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1 Homeostatic and non-homeostatic appetite control along the spectrum of physical activity
2 levels: an updated perspective

3

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18

19 **Abstract**

20 The current obesogenic environment promotes physical inactivity and food consumption in
21 excess of energy requirements, two important modifiable risk factors influencing energy
22 balance. Habitual physical activity has been shown to impact not only energy expenditure, but
23 also energy intake through mechanisms of appetite control. This review summarizes recent
24 theory and evidence underpinning the role of physical activity in the homeostatic and non-
25 homeostatic mechanisms controlling appetite. Energy intake along the spectrum of physical
26 activity levels (inactive to highly active) appears to be J-shaped, with low levels of physical
27 activity leading to dysregulated appetite and a mismatch between energy intake and
28 expenditure. At higher levels, habitual physical activity influences homeostatic appetite control
29 in a dual-process action by increasing the drive to eat through greater energy expenditure, but
30 also by enhancing post-meal satiety, allowing energy intake to better match energy
31 expenditure in response to hunger and satiety signals. There is clear presumptive evidence
32 that physical activity energy expenditure can act as a drive (determinant) of energy intake.
33 The influence of physical activity level on non-homeostatic appetite control is less clear, but
34 low levels of physical activity may amplify hedonic states and behavioural traits favouring
35 overconsumption indirectly through increased body fat. More evidence is required to
36 understand the interaction between physical activity, appetite control and diet composition on
37 passive overconsumption and energy balance. Furthermore, potential moderators of appetite
38 control along the spectrum of physical activity, such as body composition, sex, and type,
39 intensity and timing of physical activity, remain to be fully understood.

40 **Keywords:** appetite control, physical activity, energy intake, food hedonics, energy balance

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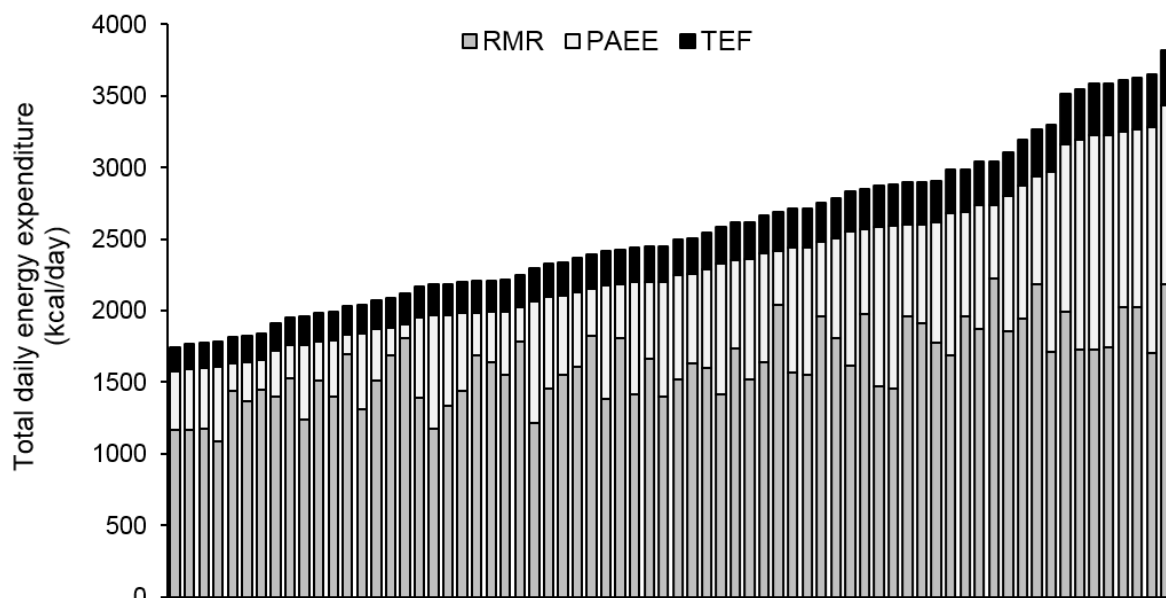
43 1. Defining energy balance and appetite control

