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Title: A randomized cross-over trial assessing the effects of acute exercise on appetite, circulating ghrelin concentrations and butyrylcholinesterase activity in normal weight males with variants of the obesity-linked *FTO* rs9939609 polymorphism.

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Short running title: Ghrelin, exercise and *FTO* rs9939609 genotype.

Abbreviations: AG, acyl-ghrelin; AUC, area under the curve; BChE, butyrylcholinesterase; BMI, body mass index; CI, confidence interval; DAG, des-acyl-ghrelin; ES, effect size; *FTO*, the fat mass and obesity-associated gene; GLP-1, glucagon-like peptide 1; PYY, peptide YY; SD, standard deviation; SEM, standard error of mean; SNP, single nucleotide polymorphism.

ClinicalTrials.gov registration: NCT03025347

Data described in the manuscript will be made available upon request pending application and approval.

1 Abstract

Background: The fat mass and obesity-associated gene (*FTO*) rs9939609 A-allele is
associated with higher acyl-ghrelin (AG) concentrations, higher energy intake and obesity,
though exercise may mitigate rs9939609 A-allele linked obesity risk. Butyrylcholinesterase
(BChE) hydrolyses AG to des-acyl-ghrelin (DAG), potentially decreasing appetite. However,
the effects of the *FTO* rs9939609 genotype and exercise on BChE activity, AG, DAG and
energy intake are unknown.

8 **Objective:** We hypothesized that individuals homozygous for the obesity-risk A-allele (AAs) 9 would exhibit higher postprandial AG and energy intake than individuals homozygous for the 10 low obesity-risk T-allele (TTs), but that exercise would increase BChE activity and diminish 11 these differences.

Methods: Twelve AA and 12 TT normal weight males completed a control (8 hours rest) and an exercise (1 hour of exercise at 70% peak oxygen uptake, 7 hours rest) trial in a randomized cross-over design. A fixed meal was consumed at 1.5 hours and an *ab libitum* buffet meal at 6.5 hours. Appetite, appetite-related hormones, BChE activity and energy intake were assessed.

17 **Results**: AAs displayed lower baseline BChE activity, higher baseline AG/DAG ratio,

18 attenuated AG suppression after a fixed meal and higher *ad libitum* energy intake than TTs

19 (ES \ge 0.76, P \le 0.049). Exercise increased delta BChE activity in both genotypes (ES = 0.41,

20 P = 0.004); however, exercise lowered AG and the AG/DAG ratio to a greater extent in AAs

21 ($P \le 0.041$), offsetting the higher AG ghrelin profile observed in AAs during the control trial

22 (ES \ge 0.88, P \le 0.048). Exercise did not elevate energy intake in either genotype (P = 0.282).

23 Conclusions: Exercise increases BChE activity, suppresses AG and the AG/DAG ratio and

24 corrects the higher AG profile observed in obesity-risk AA individuals. These findings

- 25 suggest that exercise or other methods targeting BChE activity may offer a preventative
- 26 and/or therapeutic strategy for AA individuals.
- 27
- 28 **Keywords:** exercise; ghrelin; appetite; *FTO* gene; butyrylcholinesterase; obesity

29 INTRODUCTION

5

A cluster of single nucleotide polymorphisms (SNP) within intron one of the fat mass and 30 31 obesity-associated gene (FTO) have been consistently associated with obesity (1-3). At the 32 FTO rs9939609 SNP, homozygous obesity-risk A-allele carriers (AA) have a 1.7-fold higher 33 risk for obesity compared to individuals homozygous for the T-allele (TT) (1). Compared 34 with TTs, AA individuals exhibit lower postprandial satiety and higher energy intake (4-6). 35 Karra et al. (7) also reported that AAs displayed an attenuated postprandial suppression of the 36 orexigenic hormone acyl-ghrelin (AG) and appetite compared to TTs. These findings suggest 37 the impaired postprandial suppression of AG might contribute to the higher energy intake and 38 obesity risk in AAs.

39 Acute bouts of moderate- to vigorous-intensity exercise acutely suppress both subjective 40 appetite perceptions and circulating AG concentrations (8,9). In addition, circulating 41 concentrations of the anorectic hormones PYY and GLP-1 are increased by a single exercise 42 bout (9,10). These gut hormone changes are suggested to provoke the acute anorectic effect 43 of exercise (8,9,11). Further to changes during the exercise bout, circulating AG 44 concentrations remain suppressed while PYY and GLP-1 are elevated in the hours after 45 exercise (8,9,11). Importantly, the lack of compensatory changes in hunger and appetite-46 related hormones to an energy shortfall caused by exercise results in a short-term negative 47 energy balance, which if sustained, could facilitate weight management (12).

The serine hydrolase butyrylcholinesterase (BChE) regulates circulating ghrelin
concentrations by hydrolyzing AG to des-acyl-ghrelin (DAG), which is suggest to have an
anorexigenic effect (13). Recent studies indicate that reduced BChE activity leads to a higher
AG/DAG ratio, greater food consumption and weight gain (14,15). However, less is known
about the interplay between BChE, *FTO* rs9939609 and exercise in humans. One study

indicated that a single bout of light running increases BChE activity in humans (16), but
further work is needed to examine if BChE activity is linked to *FTO* rs9939609 genotype and
exercise-dependent changes in plasma ghrelin concentrations or appetite-related outcomes in
humans.

57 Our primary aim was to investigate the effect of the *FTO* rs9939609 genotype and exercise 58 on circulating AG and DAG concentrations, BChE activity, appetite and energy intake in a 59 group of normal-weight AA males and a matched-group of TT males. As a secondary aim, 60 we examined the effect of exercise and/or the *FTO* rs9939609 genotype on plasma 61 concentrations of leptin, PYY and GLP-1. We hypothesized that AAs would exhibit higher 62 AG, appetite and energy intake compared to TTs, but exercise would increase BChE activity 63 and suppress these rs9939609-related differences.

64 PARTICIPANTS AND METHODS

65 Participants

66 The study was performed according to the principles set out in the Declaration of Helsinki 67 and was approved by the Loughborough University ethical advisory committee. We recruited 68 202 healthy, non-smoking males aged 18-50 y of mixed European descent who provided 69 written informed consent to take part in a database study. Exclusion criteria were history of 70 cardio-metabolic disease, medical or psychiatric conditions, substance abuse and food 71 allergies. Participants' height and body mass were measured, and waist circumference was 72 assessed as the narrowest portion of the torso between the xiphoid process and the naval. 73 Skinfold thickness was measured and body fat percentage was estimated (17). Habitual 74 physical activity levels were assessed using the short form International Physical Activity 75 Questionnaire (18) and eating behaviors and attitudes were assessed using the Three-Factor 76 Eating Questionnaire (19). A venous blood sample was collected and DNA was extracted. All 77 DNA extractions from peripheral blood samples were performed using the QIAamp DNA 78 Blood Midi Kit (Qiagen). Genotyping for rs9939609 was performed by LGC Limited 79 (Hertfordshire, UK) using the KASP (KBioscience Competitive Allele-Specific PCR) SNP 80 genotyping system (www.lgcgenomics.com/genotyping/kasp-genotyping-reagents/). Blind 81 duplicates were used to detect possible DNA mix-up. From the database, we recruited a 82 group of 12 AA and 12 TT participants (Table 1) for a randomized cross-over study 83 (Supplementary Figure 1). Participants provided written informed consent if they were 84 invited back and completed the study between January 2015 to February 2016. Further to the 85 criteria mentioned, to be included in this trial, participants had to be weight stable ($\leq 3 \text{ kg}$) 86 over previous 3 months) and habitually consumed breakfast on 5 or more days of the week in 87 an attempt to reduce the influence of breakfast consumption on fasting ghrelin concentrations 88 (20). Participants were also excluded if they presented any food allergies. Groups were 89 matched for anthropometric indices, age and peak oxygen uptake (Table 1). The study is registered at clinicaltrials.gov as NCT03025347. 90

91 Main trials

92 Participants attended a preliminary measures and familiarization session prior to main trials. 93 Body mass, height, body fat percentage, body mass index (BMI) and waist circumference 94 were re-measured as described to confirm no substantial changes occurred from the database 95 study. Participants performed submaximal incremental and peak oxygen uptake running tests 96 on a motorized treadmill as described elsewhere (8). Individual running speed-oxygen uptake 97 linear regression equations and peak oxygen uptake were used to calculate the running speed 98 that corresponded to 70% of each participant's peak oxygen uptake. Participants also 99 completed a food preference questionnaire and were familiarized with the buffet meal, to 100 reduce the risk of any changes in food intake due to novelty of the meal.

