in response to exercise (23) contrasts the present findings and the bulk of the 382 383 literature which has studied men (30). Regression to the mean may have been a confounding factor in the study of Larson-Meyer et al. (23) however. Furthermore, differences in the 384 analytical techniques utilised between studies may also be influential. Nonethless, despite 385 386 these noted discrepancies, ad libitum energy intake remained unchanged in each of the 387 aforementioned studies. Thus, as seen in men, single sessions of exercise do not appear to influence energy intake in women. 388

389

Although we found no differences between sexes in compensatory responses to exercise, 390 391 females participants exhibited significantly higher plasma acylated ghrelin concentrations across main trials compared with men -a finding which has been reported previously (13). 392 Despite this disparity, appetite ratings were paradoxically higher in men than women across 393 394 main trials. This difference may highlight the importance of relative changes in gut hormone concentrations, rather than absolute circulating levels which may markedly differ between 395 individuals. The similar acylated ghrelin response to exercise in both sexes may therefore 396 397 underpin the comparable appetite and energy intake responses observed. Given that acylated ghrelin and PYY₃₋₃₆ function within a network of other key appetite regulatory peptides (17), 398 399 additional research is needed to characterise the impact of the present interventions on

glucagon-like-peptide-1, oxyntomodulin, pancreatic polypeptide and leptin in men comparedwith women.

402

403 The higher appetite ratings and food intake seen in men in experiment two supports the concept that lean body mass is the primary determinant of tonic appetite ratings and energy 404 intake (1). This theory is further supported by our finding that energy intake during *ad libitum* 405 feeding did not differ between sexes when expressed per kilogram of lean body mass. 406 Although acylated ghrelin may in part mediate the episodic changes in appetite observed in 407 408 the present study, the lower tonic concentrations observed in men suggests that lean body mass may influence appetite and energy intake through an alternative mechanism. Recent 409 410 evidence suggests that resting metabolic rate may be important in this regard (1).

411

412 Our findings provide a comparative insight into the short-term appetite, energy intake and gut hormone responses to acute energy deficits in women compared with men. In accordance 413 414 with the recent findings of Caudwell et al. (5), these new data support the perspective that men and women do not exhibit different physiological or behavioural compensatory 415 responses to energy deficits (induced by exercise or food-restriction); at least during the 416 actual day when an energy deficit is imposed. Our findings therefore support the importance 417 418 of exercise for weight management in women however these data must be considered in light 419 of certain limitations. Firstly, both experiment one and two were powered to detect changes in food intake and it is possible that subtle effects of the present interventions on appetite and 420 gut hormones may not have been detected. Secondly, the implementation of prolonged and 421 422 strenuous exercise protocols, completed by recreationally active individuals, may limit the generalisability of the findings i.e. to those who are less active or less fit. The arduous 423 exercise undertaken in the present studies may therefore not be achievable by many seeking 424

425 to commence a weight loss program and additional work is needed with overweight and/or426 obese participants.

427

In conclusion, the experiments presented in this paper have provided evidence that appetite, energy intake and gut hormone responses to acute energy deficits do not differ between men and women. These data support the importance of exercise for weight management in women to reduce the compensatory responses to energy deficits achieved solely via food restriction.

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439

440 CONFLICT OF INTEREST

441 All authors declare that there are no conflicts of interest. The results of the present study do

442 not constitute endorsement by ACSM.

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444

1. Blundell JE, Caudwell P, Gibbons C, et al. Role of resting metabolic rate and energy
expenditure in hunger and appetite control: a new formulation. *Dis Model Mech.* 2012;
5: 608–13.

450

451	2. Borer KT, Wuorinen E, Chao C, Burant C. Exercise energy expenditure is not

452 consciously detected due to oro-gastric, not metabolic, basis of hunger sensation.

```
453 Appetite. 2005; 45: 177–81.
```

Behav. 1995; 58: 1067-77.

454

455 3. Brechet S, Plaisancié P, Dumoulin V, Chayvialle JA, Cuber JC, Claustre J.