44 Energy balance and resulting effects on body weight are the product of a complex
45 relationship between energy intake (EI) and energy expenditure (EE). EI is modulated by the
46 appetite control system through food consumption and eating behaviour. EI is largely
47 influenced by a combination of internal biological factors such as resting metabolic rate (RMR)
48 [1] and external nutritional factors such as energy density (kcal/g) of the food consumed [2],
49 with intake being greater at higher RMR and energy density [3]. Of the macronutrients, dietary
50 fat has the strongest influence on energy density (9 kcal/g) compared to carbohydrate and
51 protein (4 kcal/g). Because of its higher energy density, dietary fat has been shown to be less
52 satiating per unit of energy than the other macronutrients, resulting in greater energy intake
53 when consumed ad libitum, which has led to the term “passive overconsumption” [4].

54 On the other side of the energy balance equation is EE, which is composed of RMR,
55 physical activity energy expenditure (PAEE) and thermic effect of food (TEF), with typical
56 ranges of 60-70%, 15-35% and 5-15% of total daily energy expenditure (TDEE), respectively
57 [5]. PA encompasses structured exercise in addition to occupational, household,
58 transportation and other activities of daily living, termed non-exercise activity thermogenesis
59 (NEAT) [6]. In recent years, wearable technologies that estimate free-living total daily energy
60 expenditure (TDEE), and minutes spent sedentary and in different intensities of PA (i.e. light,
61 moderate and vigorous) have allowed for the objective assessment of habitual PA level, which
62 was a major limitation of past research investigating the impact of habitual PA on appetite
63 control [7]. These methods have also been improved to properly assess sedentary behaviour
64 using inclinometers, which is specifically defined as “any waking behaviour characterized by
65 an energy expenditure ≤ 1.5 metabolic equivalents (METs), while in a sitting, reclining or lying
66 posture” [8]. This is different to physical inactivity which is “an insufficient physical activity level
67 to meet present physical activity recommendations” [8]. Distinguishing between sedentary
68 behaviour and physical inactivity is of importance to better understand their influence on

69 energy balance and appetite control. This review focuses on habitual PA levels (i.e. physical
70 activity/inactivity) rather than sedentary behaviour.

71 The components of TDEE are illustrated in Figure 1, which shows the contribution of
72 RMR and PAEE towards TDEE can vary widely between individuals depending on levels of
73 PA. Indeed, within this group of 70 individuals (61% female, age: 29.5 ± 9.1 years, body mass
74 index: 22.7 ± 2.3 kg/m²; means \pm SD) [9, 10], PAEE varied in absolute values between 102
75 and 1579 kcal, and from 5% to 43% as a percentage of TDEE. Our argument is that this
76 degree of variability must have considerable implications for appetite control, energy balance
77 and body weight. Here, PAEE was calculated by subtracting measured RMR (indirect
78 calorimetry) and estimated TEF (~10% of TDEE) from measured TDEE obtained from a multi-
79 sensor accelerometry device worn for five to seven days. While TEF was estimated at 10%
80 as a generalisation for healthy individuals consuming a mixed diet, it is important to consider
81 that TEF varies according to the energy content and macronutrient composition of food [11]
82 and is perhaps reduced with obesity [12], but further well-designed studies are required to
83 confirm the latter [11].



84
85 **Figure 1** Individual profile (n=70) of the components of TDEE including resting metabolic rate
86 (RMR), physical activity energy expenditure (PAEE), thermic effect of food (TEF). TDEE is

87 composed primarily of RMR, followed by PAEE, which varies widely between individuals. From
88 Beaulieu et al. [unpublished results].