101 Next, in a randomized cross-over design stratified by rs9939609 genotype group, all 102 participants completed two main trials separated by 7-14 days: exercise and control. Further 103 to enrolling participants, the main investigator conducted the block randomization plan for 104 each genotype from the website www.randomization.com and assigned participants to the 105 order of trials completed. Participants were instructed to complete a weighed food diary in 106 the 24 h before the first trial and replicate it in the 24 h before the second trial. Participants 107 were also instructed to refrain from alcohol consumption and strenuous physical activity in 108 this period. A pizza meal (5201 kJ) was consumed by participants between 19:00-20:00 the 109 night before main trials to negate the influence of preceding food intake on morning appetite 110 and appetite-related hormone concentrations (21). Adherence to these procedures was 111 assessed by verbal confirmation.

112 A schematic representation of the main trial procedures is shown in **Figure 1**. Participants 113 arrived at the laboratory at approximately 08:30 after an overnight fast. A cannula was 114 inserted into an antecubital vein 60 min before blood sampling commenced to mitigate any 115 stress response caused by anxiety with the cannula (21). In the control trial, participants 116 rested for 8 h, while in the exercise trial, participants ran at 70% of peak oxygen uptake for 117 60 min and then rested for 7 h. Participants read, worked and watched TV through laptop and 118 tablet devices while resting. Expired gas samples were collected into Douglas bags every 15 119 min throughout the first hour in both trials for calculation of energy expenditure (22).

120 Fixed test meal and buffet meal

121 Participants consumed a standardized 5623 kJ (52% carbohydrate, 25% fat, 23% protein) test

122 meal consisting of white rolls, butter, cheese, chips, chocolate slices and milkshake at 1.5 h.

123 Participants were instructed to consume the meal within 20 minutes.

124 At 6.5 h, participants were provided with a buffet meal in a booth and instructed to eat ad libitum. Food items of the buffet meal were presented identically on each trial and included 125 126 white and brown bread, butter, chicken, ham, lettuce, tomato, yoghurts, cookies and apples. 127 Participants were instructed to eat until "comfortably full and satisfied" before leaving the eating booth. To minimize distractions that may influence food consumption, the buffet was 128 129 provided in isolation and participants were not permitted the use of mobile phones or 130 electronic devices. Items were provided in excess of expected consumption and participants 131 were provided with more food items if requested. The amount of each food item consumed 132 was calculated by measuring the weighted difference of all the food items before and after the meal. Manufacturer details were used to determine energy and macronutrient consumption. 133

134 Appetite ratings

- 135 Visual analogue scales (VAS) were used to assess subjective feelings of hunger, fullness,
- 136 prospective food consumption and hedonic wanting of food (23,24). Measures were taken
- 137 every 30 min from baseline to 5.0 h, and then at 6.5, 7.0, 7.5 and 8.0 h.

138 Blood sampling

- 139 Blood samples were collected into chilled EDTA monovettes (Sarstedt, Leicester, UK) every
- 140 30 min from baseline to 4.0 h and subsequently at 5.0, 6.5 and 7.5 h to measure circulating
- 141 concentrations of AG, DAG, total PYY and total GLP-1. Circulating leptin was measured
- 142 from fasting samples only. Plasma BChE activity was determined from samples collected at
- 143 0, 0.5 and 1 h in the control and exercise trials. All collected samples were immediately
- 144 centrifuged at 2383g for 10 min at 4°C. After centrifugation, 100 µL of 0.5 mol/L
- 145 hydrochloric acid was added per 900 µL of plasma supernatant to preserve DAG. To preserve
- 146 the stability of AG, one monovette was treated with a 50 µL solution of PBS, P-
- 147 hydroxymercuribenzoic acid and sodium hydroxide. The plasma supernatant of this sample

148	was dispensed into a storage tube and 100 μL of 1 mol/L hydrochloric acid was added per 1
149	ml of plasma. All samples were stored at -80°C until batch analysis.

150 **Biochemical analysis**

- 151 Enzyme-linked immunosorbent assays were used to measure circulating concentrations of
- 152 AG, DAG (SCETI, Tokyo, Japan), total PYY, total GLP-1 (Millipore, Watford, UK) and
- leptin (R&D Systems, Abington, UK). The intra-assay variability was 4.3%, 3.5%, 1.9%,

154 3.6% and 1.8% for AG, DAG, total PYY, total GLP-1 and leptin, respectively.

- Details of BChE analysis are documented in the Supplementary Methods. In short, BChE
 assays were performed based upon the cholinesterase assay method developed by Ellman
- 157 (25), with butyrylthiocholine iodide as the enzymatic substrate.

158 Statistical analyses

159 A sample size of 24 was chosen based on data suggesting that a 10 pmol/L reduction in 160 circulating AG during exercise could be detected with > 80% power using a two-tailed *t*-test 161 whilst assuming a SD_{diff} of 16 pmol/L and adopting an alpha value of 0.05 (26). Primary 162 outcomes measured in this trial were AG, DAG, BChE activity, appetite and *ad libitum* 163 energy intake, and secondary outcomes were total GLP-1, total PYY and leptin. To reduce 164 day-to-day variability, appetite-related hormone concentrations and BChE were analyzed and 165 presented as delta values. Appetite ratings, appetite-related hormone concentrations and BChE activity were analyzed using linear mixed models with trial (exercise or control), 166 167 genotype (AA or TT) and time included as fixed factors. Total area under the curve (AUC) was calculated using the trapezoidal rule. For blood parameters, AUC was calculated during 168 169 the intervention (0.0-1.0 h), post-test meal (1.5-3.5 h), afternoon (3.5-6.5 h) and post-buffet 170 meal (6.5-7.5 h) periods. AUC for subjective appetite ratings was calculated during the

171 intervention (0.0-1.0 h), post-test meal (1.5-3.5 h), afternoon (3.5-6.5 h) and post-buffet meal 172 (6.5-8.0 h) periods. Linear mixed models were used for trial and genotype comparisons of AUC values and food consumption at the buffet meal. Post-hoc analysis was conducted using 173 174 Holm-Bonferroni correction for multiple comparisons. Absolute standardized effect sizes (ES) were calculated by dividing the difference between the mean values (exercise vs. control 175 176 or AAs vs. TTs) with the pooled standard deviation. An ES of 0.2 was considered the 177 minimum important difference for all outcome measures, 0.5 moderate and 0.8 large (27). 178 The 95% confidence intervals (CI) for mean absolute pairwise differences between 179 experimental trials or genotype groups were calculated. Statistical significance was accepted 180 as P < 0.05. Linear mixed models were conducted with trial order as a fixed effect which 181 revealed no main or interactive effects for any outcome ($P \ge 0.073$; data not shown). Unless 182 stated otherwise, data presented in tables and figures are shown as mean \pm SEM, while 183 descriptive data are presented as mean \pm SD. Data were analyzed using IBM SPSS Statistics 184 for Windows software (version 23.0, IBM corporation, New York, USA).

185 **RESULTS**

186 **Participant characteristics**

187 There were no differences between AAs and TTs for age, height, body mass, BMI, body fat

188 %, lean body mass, waist circumference, eating behaviors, habitual physical activity levels or

189 peak oxygen uptake ($P \ge 0.121$) (Table 1). There were no differences in energy intake

190 between AAs and TTs in the 24 h before the main trials (AA: 9516 ± 595 kJ vs TT: $9630 \pm$

191 891 kJ; P = 0.716).

192 Treadmill running responses

193 We observed no between-genotype differences in exercise responses for running speed (AA:

194 11.1 ± 1.5 vs. TT: 11.3 ± 1.6 km/h; P = 0.786), heart rate (AA: 178 ± 13 vs. TT: 177 ± 12

195 beats/min; P = 0.934), gross energy expenditure (AA: 3809 ± 366 vs. TT: 3568 ± 249 kJ; P =

196 0.117) or percentage of peak oxygen uptake (AA: 71 ± 2 vs. TT: $70 \pm 2\%$; P = 0.283).