456 Involvement of beta1- and beta2- but not beta3-adrenoceptor activation in adrenergic

457 PYY secretion from the isolated colon. *J Endocrinol*. 2001; 168: 177–83.

458

4. Buffenstein R, Poppitt SD, McDevitt RM, Prentice AM. Food intake and the
menstrual cycle: a retrospective analysis, with implications for appetite research. *Physiol*

462

461

5. Caudwell P, Gibbons C, Finlayson G, Näslund E, Blundell J. Exercise and weight
loss: no sex differences in body weight response to exercise. *Exerc Sport Sci Rev.* 2014;
42: 92–101.

466

467 6. Caudwell P, Gibbons C, Hopkins M, King N, Finlayson G, Blundell J. No sex
468 difference in body fat in response to supervised and measured exercise. *Med Sci Sports*469 *Exerc.* 2013; 45: 351–8.

470

471	7. Deighton K, Batterham RL, Stensel DJ. Appetite and gut peptide responses to
472	exercise and calorie restriction: the effect of modest energy deficits. Appetite. 2014; 81:
473	52–9.
474	
475	8. Donnelly JE, Hill JO, Jacobsen DJ, et al. Effects of a 16-month randomized controlled
476	exercise trial on body weight and composition in young, overweight men and women:
477	the Midwest Exercise Trial. Arch Intern Med. 2003; 163: 1343–50.
478	
479	9. Faul F, Erdfelder E, Lang A-G, Buchner A. G*Power 3: a flexible statistical power
480	analysis program for the social, behavioral, and biomedical sciences. Behav Res
481	Methods. 2007; 39: 175–91.
482	
483	10. Food & Agricultural Organisation of the UN. Human Energy Requirements: Report
484	of a Joint FAO/WHO/UNU Expert Consultation. 2001.
485	
486	11. Flint A, Raben A, Blundell JE, Astrup A. Reproducibility, power and validity of
487	visual analogue scales in assessment of appetite sensations in single test meal studies.
488	Int J Obes Relat Metab Disord. 2000; 24: 38–48.
489	
490	12. Frayn KN. Calculation of substrate oxidation rates in vivo from gaseous exchange. J
491	Appl Physiol. 1983; 55: 628–34.
492	
493	13. Greenman Y, Golani N, Gilad S, Yaron M, Limor R, Stern N. Ghrelin secretion is
494	modulated in a nutrient- and gender-specific manner. Clin Endocrinol. 2004; 60: 382-8.
495	

496	14. Hagobian TA, Braun B. Physical activity and hormonal regulation of appetite: sex
497	differences and weight control. Exerc Sport Sci Rev. 2010; 38: 25-30.
498	
499	15. Hagobian TA, Sharoff CG, Stephens BR, et al. Effects of exercise on energy-
500	regulating hormones and appetite in men and women. Am J Physiol Regul Integr Comp
501	Physiol. 2009; 296: R233–42.
502	
503	16. Hagobian TA, Yamashiro M, Hinkel-Lipsker J, Streder K, Evero N, Hackney T.
504	Effects of acute exercise on appetite hormones and ad libitum energy intake in men and
505	women. Appl Physiol Nutr Metab. 2013; 38: 66–72.
506	
507	17. Hameed S, Dhillo WS, Bloom SR. Gut hormones and appetite control. Oral Dis.
508	2009; 15: 18-26.
509	
510	18. Hopkins M, Blundell JE, King NA. Individual variability in compensatory eating
511	following acute exercise in overweight and obese women. Br J Sports Med. 2013; 48: 1472-
512	1476.
513	
514	19. Irving BA, Weltman JY, Patrie JT, et al. Effects of exercise training intensity on
515	nocturnal growth hormone secretion in obese adults with the metabolic syndrome. J Clin
516	Endocrinol Metab. 2009; 94: 1979–86.
517	
518	20. King JA, Wasse LK, Ewens J, et al. Differential acylated ghrelin, peptide YY3-36,
519	appetite, and food intake responses to equivalent energy deficits created by exercise and
520	food restriction. J Clin Endocrinol Metab. 2011; 96: 1114–21.