89

90 In order to understand how PA influences appetite, it is necessary to have a
91 conceptualisation of the functional operations of the appetite system. Our position is that
92 appetite is controlled by several processes that form a psychobiological system that signals
93 hunger (drives feeding), satiation (terminates feeding) and satiety (post-meal suppression of
94 hunger), which in turn determine food (energy) intake [13]. These processes are influenced
95 by episodic and tonic signals. Episodic signals occur on a meal-to-meal basis and diurnal
96 variations in these signals reflect the size, pattern and frequency of meals and eating
97 episodes. Episodic signals can be excitatory or inhibitory, and are related to meal initiation,
98 termination and satiety. Tonic signals stem from body tissues and cellular metabolism, and
99 convey information relating to energy availability and energy needs to the central nervous
100 system [14]. These homeostatic mechanisms interact with non-homeostatic processes, such
101 as food hedonics and behavioural traits, in the overall expression of appetite [15]. The complex
102 relationships between homeostatic and non-homeostatic inputs, coupled with the current
103 obesogenic food environment, can make individuals vulnerable to overconsumption and
104 weight gain. However, there has been increasing interest and research on the influence and
105 benefits of PA on appetite control as it plays an integral (and readily modifiable) part in energy
106 balance and body weight [16-20]. The aim of this review is to summarize recent theory and
107 evidence underpinning the role of habitual PA in the mechanisms controlling appetite and its
108 impact on energy balance. In principle, PA could influence tonic and episodic appetite control
109 by adjusting the biological components of fat-free mass, fat mass, RMR and postprandial
110 peptides, for example. In addition, our position is that PA will influence appetite control via
111 both direct and indirect mechanisms; the direct effect of PAEE will drive appetite in a similar
112 manner to RMR, i.e. by increasing TDEE and associated energy demand.

113

114 **1.1 A note on physical activity, energy expenditure and body composition**

115 Common beliefs regarding TDEE assume that it increases linearly with PA in a dose-
116 dependent manner, whereby greater PA levels lead to greater TDEE [21]. Recently, whether
117 TDEE increases in proportion to PA level has been debated, and a constrained EE model has
118 been proposed by Pontzer et al. [22]. Using doubly-labelled water to measure TDEE and
119 accelerometers to measure PA in several populations of adults, these authors demonstrated
120 that at lower levels of PA, TDEE increases linearly with PA, but at a certain threshold of much
121 higher PA, there was a plateau in TDEE. Thus, it was suggested that TDEE is regulated such
122 that compensatory reductions in other metabolic processes or components of TDEE occur
123 with increasing PA to maintain TDEE within a certain narrow range [22]. This could be related
124 to a reduction in NEAT or enhanced metabolic economy/efficiency of PA at higher levels of
125 PA. However, further evidence is required to validate this model, and more convincing study
126 designs are needed to elucidate the mechanisms that could regulate TDEE and the specific
127 components of TDEE affected at very high levels of PA.

128 It is also important to emphasise that within the general population, which is highly
129 inactive and at the low end of the PA spectrum [23], an increase in PA will likely lead to an
130 increase in TDEE. Therefore, PA should remain a key component in weight management and
131 health promotion for everyone.¹ Indeed, data from our group have shown significant inverse
132 associations between objectively-measured PA and adiposity in individuals ranging in body
133 mass index [24] as well as a significant body fat loss in response to a 12-week exercise
134 intervention in inactive men and women with overweight and obesity [25], in line with
135 systematic reviews [26, 27].

136

¹ It should be kept in mind that some studies have shown unchanged TDEE despite increased PAEE in certain inactive populations; see Melanson [21] for review.