197 Circulating appetite-related hormones and BChE activity

198 Fasting concentrations of AG, DAG, total GLP-1, total PYY and leptin at baseline were not

199 different between genotype groups ($P \ge 0.127$) or between trials ($P \ge 0.259$) (**Table 2**). The

200 fasting AG/DAG ratio and BChE activity were similar between trials (P > 0.319), but the

201 AG/DAG ratio and BChE were higher and lower, respectively, in AAs than TTs (ES \ge 0.76,

202 $P \le 0.047$) (Table 2).

203 Linear mixed models for delta AG identified a main effect of trial (P < 0.001) and time (P < 0.001)

0.001) but not genotype (mean difference: -0.02 pmol/L, 95% CI -2, 2 pmol/L, P = 0.988)

205 (Figure 2A). The main effect of trial revealed lower delta AG concentrations in the exercise

than control trial (mean difference: -5 pmol/L, 95% CI -6, -5 pmol/L, ES = 0.79). Analysis

207 also identified a genotype-by-time interaction (P = 0.007), but post-hoc analysis revealed no

208 differences after Holm-Bonferroni adjustment ($P \ge 0.060$). The AUC for delta AG was lower

in the exercise than control trial during the intervention (0.0-1.0 h), post-test meal (1.5-3.5 h)

and afternoon (3.5-6.5 h) periods (all ES \ge 0.53, P \le 0.001) (**Table 3**). The magnitude of

211 reduction in AUC for delta AG after exercise was greater in AAs than TTs during the post-

212 test meal period (1.5-3.5 h; -23.98 pmol/L \cdot h (ES = 2.49) vs. -14.3 pmol/L \cdot h (ES = 1.62),

respectively; genotype-by-trial interaction P = 0.041) (Table 3). Post-hoc analysis of the post-

test meal period revealed higher AUC delta AG in AAs compared to TTs in the control trial

215 (ES = 1.25, P = 0.011), but no between-genotype differences were seen in the exercise trial 216 (ES = 0.03, P = 0.951).

217	There was a main effect of trial ($P < 0.001$) and time ($P < 0.001$) but not genotype (mean
218	difference: 9 pmol/L, 95% CI -5, 24 pmol/L, $P = 0.197$) for delta DAG (Figure 2B). The
219	main effect of trial revealed lower delta DAG concentrations in the exercise than control trial
220	(mean difference: -17 pmol/L, 95% CI -20, -14 pmol/L, $ES = 0.44$). The magnitude of
221	reduction in delta DAG concentrations after exercise was greater in TTs than AAs (-25
222	pmol/L (ES = 0.58) vs9 pmol/L (ES = 0.26), respectively; genotype-by-trial interaction P $<$
223	0.001). The AUC for delta DAG was lower in the exercise than control trial during the
224	intervention (0.0-1.0 h), post-test meal (1.5-3.5 h) and afternoon (3.5-6.5 h) periods (all ES \geq
225	0.29, P \leq 0.028) (Table 3). The magnitude of reduction in AUC for delta DAG after exercise
226	was greater in TTs than AAs during the intervention period (0.0-1.0 h; -82.4 pmol/L·h (ES =
227	2.47) vs46.2 pmol/L·h (ES = 1.66), respectively; genotype-by-trial interaction $P = 0.042$)
228	and post-test meal period (1.5-3.5 h; -100.8 pmol/L \cdot h (ES = 1.66) vs39.0 (ES = 0.59),
229	respectively; genotype-by-trial interaction $P = 0.025$) (Table 3).
230	Linear mixed models for the delta AG/DAG ratio identified a main effect of trial ($P < 0.001$)
231	and time (P < 0.001) but not genotype (mean difference: -0.006, 95% CI -0.015, 0.003, P =
232	0.192) (Figure 2C). The main effect of trial revealed the delta AG/DAG ratio was lower in
233	the exercise than control trial (mean difference: -0.025 , 95% CI -0.029 , -0.022 , ES = 0.88).
234	The magnitude of reduction in the delta AG/DAG ratio after exercise was greater in AAs than
235	TTs at time points between 0.5 h to 2.5 h (genotype-by-trial-by-time interaction, $P = 0.004$).
236	The AUC for the AG/DAG ratio was lower in the exercise than control trial during the
237	intervention, post-test meal, and post-buffet meal periods (all ES \ge 0.54, P \le 0.006) (Table 3).
238	The magnitude of reduction in AUC for the delta AG/DAG ratio after exercise was greater in

239 AAs than TTs during the intervention period (0.0-1.0 h; -0.12 (ES = 5.18) vs. -0.07 (ES = 240 1.63), respectively; genotype-by-trial interaction P = 0.004) and post-test meal period (1.5-3.5 h; -0.16 (ES = 2.72) vs. -0.02 (ES = 0.28), respectively; genotype-by-trial interaction P =241 242 0.001) (Table 3). Post-hoc analysis of the intervention period revealed a similar AUC delta 243 AG/DAG ratio between groups in the control trial (ES = 0.27, P = 0.518), but the AG/DAG 244 ratio was lower in AAs compared to TTs in the exercise trial (ES = 1.75, P < 0.001). Post-hoc 245 analysis in the post-test meal period indicated that AAs exhibited higher AUC delta AG/DAG in the control trial (ES = 0.88, P = 0.048) but lower AUC delta AG/DAG in the exercise trial 246 247 (ES = 1.17, P = 0.018) compared to TTs.

There was a main effect of trial (P < 0.001) and time (P < 0.001) but not genotype (mean

difference: 2 pmol/L, 95% CI -2, 7 pmol/L, P = 0.335) for delta total GLP-1 (**Figure 3A**).

250 The main effect of trial revealed higher delta total GLP-1 concentrations in the exercise than

control trial (mean difference: 14 pmol/L, 95% CI 12, 15 pmol/L, ES = 1.14). Analysis also

identified a genotype-by-time interaction (P = 0.002), but post hoc analysis showed no

253 differences after Holm-Bonferroni adjustment ($P \ge 0.092$). The AUC for delta total GLP-1

254 was higher in the exercise than control trial during all time periods (all ES \ge 0.50, P \le 0.044),

and higher in AAs than TTs during the post-buffet meal period (6.5-7.5 h; ES = 0.92, P =

256 0.011) (**Table 4**).

A main effect of trial (P < 0.001) and time (P < 0.001) but not genotype (mean difference: 10 pmol/L, 95% CI -9, 29 pg/mL, P = 0.278) was detected for delta total PYY (**Figure 3B**). The main effect of trial revealed higher delta total PYY concentrations in the exercise than control trial (mean difference: 25 pg/mL, 95% CI 20, 30 pmol/L, ES = 0.50). The AUC for delta total PYY was higher in the exercise than control trial during the intervention (0.0-1.0 h; ES = 262 3.08, P < 0.001) and post-test meal (1.5-3.5 h; ES = 1.56, P < 0.001) periods, and higher in AAs than TTs during the post-buffet meal period (6.5-7.5 h; ES = 0.78, P = 0.029) (Table 4). 263 264 Analysis for delta BChE identified a main effect of time (P < 0.001) and trial (P = 0.004), 265 with elevated BChE activity in the exercise trial compared to the control trial (mean 266 difference: 0.072 KU/L, 95% CI 0.024, 0.120 KU/L, ES = 0.41) (Figure 4). There was, 267 conversely, no main effect of genotype (mean difference: -0.016 KU/L, 95% CI -0.095, 268 0.063, ES = 0.09, P = 0.681), and no two-way or three-way interactions for BChE activity (P 269 \geq 0.094) (Figure 4).