522	21. King JA, Wasse LW, Stensel DJ, Nimmo MA. Exercise and ghrelin. A narrative
523	overview of research. Appetite. 2013; 68: 83-91.
524	
525	22. King NA, Snell L, Smith RD, Blundell JE. Effects of short-term exercise on appetite
526	responses in unrestrained females. Eur J Clin Nutr. 1996. 50: 663-667.
527	
528	23. Larson-Mayer DE, Palm S, Bansal A, Austin KJ, Marie Hart A, Alexander BM. Influence
529	of running and walking on hormonal regulators of appetite in women. J Obes. 2012. Article
530	ID 730409.
531	
532	24. Le Roux CW, Batterham RL, Avlwin SJB, et al. Attenuated peptide YY release in
533	obese subjects is associated with reduced satiety. Endocrinology. 2006; 147: 3-8.
534	
535	25. Le Roux CW, Patterson M, Vincent RP, Hunt C, Ghatei MA, Bloom SR.
536	Postprandial Plasma Ghrelin Is Suppressed Proportional to Meal Calorie Content in
537	Normal-Weight But Not Obese Subjects. J Clin Endocrinol Metab. 2005; 90: 1068-
538	1071.
539	
540	26. Long SJ, Hart K, Morgan LM. The ability of habitual exercise to influence appetite and
541	food intake in response to high-and low-energy preloads in man. Br J Nutr. 2002. 87: 517-
542	523.
543	
544	27. Martins C, Truby H, Morgan LM. Short-term appetite control in response to a 6-week
545	exercise program in sedentary volunteers. Br J Nutr. 2007. 98: 834-842.

547	28. Mifflin MD, St Jeor ST, Hill LA, Scott BJ, Daugherty SA, Koh YO. A new
548	predictive equation for resting energy expenditure in healthy individuals. Am J Clin
549	Nutr. 1990; 51: 241–7.
550	
551	29. Rowell LB. Human cardiovascular adjustments to exercise and thermal stress.
552	<i>Physiol Revs.</i> 1974; 54: 75–159.
553	
554	30. Schubert MM, Sabapathy S, Leveritt M, Desbrow B. Acute exercise and hormones
555	related to appetite regulation. Sports Med. 2014; 44: 387-403.
556	
557	31. Stensel DJ. Exercise, appetite and appetite regulating hormones: implications for
558	food intake and weight control. Ann Nutr Metab. 2010; 57 (Supplement 2): 36-42.
559	
560	32. Stubbs RJ, Hughes DA, Johnstone AM, et al. The use of visual analogue scales to
561	assess motivation to eat in human subjects: a review of their reliability and validity with
562	an evaluation of new hand-held computerized systems for temporal tracking of appetite
563	ratings. Br J Nutr. 2000; 84:405-15.
564	
565	33. Stunkard AJ, Messick S. The three-factor eating questionnaire to measure dietary
566	restraint, disinhibition and hunger. J Psychosom Res. 1985; 29: 71-83.
567	
568	34. Taylor Hl, Buskirk E, Henschel A. Maximal oxygen intake as an objective measure
569	of cardio-respiratory performance. J Appl Physiol. 1955; 8: 73-80.
570	

571	35. Westerterp KR, Meijer GA, Janssen EM, Saris WH, Ten Hoor F. Long-term effect
572	of physical activity on energy balance and body composition. Br J Nutr. 1992; 68: 21-
573	30.
574	
575	36. Wijlens AG, Erkner A, Alexander E, Mars M, Smeets PA, de Graaf C. Effects of oral and
576	gastric stimulation on appetite and energy intake. Obesity. 20: 2226-2232.
577	
578	37. Yang J, Brown MS, Liang G, Grishin NV, Goldstein JL. Identification of the
579	acyltransferase that octanoylates ghrelin, an appetite-stimulating peptide hormone. Cell.
580	2008; 132: 387–96.
581	
582	38. Zhang T, Uchida T, Gomez G, Lluis F, Thompson JC, Greeley GH. Neural
583	regulation of peptide YY secretion. Regul Pept. 1993; 48: 321-8.
584	
505	
585	
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Table 1: Participant characteristics in experiment one and two