137 **2. Homeostatic mechanisms of appetite control**

138 The homeostatic control of appetite can be conceptualised as a matrix of events and
139 interactions occurring in the three levels of the psychobiological system. These include
140 psychological and behavioural events, peripheral physiological and metabolic events, and
141 neurotransmitter and metabolic interactions in the brain [13]. It has been argued that the
142 desynchronisation of these three levels is associated with a disruption of appetite, for example
143 with eating disorders (and perhaps with physical inactivity and obesity). The system
144 incorporates the events and behaviours that lead to, arise during, terminate and occur after
145 food consumption, which have been termed the Satiety Cascade [13]. During and shortly after
146 food intake and gastric emptying, the secretion of the orexigenic (appetite stimulating) peptide
147 ghrelin is suppressed and a variety of anorectic (appetite inhibiting) peptides, such as
148 cholecystokinin (CCK), glucagon-like peptide 1 (GLP-1), and peptide YY (PYY), among
149 others, are released from the gut to promote meal termination (satiety) and the post-meal
150 suppression of hunger (satiety), which in turn, coordinate meal size and frequency [13, 28,
151 29]. In addition to its episodic role, ghrelin may also be involved in tonic appetite control as a
152 compensatory hormone to restore body weight status; with obesity, ghrelin concentrations
153 appear to be lower, whereas with weight loss, they increase [30]. Other tonic signals such as
154 leptin and insulin have been hypothesised to act directly in the central nervous system to
155 reduce appetite and energy intake [31]. However, with body fat accumulation there appears
156 to be a resistance to the direct negative feedback action of leptin and insulin in the
157 hypothalamus [14, 31]. Moreover, it is believed that an interaction between episodic and tonic
158 peptides exists, with a reduced sensitivity to leptin or insulin with increased body fat leading
159 to blunted signalling of satiety peptides such as CCK and GLP-1 [14, 32, 33]. Indeed, the
160 postprandial response of insulin has been associated with satiety in lean individuals, but not
161 in individuals with obesity [32, 34-37].

162

163 **3. Non-homeostatic factors involved in appetite control and eating behaviour**

164 In addition to homeostatic mechanisms, non-homeostatic factors involved in appetite
165 control include food hedonics and eating behaviour traits [38, 39]. Hedonic thoughts about
166 food and the sensory appreciation of certain food attributes like salt, sugar and fat determine
167 food preference and choice, and thereby contribute to meal size and frequency [40]. Food
168 hedonics reflect the separate processes of 'liking' and 'wanting' [41]. Liking can be defined as
169 the degree of sensory pleasure obtained from foods, whereas wanting is the motivation or
170 attraction towards certain foods [42]. While both processes are involved in the motivation to
171 eat, they operate as distinct entities where an increase in wanting may not necessarily predict
172 an increase in liking and vice versa [43]. Wanting may be more important for overconsumption
173 and maintenance of obesity than liking, which tends to remain stable within an individual and
174 does not appear to be influenced by obesity [39, 40, 44]. Eating behaviour traits such as dietary
175 restraint (i.e. concern over weight gain and the attempt to reduce food intake), disinhibition
176 (i.e. tendency of an individual to overeat and to eat opportunistically in the obesogenic
177 environment) [45], binge eating (i.e. excessive consumption of food in a discrete period of time
178 often accompanied by feelings of guilt and loss of control over eating) [46] and control over
179 food cravings (i.e. frequency, intensity and type of food cravings) [47] are also considered as
180 risk factors for overconsumption and weight gain [48].

181 In today's obesogenic environment, the availability of highly palatable and often
182 energy-dense foods raises the importance of hedonic influences on the control of food intake
183 that occur independently from and/or in opposition to the energy need or weight status of an
184 individual [49]. Indeed, there is growing evidence to support the considerable functional
185 overlap between the homeostatic and hedonic mechanisms of appetite control [15, 50], which
186 could be linked by GLP-1 [51], ghrelin [52, 53], insulin and/or leptin [14]. Consequently,
187 hedonic signals occurring when palatable and energy-dense foods are ingested can disrupt
188 or override homeostatic satiety signals and lead to overconsumption [52]. Indeed, it has been
189 suggested that chronic high-fat intake attenuates the satiating properties of CCK through
190 reduced sensitivity of the vagal receptors [54, 55]. Moreover, it appears the PYY response to

191 dietary fat is attenuated in individuals with obesity compared to those with a healthy weight
192 [56]. This may be mediated by an accumulation of body fat which has been proposed to
193 weaken satiety signalling [14, 32, 33], perpetuating overeating in individuals with excess body
194 fat and obesity. However, it is important to note that palatability of food per se may not lead to
195 overconsumption but is it rather the high energy density associated with palatable foods rich
196 in fat and sugar that is driving the increase in EI [39]. For example, consumption of highly
197 palatable artificially sweetened low-calorie foods may not cause overconsumption of energy
198 at a particular meal (but some studies have shown that consuming artificial sweeteners can
199 lead to overconsumption at subsequent meals [57]).