270 Appetite ratings

271 Linear mixed models for each appetite perception identified a main effect of trial (P = 0.002) 272 and time (P < 0.001) but not genotype ($P \ge 0.072$) (Figure 5). The main effect of trial for 273 each perception revealed suppressed appetite in the exercise compared with the control trial 274 (all $ES \ge 0.12$). Analysis also identified a genotype-by-time interaction for each appetite 275 perception (P < 0.001) (Figure 5). Post-hoc analysis of the genotype-by-time interaction 276 revealed higher ratings of hunger and hedonic wanting of food and lower ratings of fullness 277 in AAs than TTs at time points between 3.0 to 4.0 h (all ES \ge 1.04, P \le 0.033). There were no 278 between-genotype differences at any time point for prospective food consumption after 279 Holm-Bonferroni correction ($P \ge 0.130$). A main effect of trial for AUC values in the 280 intervention period (0.0-1.0 h) revealed lower ratings of hunger, prospective food 281 consumption and hedonic wanting of food and higher ratings of fullness in the exercise than 282 control trial (all ES \geq 1.14, P < 0.001) (**Table 5**). A main effect of genotype for AUC values 283 in the post-test meal (1.5-3.5 h) and afternoon (3.5-6.5 h) periods revealed higher ratings of hunger, prospective food consumption and hedonic wanting of food but lower ratings of 284 285 fullness in AAs than TTs (all ES \ge 0.81, P \le 0.045) (Table 5).

286 **Buffet meal**

287 Absolute energy intake was greater in AAs than TTs (ES = 0.86, P = 0.049), but was similar 288 between the exercise and control trials (P = 0.282) (**Table 6**). Relative energy intake was 289 substantially lower in the exercise than control trial (ES = 1.84, P < 0.001), and tended to be 290 greater in AAs than TTs (ES = 0.80, P = 0.081). Protein intake was higher in AAs than TTs 291 (ES = 0.93, P = 0.033), and intakes of carbohydrate (ES = 0.73, P = 0.075) and fat (ES = 292 0.82, P = 0.072) were meaningfully, albeit not statistically, greater in AAs than TTs. Linear 293 mixed models revealed no genotype-by-trial interactions for energy or macronutrient intakes 294 (P > 0.207).

295 **DISCUSSION**

296 The primary findings of this study are that normal weight males homozygous for the obesity-297 risk FTO rs9939609 A-allele displayed lower fasting BChE activity and higher postprandial 298 AG and AG/DAG ratio which coincided with higher postprandial appetite and *ad libitum* 299 energy intake compared to TTs. A single bout of exercise increased BChE activity and 300 suppressed circulating AG. Importantly, the exercise-induced suppression of the AG/DAG 301 ratio was greater in AA versus TT individuals, negating the differences in ghrelin seen in the 302 control trial. Exercise transiently suppressed appetite and did not lead to compensatory 303 increases in appetite or energy intake after the test meal in either genotype group.

Elevated AG and AG/total ghrelin ratio profiles in AAs have been implicated in their higher obesity risk (7,28). More recently, DAG has been shown to antagonize the orexigenic effects of AG, and the AG/DAG ratio has been suggested as a key determinant of appetite, energy intake and body weight (29,30). Thus, our novel finding of a higher AG/DAG ratio in AAs compared to TTs supports the concept that ghrelin may play an aetiopathogenic role in the higher energy intake and obesity-risk associated with the A-allele of rs9939609. However, we 310 showed that exercise suppresses AG and the AG/DAG ratio and offsets these rs9939609 311 genotype differences. An acute reduction in AG during exercise has been shown before (8), 312 but our study is the first to show differences between AA and TT individuals during exercise 313 and immediately after the test meal. Specifically, in response to exercise, we found a greater 314 reduction in the AG/DAG ratio during the exercise intervention period, and in AG and the 315 AG/DAG ratio after provision of the test meal (1.5-3.5 h) in AAs compared with TTs. 316 Physical activity attenuates the effect of rs9939609 A obesity-risk allele on adiposity (31), 317 but our study may offer insights into the mechanisms of this genotype-lifestyle interaction 318 (31). That is, the greater exercise-induced suppression of AG and the AG/DAG ratio in AAs 319 could partly explain the greater weight loss seen in carriers of the risk genotype with exercise 320 interventions (32,33).

321 The elevation in BChE activity in response to exercise supports previous findings suggesting 322 that an acute bout of walking/running elevated plasma BChE activity (16). The mechanisms 323 underlying this response require further study, though it may be that the transient increase in 324 inflammatory markers could be implicated (34). It is possible that the elevation in BChE 325 activity during exercise increased AG hydrolysis to DAG, providing a plausible mechanism 326 for the exercise-induced reduction of plasma AG concentrations. However, we also showed 327 that plasma DAG concentrations were suppressed during exercise, indicating that an 328 attenuation of ghrelin release may also be implicated in response to exercise. Therefore, it is 329 likely that several mechanisms are involved in the exercise-stimulated suppression of AG.

Another novel finding of lower fasting BChE activity in AA compared to TT individuals
offers a potential explanation for the higher AG/DAG ratio and energy intake observed in AA
versus TT individuals. BChE activity increases AG hydrolysis in plasma, leading to greater
DAG and a lower AG/DAG ratio, which has been linked to lower energy consumption and

lower adiposity in mice (14). In contrast to our findings, the *FTO* rs9939609 A-allele has
previously been associated with higher BChE activity, yet this relationship was diminished
when BMI was controlled (35). The careful matching of AAs and TTs in our study may have
improved the sensitivity to detect differences in the *FTO* rs9939609 genotype, particularly as
age, sex, substance abuse, physical activity and smoking have been shown to affect BChE
activity (36,37).

340 Our findings may expound a complex set of mechanisms that link FTO and obesity. FTO 341 encodes FTO protein, which demethylates the nucleoside N6-methyladenosine in RNA and, 342 in turn, regulates mRNA export, RNA metabolism and RNA splicing (7,38). Ghrelin, ghrelin-343 O-acyltransferase and BChE mRNA have all been identified as targets for FTO 344 demethylation and this could offer a mechanistic link between FTO rs9939609 and our 345 findings (7). Indeed, AAs have been reported to exhibit higher FTO protein expression 346 compared to TTs, indicating a potential direct mechanistic link between rs9939609 A-allele, 347 the FTO protein, circulating ghrelin, lower BChE activity, higher energy intake and obesity. 348 Taken together, this could suggest that therapeutic interventions augmenting BChE activity 349 may offer a potential strategy that could assist with weight management in AA individuals.

350 Acute studies report that appetite is transiently suppressed during exercise and compensatory 351 changes in these perceptions and energy intake do not occur (8–10). Our results are 352 consonant with these findings, and we demonstrated that the appetite suppression during 353 exercise was comparable in AAs and TTs and ad libitum energy intake was unaltered after 354 exercise in both genotype groups. We also showed that AAs exhibited greater perceptions of 355 appetite in the 4.5 hours after the test meal and consumed a higher energy intake and protein 356 at the buffet meal. Our results are in agreement with studies indicating that individuals with 357 the A-allele of rs9939609 exhibit reduced satiety (4,7,39), higher food intake (5,6) and

elevated protein intake (40). It seems likely that the greater postprandial appetite displayed by AAs plays a role in the higher energy intake exhibited by this group. The *FTO*-linked change in protein consumption could be related to the role *FTO* plays in sensing amino acids (41). It is, nevertheless, noteworthy that there was a tendency for AA individuals to consume more carbohydrate and fat at the buffet meal. This indicates that the *FTO* rs9939609 A-allele is associated with a higher intake of all macronutrients and this may have been detected with a larger sample size.

In line with previous studies, total GLP-1 and total PYY concentrations were elevated during 365 366 and immediately after exercise (9,11), and this rise was similar in AAs and TTs. At most 367 periods of the day, concentrations of the satiety hormones, leptin, total GLP-1 and total PYY were not influenced by the FTO rs9939609 variant, supporting previous research (7). The 368 369 only exception was after the buffet meal, where the elevations in total GLP-1 and total PYY 370 were greater in AAs than TTs. However, rather than any effect of the FTO rs9939609 variant, 371 this is likely to reflect the greater energy and protein intake seen in AAs at the buffet meal 372 (42,43). Our data therefore bolster evidence suggesting that AAs and TTs exhibit no 373 differences in circulating PYY and GLP-1 concentrations after standardized food intake (7).