	Experiment 1	Experi	nent 2
	(Females)	(Females)	(Males)
Participant number (n)	12 [‡]	10 ⁺	10 ⁺
Age (y)	22.4 (2.1)	22.3 (2.5)	22.6 (3.8)
Height (cm)	165.6 (5.4)	166.6 (5.4)	180.5 (6.2)*
Body mass (kg)	60.4 (4.2)	61.9 (7.3)	75.4 (9.4)*
BMI (kg/m²)	22.0 (1.1)	22.3 (2.32)	23.1 (2.1)
Body Fat (%)	24.1 (2.8)	22.4 (5.5)	10.1 (4.2)*
Lean mass (kg)	45.9 (3.7)	47.4 (1.4)	67.5 (3.3)*
VO2 max (mL/kg/min)	50.4 (4.3)	48.8 (6.1)	66.1 (9.2)*

*significantly different between males and females (P < 0.005) [†]acylated ghrelin and PYY₃₋₃₆ data available for 11 participants

⁺ acylated ghrelin data available for 8 participants

Table 2. Energy and macronutrient intakes of men and women during the buffet meal in the controland exercise trials.

	Cor	ntrol	Exercise	
	Men	Women	Men	Women
Fat (kJ)	355 ± 274	175 ± 142	348 ± 245	168 ± 142
Fat (kJ.kg lean mass ⁻ ¹)	5 ± 5	4 ± 3	5 ± 3	4 ± 3
Carbohydrate (kJ) [‡]	680 ± 318	434 ± 174	788 ± 322	446 ± 201
Carbohydrate (kJ.kg lean mass ⁻¹)	10 ± 6	9 ± 4	12 ± 6	10 ± 5
Protein (kJ)	148 ± 111	87 ± 75	149 ± 100	95 ± 68
Protein (kJ.kg lean mass ⁻¹)	2 ± 1	2 ± 1	2 ± 1	2 ± 1
Energy intake (kJ) [‡]	4971 ± 2644	2916 ± 1506	5385 ± 2423	2979 ± 1586
Energy intake (kJ.kg lean mass ⁻¹)	75 ± 38	63 ± 25	84 ± 38	63 ± 38

604 Values are mean (SD). Females n=10; males n=10. \ddagger Significantly higher in men than women (P <

605 0.05).

607 FIGURE CAPTIONS

Figure 1. Time-averaged appetite (a), circulating acylated ghrelin (b) and peptide YY_{3-36} (c) AUC for each 9 h trial. *Food-Def significantly different from Ex-Def and control; † Ex-Def significantly different from Food-Def and control (experiment one – female participants only). Values are mean (SEM), N = 12 for appetite and 11 for acylated ghrelin and peptide YY₃₋₃₆.

Figure 2. Appetite (a), circulating acylated ghrelin (b) and peptide YY_{3-36} (c) concentrations across the Con (∇), Ex-Def (\bullet) and Food-Def (\circ) trials (experiment one – female participants only). Hatched shaded rectangles indicate standardised test meals, lightly shaded rectangle indicates exercise, black rectangle indicates *ad libitum* meal. Values are mean (SEM), N = 12 for appetite and 11 for acylated ghrelin and peptide YY₃₋₃₆.

Figure 3. (a) Appetite ratings in Male Con (\circ), Male Ex (\bullet), Female Con (∇) and Female Ex ($\mathbf{\nabla}$) (experiment two – male and female participants). Hatched shaded rectangles indicate standardised test meal, lightly shaded rectangle indicates exercise, black rectangle indicates *ad libitum* meal. (b) Time-averaged appetite AUC for each 7 h trial. ‡ Males significant different than females. § Control significantly different than exercise. Values are mean (SEM). Females N=10; males N=10.

Figure 4. (a) Plasma acylated ghrelin concentrations in Male Con (\circ), Male Ex (\bullet), Female Con (∇) and Female Ex ($\mathbf{\nabla}$) (experiment two – male and female participants). Hatched shaded rectangles indicate standardised test meal, lightly shaded rectangle indicates exercise, black rectangle indicates *ad libitum* meal. (b) Time-averaged acylated ghrelin AUC for each 7 h trial. ¶ Females significantly different than males. § Control significantly different than exercise. Values are mean (SEM). Females N=8; males N=8.