200

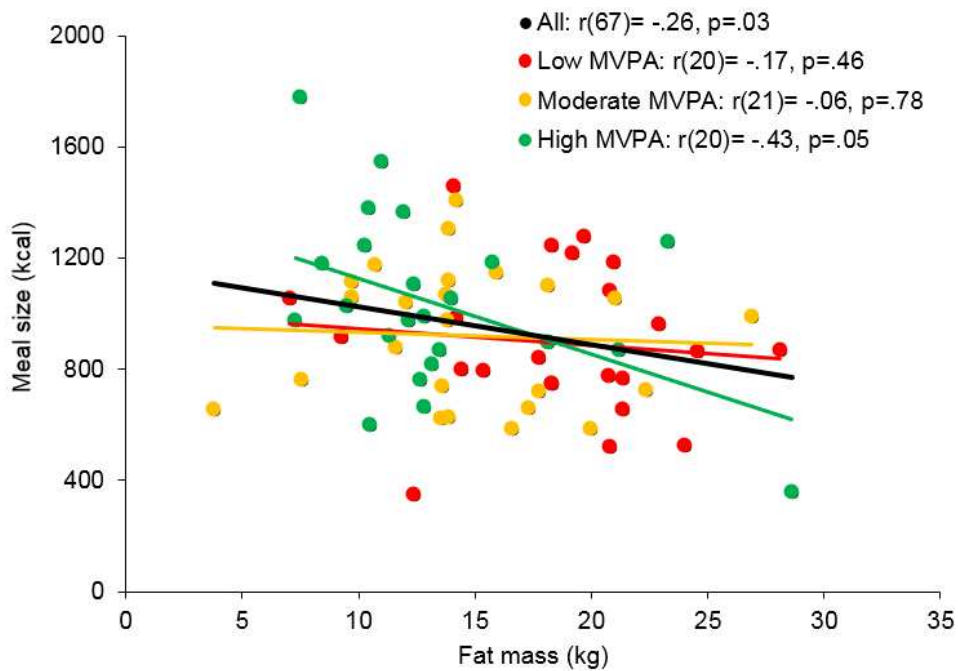
201 **4. Energy intake and appetite control along the spectrum of physical activity levels**

202 While there is evidence for the negative feedback mechanisms involved in satiation
203 and satiety based on the interaction between tonic adiposity and episodic gut signals, less is
204 known on the factors that drive hunger and food intake. Whether EE and PA are drivers of EI
205 has not been well understood [58]. The relationship between EE, PA and EI in humans was
206 examined over 50 years ago [59-61]. Mayer et al. demonstrated a relationship between
207 occupational PA and daily EI in Bengali jute mill workers whose daily occupations ranged from
208 “sedentary” to “very heavy work” whereby those performing very heavy work consumed more
209 than those performing light work [61]. In line with Mayer, Edholm et al. [60] found a strong
210 relationship between TDEE and daily EI in army cadets over three weeks. Despite providing
211 initial evidence for physiological processes and behavioural activities impacting on appetite
212 and providing a demand for food intake, this concept was left dormant for several decades.