374 Our study is not without limitations. First, we studied normal weight males who exhibited 375 high peak oxygen uptake. It is unclear if the responses observed would be evident in other 376 populations such as women, older adults, and in cohorts with overweight and obesity. It is also not known if the changes observed in response to exercise would be seen during exercise 377 378 protocols lower in time and intensity. Hence, though our results may be important for obesity 379 prevention, additional work is needed in other populations and in response to exercise 380 regimens performed more frequently amongst the general population, especially in those who 381 are overweight or obese. Second, we only examined BChE activity during the first hour of

the main trials. Although this allowed us to evaluate the transient influence of exercise,

383 further work is needed to elucidate the longer-term changes in BChE activity after exercise.

In conclusion, our study showed carriers of the *FTO* rs9939609 A-allele display lower fasting

385 BChE activity, higher post-meal AG and AG/DAG ratio, and higher energy intake compared

to TTs. However, a single bout of exercise enhances BChE activity, and corrects the

387 attenuated meal-induced suppression of AG in AAs, while the energy cost of exercise did not

388 engender an increase in energy intake in either genotype group. These findings suggest that

389 exercise could be a strategy to ameliorate the adiposity-related traits mediated by the obesity-

390 linked *FTO* rs9939609 SNP.

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399 Author contributions: JD, RLB and DJS designed the research; JD, JAK and DJS

400 conducted the research; JD, JJ and AP conducted DNA extraction; JD, DJC, JJ, WGC and

401 RLB conducted biochemical analysis; JD, JAK, AET, RLB and DJS analyzed data and

402 performed statistical analysis; JD, AET, RLB and DJS wrote the paper; JD, RLB and DJS

403 had primary responsibility for final content. All authors read and approved the final

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References

- Frayling TM, Timpson NJ, Weedon MN, Zeggini E, Freathy RM, Lindgren CM, Perry JRB, Elliott KS, Lango H, Rayner NW, et al. A common variant in the FTO gene is associated with body mass index and predisposes to childhood and adult obesity. Science (80-). 2007;316:889–94.
- Scuteri A, Sanna S, Chen WM, Uda M, Albai G, Strait J, Najjar S, Usala G, Dei M, Lai S, et al. Genome-wide association scan shows genetic variants in the FTO gene are associated with obesity-related traits. PLoS Genet. 2007;3:1200–10.
- Speliotes EK, Willer CJ, Berndt SI, Monda KL, Thorleifsson G, Jackson AU, Allen HL, Lindgren CM, Luan J, Mägi R, et al. Association analyses of 249,796 individuals reveal 18 new loci associated with body mass index. Nat Genet. 2010;42:937–48.
- Rutters F, Lemmens SGT, Born JM, Bouwman F, Nieuwenhuizen AG, Mariman E, Westerterp-Plantenga MS. Genetic associations with acute stress-related changes in eating in the absence of hunger. Patient Educ Couns. 2010;79:367–71.
- Cecil JE, Tavendale R, Watt P, Hetherington MM, Palmer CNA. An obesityassociated FTO gene variant and increased energy intake in children. N Engl J Med. 2008;359:2558–66.
- Wardle J, Llewellyn C, Sanderson S, Plomin R. The FTO gene and measured food intake in children. Int J Obes (Lond). 2009;33:42–5.
- Karra E, Daly OGO, Choudhury AI, Yousseif A, Millership S, Neary MT, Scott WR, Chandarana K, Manning S, Hess ME, et al. A link between FTO, ghrelin, and impaired brain food-cue responsivity. J Clin Invest. 2013;123:1–13.
- 8. Broom DR, Stensel DJ, Bishop NC, Burns SF, Miyashita M. Exercise-induced suppression of acylated ghrelin in humans. J Appl Physiol. 2007;102:2165–71.
- 9. Broom DR, Batterham RL, King JA, Stensel DJ. Influence of resistance and aerobic

exercise on hunger, circulating levels of acylated ghrelin, and peptide YY in healthy males. Am J Physiol Regul Integr Comp Physiol. 2009;296:R29–35.

- King JA, Wasse LK, Ewens J, Crystallis K, Emmanuel J, Batterham RL, Stensel DJ. Differential acylated ghrelin, peptide YY3-36, appetite, and food intake responses to equivalent energy deficits created by exercise and food restriction. J Clin Endocrinol Metab. 2011;96:1114–21.
- Martins C, Morgan LM, Bloom SR, Robertson MD. Effects of exercise on gut peptides, energy intake and appetite. J Endocrinol. 2007;193:251–8.
- Manning S, Batterham RL. The role of gut hormone peptide YY in energy and glucose homeostasis: Twelve years on. Annu Rev Physiol. 2014;76:585–608.
- De Vriese C, Gregoire F, Lema-Kisoka R, Waelbroeck M, Robberecht P, Delporte C. Ghrelin degradation by serum and tissue homogenates: Identification of the cleavage sites. Endocrinology. 2004;145:4997–5005.
- Chen VP, Gao Y, Geng L, Brimijoin S. Butyrylcholinesterase regulates central ghrelin signaling and has an impact on food intake and glucose homeostasis. Int J Obes. 2017;41:1413–9.
- Chen VP, Gao Y, Geng L, Brimijoin S. Butyrylcholinesterase gene transfer in obese mice prevents postdieting body weight rebound by suppressing ghrelin signaling. Proc Natl Acad Sci. 2017;114:10960–5.
- Zimmer KR, Lencina CL, Zimmer AR, Thiesen FV. Influence of physical exercise and gender on acetylcholinesterase and butyrylcholinesterase activity in human blood samples. Int J Environ Health Res. 2012;22:279–86.
- Durnin J, Womersley J. Body fat assessed from total body density and its estimation from skinfold thickness : measurements on 481 men and women aged from 16 to 72 years. Br J Nutr. 1973;32:77–97.

- Craig CL, Marshall AL, Sjöström M, Bauman AE, Booth ML, Ainsworth BE, Pratt M, Ekelund U, Yngve A, Sallis JF, et al. International physical activity questionnaire: 12-Country reliability and validity. Med Sci Sports Exerc. 2003;35:1381–95.
- 19. Stunkard AJ, Messick S. The three-factor eating questionnaire to measure dietary restraint, disinhibition and hunger. J Psychosom Res. 1985;29:71–83.
- Frecka JM, Mattes RD. Possible entrainment of ghrelin to habitual meal patterns in humans. AJP Gastrointest Liver Physiol. 2008;294:G699–707.
- Chandarana K, Drew ME, Emmanuel J, Karra E, Gelegen C, Chan P, Cron NJ, Batterham RL. Subject standardization, acclimatization, and sample processing affect gut hormone levels and appetite in humans. Gastroenterology. 2009;136:2115–26.
- Frayn KN. Calculation of substrate oxidation rates in vivo from gaseous exchange. J Appl Physiol. 1983;55:628–34.
- 23. Flint A, Raben A, Blundell JE, Astrup A. Reproducibility, power and validity of visual analogue scales in assessment of appetite sensations in single test meal studies. Int J Obes Relat Metab Dsorders J Int Assoc Study Obes. 2000;24:38–48.
- Batterham RL, Ffytche DH, Rosenthal JM, Zelaya FO, Barker GJ, Withers DJ,
 Williams SCR. PYY modulation of cortical and hypothalamic brain areas predicts feeding behaviour in humans. Nature. 2007;450:106–9.
- 25. Ellman GL, Courtney KD, Andres V, Featherstone RM. A new and rapid colorimetric determination of acetylcholinesterase activity. Biochem Pharmacol. 1961;7:88–95.
- 26. Wasse LK, Sunderland C, King JA, Miyashita M, Stensel DJ. The influence of vigorous running and cycling exercise on hunger perceptions and plasma acylated ghrelin concentrations in lean young men. Appl Physiol Nutr Metab. 2013;38:1–6.
- Cohen J. Statistical power analysis for the behavioral sciences. Statistical Power Analysis for the Behavioral Sciences. 1988. p. 567.