213 The roles of body composition and EE in driving food intake have recently been re-
214 examined and have gathered attention within the scientific community [62, 63]. Fat-free mass
215 has been found to be strongly positively associated with EI in lean and overweight/obese
216 individuals [64, 65], corroborating findings from earlier but less known studies [66, 67]. In

217 contrast, an inverse relationship between fat mass and EI, and hunger and EI was found in
218 lean but these associations were weaker and less consistent in overweight and obese
219 individuals [64, 66-68]. These findings are in line with the proposition that negative feedback
220 signals reflecting energy stores inhibiting food intake are blunted with higher body fat [14, 31].
221 In addition to fat-free mass, RMR has also been shown to predict EI and hunger [1], which led
222 to the suggestion that RMR (largely determined by fat-free mass) exerts a tonic day-to-day
223 signal for hunger and the drive to eat [69]. It has recently been shown that the associations
224 between fat-free mass and EI are mediated by RMR [3] and TDEE [70], suggesting that the
225 associations between fat-free mass and EI reflect the energetic demands of metabolically
226 active tissue.

227 In the 70 individuals from Figure 1, we found that both fat-free mass and fat mass were
228 significantly positively and inversely associated with meal size, respectively, corroborating
229 prior studies in lean individuals [62, 66]. Interestingly, exploratory analyses suggested that the
230 strength of the association between fat mass and meal size may be moderated by PA level as
231 the association was strongest in those with the highest time spent in moderate-to-vigorous PA
232 (MVPA) when divided by sex-stratified tertiles (Figure 2) [Beaulieu et al., unpublished results].
233 However, the mechanisms responsible for this effect are unknown and whether these stem
234 from a direct effect of PA on fat mass or indirectly through other physiological, behavioural or
235 psychological factors remains to be elucidated.



236

237 **Figure 2** Relationship between fat mass and meal size within sex-stratified tertiles of
 238 moderate-to-vigorous PA (MVPA; n=70). The strength of the inverse association between fat
 239 mass and energy intake was found to be strongest in those with the highest levels of PA.
 240 From Beaulieu et al. [unpublished results].

241

242 The contribution of PA per se towards the drive to eat is less apparent. In comparison
 243 to RMR, PAEE makes up a smaller portion of TDEE and is more variable; therefore, its impact
 244 on EI may be harder to quantify. However, it can be proposed that PAEE (behavioural EE) will
 245 be a determinant of EI just like RMR (but weaker overall and with great individual variability –
 246 see Figure 1). Recently, a systematic review and a meta-analysis concluded that there is little
 247 evidence that PA or exercise, whether acute or chronic, leads to changes in EI [71, 72].
 248 However, the acute or relatively short-term nature of these studies may not have been long
 249 enough to demonstrate a compensatory rise in EI with habitual PA [73] and as originally
 250 demonstrated by Mayer et al. [61]. Indeed, a strong relationship was found between

251 objectively-measured PA (activity counts) and EI (food records) in 300 middle-aged women
252 [74]. It is important to note that the study by Mayer et al. revealed two separate effects of
253 habitual PA level on EI, characterised by a J-shape relationship. In the jute mill workers with
254 higher levels of occupational PA (e.g. “medium” to “very heavy” work), daily EE and EI were
255 closely matched, but at low levels of occupational PA where body mass was also greater, this
256 coupling was lost, such that daily EI exceeded EE in those performing “sedentary” to “light”
257 work [61].

258

259 **4.1 The zones of appetite control**

260 Based on the study by Mayer et al. [61], Blundell proposed that appetite control is
261 enhanced with increasing levels of PA [75]. In contrast, physical inactivity could not only
262 reduce TDEE but also lead to appetite dysregulation, overconsumption and eventually weight
263 gain [75]. Indeed, according to Jacobs [76], “the late Henry L Taylor favoured a model that
264 linked EI to EE in a J-shaped curve (personal communication, late 1970s). The first part of his
265 concept was that EI is in exact homeostasis with EE under conditions of high EE. The second
266 part was that there is a failure of homeostasis in a sedentary lifestyle because of its
267 accompanying low EE. He postulated that body signals go awry in sedentary lifestyles; when
268 a person does no physical work, the body will not recognize that it is being overfed. Sedentary
269 persons may lose the innate ability to compensate for inactivity by reducing their eating”
270 (p.189). It is important to note here that “sedentary” lifestyles used by this author should in fact
271 be interpreted as inactive lifestyle in light of current definitions. Thus, Blundell revisited the
272 Mayer J-shaped curve and suggested that individuals with low levels of PA could be
273 considered as being within a “non-regulated zone” of appetite control, whereas those with
274 higher levels of PA could be within a “regulated zone” of appetite control [75].

275 While this model of appetite control and EI along the spectrum of PA levels was
276 originally based on limited evidence, it has recently been supported [7, 77]. In a systematic

277 review using data from 10 cross-sectional studies that compared EI in active and inactive
278 individuals, we plotted standardized EI (z-scores) according to four PA levels ranging from low
279 to very high. This analysis revealed a clear J-shape relationship between PA level and EI [7].
280 Similarly, Shook et al. estimated EI based on changes in body composition across quintiles of
281 PA in a large sample of young adults and again demonstrated appetite dysregulation in those
282 with the lowest PA [77].

283

284 **5. The impact of physical activity and exercise on the mechanisms of appetite control**

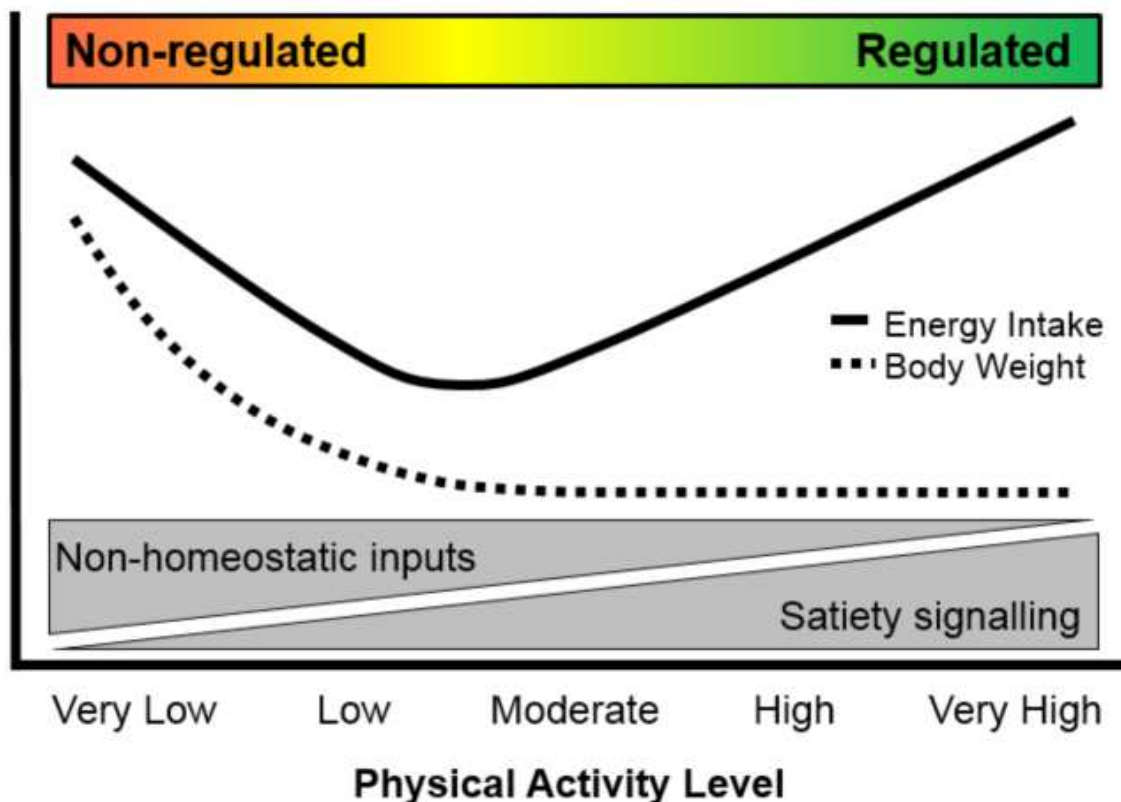
285 Emerging studies are shedding light on the mechanisms contributing to the proposed
286 dysregulation of appetite at lower levels of PA and more sensitive appetite control at higher
287 levels of PA. These mechanisms may not necessarily be the same along the whole spectrum
288 of PA levels. Acutely, exercise has been shown to influence gastric emptying [78], attenuate
289 the release of ghrelin and increase the secretion of PYY, GLP-1 and pancreatic polypeptide
290 [79]. Chronic exercise may increase the postprandial secretion of GLP-1 and PYY [80].
291 Therefore, habitual PA (and exercise) may interact with food intake to enhance hormonal
292 satiety signalling [81]. Moreover, regular PA and exercise training are associated with several
293 other physiological adaptations such as improved sensitivity to insulin [82] and leptin [83, 84],
294 substrate metabolism [85], and body composition [86], which have been proposed as
295 mechanisms involved in food intake and eating behaviour [87, 88]. Evidence suggests that PA
296 influences appetite control through a dual-process action which increases the drive to eat but
297 also post-meal satiety [89]. Several studies have now demonstrated that physically active
298 individuals show better energy compensation than their less active counterparts following
299 consumption of preloads differing in energy content such that they reduce EI to offset the
300 difference in energy consumed from the preloads [10, 90-94]. This preload-test meal paradigm
301 is effective in measuring the strength of satiety [95]. These improvements in satiety may be
302 associated with exercise-induced adaptations in episodic satiety signalling [80, 91, 96] or
303 gastric emptying [97]. In contrast, we have shown that satiation does not appear to be