- 28. Benedict C, Axelsson T, Söderberg S, Larsson A, Ingelsson E, Lind L, Schiöth HB. Brief communication : The fat mass and obesity-associated gene (FTO) is linked to higher plasma levels of the hunger hormone ghrelin and lower serum levels of the satiety hormone leptin in older adults. Diabetes. 2014;63:3955–9.
- 29. Delhanty PJD, Neggers SJ, van der Lely AJ. Ghrelin: The differences between acyland des-acyl ghrelin. European Journal of Endocrinology. 2012. p. 601–8.
- Kuppens RJ, Diène G, Bakker NE, Molinas C, Faye S, Nicolino M, Bernoux D, Delhanty PJD, van der Lely AJ, Allas S, et al. Elevated ratio of acylated to unacylated ghrelin in children and young adults with Prader–Willi syndrome. Endocrine. 2015;50:633–42.
- 31. Kilpeläinen TO, Qi L, Brage S, Sharp SJ, Sonestedt E, Demerath E, Ahmad T, Mora S, Kaakinen M, Sandholt CH, et al. Physical activity attenuates the influence of FTO variants on obesity risk: A meta-analysis of 218,166 adults and 19,268 children. PLoS Med. 2011;8:2–14.
- Mitchell JA, Church TS, Rankinen T, Earnest CP, Sui X, Blair SN. FTO genotype and the weight loss benefits of moderate intensity exercise. Obesity (Silver Spring). 2010;18:641–3.
- Xiang L, Wu H, Pan A, Patel B, Xiang G, Qi L, Kaplan RC, Hu F, Wylie-Rosett J, Qi
 Q. FTO genotype and weight loss in diet and lifestyle interventions: a systematic review and meta-analysis. Am J Clnical Nutr. 2016;103:1162–7.
- 34. Walsh NP, Gleeson M, Shephard RJ, Gleeson M, Woods JA, Bishop NC, Fleshner M, Green C, Pedersen BK, Hoffman-Goetz L, et al. Position statement part one: Immune function and exercise. Exerc Immunol Rev. 2011;17:6–63.
- Benyamin B, Middelberg RP, Lind PA, Valle AM, Gordon S, Nyholt DR, Medland
 SE, Henders AK, Heath AC, Madden PAF, et al. GWAS of butyrylcholinesterase

activity identifies four novel loci, independent effects within BCHE and secondary associations with metabolic risk factors. Hum Mol Genet. 2011;20:4504–14.

- Karasova JZ, Maderycova Z, Tumova M, Jun D, Rehacek V, Kuca K, Misik J. Activity of cholinesterases in a young and healthy middle-European population: Relevance for toxicology, pharmacology and clinical praxis. Toxicol Lett. 2017;277:24–31.
- 37. Sato KK, Hayashi T, Maeda I, Koh H, Harita N, Uehara S, Onishi Y, Oue K,
 Nakamura Y, Endo G, et al. Serum butyrylcholinesterase and the risk of future type 2
 diabetes: The Kansai Healthcare Study. Clin Endocrinol (Oxf). 2014;80:362–7.
- 38. Jia G, Fu Y, Zhao X, Dai Q, Zheng G, Yang Y, Yi C, Lindahl T, Pan T, Yang YG, et al. N6-Methyladenosine in nuclear RNA is a major substrate of the obesity-associated FTO. Nat Chem Biol. 2011;7:885–7.
- 39. Dougkas A, Yaqoob P, Givens DI, Reynolds CK, Minihane AM. The impact of obesity-related SNP on appetite and energy intake. Br J Nutr. 2013;110:1151–6.
- 40. Tanaka T, Ngwa JS, van Rooij FJ a, Zillikens MC, Wojczynski MK, Frazier-Wood AC, Houston DK, Kanoni S, Lemaitre RN, Luan J, et al. Genome-wide meta-analysis of observational studies shows common genetic variants associated with macronutrient intake. Am J Clnical Nutr. 2013;97:1395–402.
- Speakman JR. The "Fat Mass and Obesity Related" (FTO) gene: Mechanisms of Impact on Obesity and Energy Balance. Curr Obes Rep. 2015;4:73–91.
- 42. Le Roux CW, Batterham RL, Aylwin SJB, Patterson M, Borg CM, Wynne KJ, Kent A, Vincent RP, Gardiner J, Ghatei MA, et al. Attenuated peptide YY release in obese subjects is associated with reduced satiety. Endocrinology. 2006;147:3–8.
- 43. Stanley S, Wynne K, Bloom S. Gastrointestinal satiety signals III. Glucagon-like peptide 1, oxyntomodulin, peptide YY, and pancreatic polypeptide. Am J Physiol

Gastrointest Liver Physiol. 2004;286:G693-7.

	AA (n = 12)	TT (n = 12)	Main effect genotype TT vs AA Mean difference (95% CI ¹)
Age (years)	20.9 ± 3.5	21.3 ± 3.6	-0.4 (-3.4, 2.6)
Height (cm)	181.6 ± 5.8	177.5 ± 6.5	4.1 (-1.2, 9.3)
Body mass (kg)	77.6 ±11.3	73.8 ± 7.0	3.9 (-4.1, 11.8)
BMI (kg/m ²)	23.5 ± 2.7	23.5 ± 2.3	0.1 (-2.1, 2.1)
Body fat (%)	15.6 ± 5.1	13.9 ± 4.7	1.7 (-2.4, 5.9)
Lean body mass (kg)	65.2 ± 7.4	63.3 ± 4.2	1.9 (-3.2, 7.0)
Waist circumference (cm)	80.3 ± 6.1	78.1 ± 4.1	2.1 (-2.2, 6.6)
Three-Factor Eating Questionnaire			
Dietary restraint	7.7 ± 4.5	7.6 ± 3.9	0.1 (-3.5, 3.6)
Dietary disinhibition	6.3 ± 2.3	6.6 ± 1.6	-0.3 (-1.9, 1.4)
Hunger	6.5 ± 2.1	6.9 ± 1.7	-0.4 (-2.0, 1.2)
Total physical activity (metabolic equivalent minutes/week)	4368 ± 1968	4790 ± 2728	-423 (-2436, 1591)
Peak oxygen uptake (mL/kg/min)	55.8 ± 5.8	56.6 ± 4.9	-0.8 (-5.4, 3.7)

Table 1. Characteristics of the AA and TT participants.

Values are mean \pm SD. Data were analyzed using linear mixed models with genotype (AA or TT) included as a fixed factor.

¹95% confidence interval of the mean absolute difference between the genotype groups. No differences were identified between genotype groups ($P \ge 0.121$).

	AA (n = 12)		TT (n	= 12)	Main effect trial	Main effect genotype
	Control	Exercise	Control	Exercise	Mean difference (95% CI ¹)	TT vs AA Mean difference (95% CI ²)
Acyl-ghrelin (pmol/L)	22.4 ± 1.4	22.5 ± 1.3	20.9 ± 1.5	21.1 ± 1.5	0.1 (-0.4, 0.6)	1.4 (-2.7, 5.6)
Des-acyl-ghrelin (pmol/L)	135.0 ± 9.3	134.1 ± 8.7	156.3 ± 10.6	155.4 ± 10.0	-0.9 (-6.1, 4.3)	-21.3 (-49.1, 6.5)
Acyl-/des-acyl-ghrelin ratio	0.167 ± 0.005	0.169 ± 0.006	0.134 ± 0.004	0.135 ± 0.003	0.002 (-0.002, 0.006)	$0.034 (0.021, 0.047)^3$
Total GLP-1 (pmol/L)	26.2 ± 2.2	25.3 ± 2.2	32.3 ± 3.3	31.7 ± 3.5	-0.8 (-2.1, 0.6)	-6.2 (-14.6, 2.1)
Total PYY (pg/mL)	156.2 ± 12.2	163.1 ± 12.7	187.4 ± 20.8	185.4 ± 17.8	2.5 (-11.3, 16.3)	-26.8 (-72.5, 18.9)
Leptin (pg/mL)	1216.5 ± 167.0	1358.1 ±182.8	1343.3 ± 261.3	1267.0 ± 205.0	32.7 (-133.0, 198.3)	-17.9 (-657.9, 622.2)
Butyrylcholinesterase activity (KU/L)	1.481 ± 0.060	1.404 ± 0.062	1.613 ± 0.060	1.635 ± 0.062	-0.027 (-0.129, 0.074)	-0.181 (-0.360, -0.003) ³

Table 2. Fasting appetite-related hormone concentrations and butyrylcholinesterase activity at baseline for AAs and TTs in the control and exercise trials.

Values are mean \pm SEM. Data were analyzed using linear mixed models with trial (exercise or control) and genotype (AA or TT) included as fixed factors.

¹95% confidence interval of the mean absolute difference between the experimental trials.

²95% confidence interval of the mean absolute difference between the genotype groups.

³ Significant main effect of genotype (P < 0.05).

Linear mixed models revealed no main effects of trial ($P \ge 0.127$) and no genotype-by-trial interactions ($P \ge 0.319$).

GLP-1, glucagon-like peptide-1; PYY, peptide YY.

	AA (n	i = 12)	TT (n = 12)		Main effect trial	Main effect genotype	
	Control	Exercise	Control	Exercise	Control vs exercise Mean difference (95% CI ¹)	TT vs AA Mean difference (95% CI ²)	
Δ AG (pmol/L·h)							
Intervention period	3.8 ± 0.7	-17.4 ± 1.6	5.2 ± 0.9	-15.0 ± 1.5	-20.7 (-23.4, -18.0) ³	-1.9 (-4.3, 0.5)	
Post-test meal	-5.6 ± 1.9	-29.6 ± 3.5	-15.0 ± 2.4	-29.3 ± 2.7	-19.2 (-23.3, -15.1) ^{3,4}	4.5 (-2.1, 11.2) ⁴	
Afternoon	-38.9 ± 6.9	-52.1 ± 8.6	-40.2 ± 7.6	-53.9 ± 7.1	-13.4 (-19.8, -7.1) ³	1.6 (-19.7, 22.9)	
Post-buffet meal	-8.9 ± 2.4	-11.6 ± 2.7	-7.5 ± 2.7	-10.1 ± 1.8	-2.6 (-5.2, 0.1)	-1.4 (-8.1, 5.2)	
Δ DAG (pmol/L·h)							
Intervention period	18.0 ± 3.5	-28.2 ± 10.8	27.0 ± 7.8	-55.4 ± 11.2	-64.3 (-81.7, -46.9) ^{3,4}	9.1 (-10.3, 28.5) ⁴	
Post-test meal	-62.4 ± 13.3	-101.4 ± 23.4	-66.6 ± 16.6	-167.4 ± 18.4	-67.9 (-96.2, -39.7) ^{3,4}	33.2 (-13.1, 79.4) ⁴	
Afternoon	-255.6 ± 49.1	-271.4 ± 48.5	-317.4 ± 54.5	-407.6 ± 61.2	-53.0 (-99.6, -6.4) ³	99.0 (-51.1, 249.1)	
Post-buffet meal	-73.2 ± 18.6	-46.3 ± 13.2	-76.7 ± 21.6	-74.7 ± 15.9	12.3 (-5.8, 30.5)	11. 8 (-37.1, 60.6)	
Δ AG/DAG ratio							
Intervention period	0.01 ± 0.01	-0.11 ± 0.01	0.01 ± 0.01	-0.06 ± 0.01	-0.09 (-0.11, -0.08) ^{3,4}	-0.03 (-0.05, -0.02) ^{4,5}	
Post-test meal	0.03 ± 0.02	-0.12 ± 0.02	-0.05 ± 0.02	-0.06 ± 0.01	-0.09 (-0.12 , -0.05) ^{3,4}	$0.01 (-0.02, 0.04)^4$	
Afternoon	0.04 ± 0.04	-0.09 ± 0.04	0.02 ± 0.04	0.02 ± 0.04	-0.7 (-0.14, 0.00)	-0.04 (-0.13, 0.05)	
Post-buffet meal	0.04 ± 0.02	-0.04 ± 0.02	0.02 ± 0.01	0.00 ± 0.01	$-0.05 (-0.09, -0.02)^3$	-0.01 (-0.04, 0.02)	

Table 3. Time-averaged total area under the curve for delta acyl-ghrelin, des-acyl-ghrelin and the acyl-/des-acyl-ghrelin ratio for AAs and TTs in the exercise and control trials.

Values are mean \pm SEM. Intervention period covers 0.0-1.0 h; post-test meal covers 1.5-3.5 h; afternoon period covers 3.5-6.5 h; post-buffet meal covers 6.5-7.5 h. Data were analyzed using linear mixed models with trial (exercise or control) and genotype (AA or TT) included as fixed factors.

¹95% confidence interval of the mean absolute difference between the experimental trials.

² 95% confidence interval of the mean absolute difference between the genotype groups.

³ Significant main effect of trial (P < 0.05). ⁴ Significant genotype-by-trial interaction (P < 0.05). ⁵ Significant main effect of genotype (P < 0.05).

AG, acyl-ghrelin; DAG, des-acyl-ghrelin.

	AA (n = 12)		TT (n = 12)		Main effect trial	Main effect genotype	
	Control	Exercise	Control	Exercise	Control vs exercise Mean difference (95% CI ¹)	TT vs AA Mean difference (95% CI ²)	
Δ Total GLP-1 (pmol/L·h)							
Intervention period	-3.8 ± 0.9	15.0 ± 1.8	-6.5 ± 1.2	10.7 ± 2.4	$18.0(14.7, 21.3)^3$	3.5 (-0.2, 7.2)	
Post-test meal	34.2 ± 7.9	107.0 ± 12.1	21.4 ± 7.0	112.3 ± 8.0	81.6 (64.9, 98.3) ³	3.9 (-16.8, 24.8)	
Afternoon	97.0 ± 22.4	142.8 ± 15.2	80.0 ± 17.4	144.6 ± 15.4	55.2 (27.0, 83.4) ³	7.6 (-36.5, 51.7)	
Post-buffet meal	33.0 ± 7.5	44.6 ± 5.2	15.7 ± 4.8	25.0 ± 5.6	$10.4 (0.3, 20.5)^3$	$18.6 (4.7, 32.4)^4$	
Δ Total PYY (pg/mL·h)							
Intervention period	-14.7 ± 8.3	51.5 ± 13.3	-18.3 ± 3.8	53.7 ± 13.3	$69.1 (48.2, 90.0)^3$	0.7 (-21.8, 23.1)	
Post-test meal	105.7 ± 22.9	215.2 ± 34.6	61.1 ± 24.7	207.3 ± 30.7	$128.4 (74.3, 182.6)^3$	25.7 (-40.0, 91.3)	
Afternoon	507.5 ± 82.9	536.4 ± 85.8	394.0 ± 85.4	458.7 ± 67.8	46.8 (-76.5, 170.0)	95.6 (-106.9, 298.1)	
Post-buffet meal	198.4 ± 23.5	166.6 ± 21.7	108.9 ± 22.0	131.8 ± 23.7	-4.0 (-43.2, 35.3)	61.6 (7.1, 116.2) ⁴	

Table 4. Time-averaged total area under the curve for delta concentrations of total glucagon-like peptide 1 and total peptide YY for AAs and TTs in the exercise and control trials.

Values are mean \pm SEM. Intervention period covers 0.0-1.0 h; post-test meal covers 1.5-3.5 h; afternoon period covers 3.5-6.5 h; post-buffet meal covers 6.5-7.5 h. Data were analyzed using linear mixed models with trial (exercise or control) and genotype (AA or TT) included as fixed factors.

¹95% confidence interval of the mean absolute difference between the experimental trials.

²95% confidence interval of the mean absolute difference between the genotype groups.

³ Significant main effect of trial (P < 0.05).

⁴ Significant main effect of genotype (P < 0.05).

Linear mixed models revealed no genotype-by-trial interactions ($P \ge 0.169$).

GLP-1, glucagon-like peptide-1, PYY, peptide YY.

	AA (n = 12)		TT (n =12)		Main effect trial	Main effect genotype
	Control	Exercise	Control	Exercise	Control vs exercise Mean difference (95% CI ¹)	$\begin{array}{c} \text{TT vs AA} \\ \text{Mean difference} \\ (95\% \text{ CI}^2) \end{array}$
Hunger (mm·h)						
Intervention	68 ± 4	39 ± 5	80 ± 3	53 ± 6	-27 (-37,-18) ³	-10 (-20, 1)
Post-test meal	83 ± 8	87 ± 6	60 ± 6	60 ± 5	2 (-9, 13)	25 (10, 40) ⁴
Afternoon	172 ± 14	192 ± 13	138 ± 10	144 ± 14	13 (-4, 30)	$41(7,74)^4$
Post-buffet meal	35 ± 4	44 ± 4	32 ± 3	31 ± 3	-1.6 (-2, 9)	8.0 (-1, 17)
Fullness (mm·h)						
Intervention	21 ± 4	39 ± 5	13 ± 3	25 ± 5	15 (9, 21) ³	11 (0, 23)
Post-test meal	112 ± 7	115 ± 7	132 ± 6	137 ± 5	4 (-6, 15)	-20 (-37, -3) ⁴
Afternoon	108 ± 13	102 ± 13	142 ± 12	141 ± 12	-4 (-27, 19)	-37 (-66, -8) ⁴
Post-buffet meal	99 ± 4	101 ± 3	112 ± 3	110 ± 3	0 (-4, 3)	-11 (-20, -2) ⁴
Prospective food consumption (mm·h)						
Intervention	77 ± 4	51 ± 5	80 ± 4	58 ± 6	-24 (-32, -16) ³	-6 (-17, 6)
Post-test meal	99 ± 8	102 ± 7	77 ± 8	71 ± 9	-2 (-11, 8)	26 (5, 48) ⁴
Afternoon	186 ± 14	205 ± 11	163 ± 12	157 ± 16	6 (-10, 23)	$(1, 71)^4$
Post-buffet meal	46 ± 5	52 ± 5	39 ± 3	43 ± 6	5 (-1, 11)	7 (-6, 20)
Hedonic wanting of food (mm·h)						
Intervention	78 ± 4	49 ± 6	83 ± 4	57 ± 6	-28 (-38, -19) ³	-7 (-19, 6)
Post-test meal	107 ± 10	107 ± 6	80 ± 9	78 ± 10	-2 (-12, 8)	$28(4,52)^4$
Afternoon	201 ± 12	$219\pm\!9$	161 ± 13	158 ± 17	8 (-11, 26)	51 (17, 84) ⁴
Post-buffet meal	55 ± 7	61 ± 5	52 ± 6	51 ± 7	2 (-5, 10)	7 (-10, 23)

Table 5. Time-averaged total area under the curve for appetite perceptions for AAs and TTs in the control and exercise trials.

Values are mean \pm SEM. Intervention period covers 0.0-1.0 h; post-test meal covers 1.5-3.5 h; afternoon period covers 3.5-6.5 h; post-buffet meal covers 6.5-8.0 h. Data were analyzed using linear mixed models with trial (exercise or control) and genotype (AA or TT) included as fixed factors.

¹95% confidence interval of the mean absolute difference between the experimental trials.

²95% confidence interval of the mean absolute difference between the genotype groups.

³ Significant main effect of trial (P < 0.05).

⁴ Significant main effect of genotype (P < 0.05).

Linear mixed models revealed no genotype-by-trial interactions ($P \ge 0.061$).

	AA (n = 12)		TT (n	= 12)	Main effect trial	Main effect	
	Control	Exercise	Control	Exercise	Control vs exercise Mean difference (95% CI ¹)	genotype TT vs AA Mean difference (95% CI ²)	
Absolute energy intake (kJ)	5229 ± 576	5554 ± 627	3788 ± 463	$\begin{array}{r} 3897 \pm \\ 490 \end{array}$	217 (-191, 625)	1549 (10, 3088) ³	
Relative energy intake (kJ)	5139 ± 571	1888 ± 642	3710 ± 429	532 ± 467	-3214 (-3674, - 2755) ⁴	1393 (-186, 2973)	
Carbohydrate (g)	$\frac{160 \pm 18}{18}$	162 ± 17	117 ± 16	119 ± 17	3 (-12,18)	43 (-5, 90)	
Protein (g)	48 ± 4	52 ± 5	36 ± 4	37 ± 5	3 (-1, 7)	$14(1, 26)^3$	
Fat (g)	47 ±7	52 ± 8	33 ± 4	34 ± 4	3 (0, 7)	16 (-1, 34)	

Table 6. Energy and macronutrient intakes at the buffet meal for AAs and TTs in the exercise and control trials.

Values are mean \pm SEM. Relative energy intake is energy intake at the buffet meal minus the gross energy expenditure of the intervention period (0.0-1.0 h). Data were analyzed using linear mixed models with trial (exercise or control) and genotype (AA or TT) included as fixed factors.

¹95% confidence interval of the mean absolute difference between the experimental trials.

²95% confidence interval of the mean absolute difference between the genotype groups.

³ Significant main effect of genotype (P < 0.05).

⁴ Significant main effect of trial (P < 0.05).

Linear mixed models revealed no genotype-by-trial interactions ($P \ge 0.207$).

Figure legends

Figure 1. Schematic representation of the main trials.

Figure 2. Δ AG concentrations (A), DAG concentrations (B) and AG/DAG ratio (C) in AAs (n = 12) and TTs (n = 12) during the control and exercise trials. *Dotted rectangle* indicates exercise, *horizontally dashed rectangle* indicates standardized test meal, *vertically dashed rectangle* indicates buffet meal. Data are represented as mean ± SEM. Data were analyzed using linear mixed models with trial (exercise or control), genotype (AA or TT) and time included as fixed factors. Δ AG: main effect trial P < 0.001, main effect time P < 0.001, genotype-by-time interaction P = 0.007; Δ DAG: main effect trial P < 0.001, main effect trial P < 0.001, main effect trial P < 0.001, genotype-by-trial interaction P < 0.001; Δ AG/DAG ratio: main effect trial P < 0.001, genotype-by-time interaction P < 0.001, genotype-by-trial interaction P < 0.001; Δ AG/DAG ratio: main effect trial P < 0.001, genotype-by-time interaction P < 0.001, genotype-by-trial interaction P < 0.001; Δ AG/DAG ratio: main effect trial P < 0.001, genotype-by-trial interaction P < 0.001, genotype-by-trial interaction P < 0.001; Δ AG/DAG ratio: main effect trial P < 0.001, genotype-by-trial-by-time interaction P = 0.004. Linear mixed models for Δ AG, Δ DAG and Δ AG/DAG ratio revealed no main effect of genotype (all P ≥ 0.192) or other interactive effects (P ≥ 0.083). AG, acyl-ghrelin; DAG, des-acyl-ghrelin.

Figure 3. Δ Total GLP-1 (A) and total PYY (B) concentrations in AAs (n = 12) and TTs (n = 12) during the control and exercise trials. *Dotted rectangle* indicates exercise, *horizontally dashed rectangle* indicates standardized test meal, *vertically dashed rectangle* indicates buffet meal. Data are represented as mean ± SEM. Data were analyzed using linear mixed models with trial (exercise or control), genotype (AA or TT) and time included as fixed factors. Δ total GLP-1: main effect trial P < 0.001, main effect time P < 0.001, genotype-by-time interaction P = 0.002; Δ total GLP-1 and Δ total PYY revealed no main effect of genotype (all P ≥ 0.278) or other interactive effects (P ≥ 0.089). GLP-1, glucagon-like peptide-1, PYY, peptide YY.

Figure 4. Δ Plasma BChE activity in AAs (n = 12) and TTs (n =12) during the control and exercise trials at 0.5 and 1.0 h. *Dotted rectangle* indicates exercise. * *P* = 0.004 for main effect of trial. Data are represented as mean ± SEM. Data were analyzed using linear mixed models with trial (exercise or control), genotype (AA or TT) and time included as fixed factors. Δ BChE activity: main effect trial P = 0.004, main effect time P < 0.001. Linear mixed models for Δ BChE activity revealed no main effect of genotype (P = 0.681) or interactive effects (P ≥ 0.094). BChE, butyrylcholinesterase.

Figure 5. Hunger (A), fullness (B), prospective food consumption (C) and hedonic wanting of food (D) in AAs (n = 12) and TTs (n = 12) during the control and exercise trials. *Dotted rectangle* indicates exercise, *horizontally dashed rectangle* indicates standardized test meal, *vertically dashed rectangle* indicates buffet meal. Data are represented as mean \pm SEM. Data were analyzed using linear mixed models with trial (exercise or control), genotype (AA or TT) and time included as fixed factors. All appetite perceptions: main effect trial P = 0.002, main effect time P < 0.001, genotype-by-time interaction P < 0.001. Linear mixed models for each appetite perception revealed no main effect of genotype (P \ge 0.072) or other interactive effects (P \ge 0.094).