304 influenced by PA level in non-obese individuals when measured with a passive
305 overconsumption paradigm comparing ad libitum EI at meals high in fat or carbohydrate [9].
306 However, in overweight and obese individuals, Caudwell et al. [1] showed that exercise
307 training led to a reduction in EI at a high-fat/energy dense test meal. This may be associated
308 with enhanced satiation, but given the homeostatic and non-homeostatic determinants of food
309 intake, may also reflect changes in food hedonics or behavioural traits.

310 While there is strong support that habitual PA affects homeostatic mechanisms
311 controlling food intake, less is known on its effect on non-homeostatic processes, and their
312 contribution to appetite control along the spectrum of PA levels. Indeed, secondary analyses
313 from the study from Caudwell et al. [1] revealed a reduction in hedonic wanting for high-fat
314 foods relative to low-fat foods (Leeds Food Preference Questionnaire), disinhibition and binge
315 eating following exercise training [Beaulieu et al., unpublished results]. Furthermore, another
316 study from our group found inverse associations between time spent in MVPA and disinhibition
317 and binge eating, but these did not remain significant after controlling for body fat [24]. This
318 corroborates a study by Shook et al. who found greater disinhibition in their lowest quintile of
319 MVPA but not when controlling for body weight [77], and the aforementioned reduction in
320 disinhibition and binge eating score following 12 weeks of exercise training which did not
321 remain significant after controlling for change in body fat [Beaulieu et al., unpublished results].
322 These differences in disinhibition and binge eating were not apparent in non-obese individuals
323 varying in PA levels [9, 10], suggesting the influence of habitual PA on eating behaviour traits
324 may be more strongly influenced by body composition. In terms of food hedonics, differences
325 in the rewarding value of foods (liking and wanting) have been observed in lean active
326 compared to overweight inactive males [98], but in non-obese individuals, PA level did not
327 influence liking and wanting for high-fat food in the hungry or fed states [9, 10]. In inactive
328 individuals with overweight and obesity, 12 weeks of exercise training (125-250 kcal per
329 exercise session) did not affect liking or wanting [99], whereas another 12-week intervention
330 at a higher dose of exercise (500 kcal per exercise session) reduced the hedonic wanting for

331 high-fat food independent of changes in body fat [Beaulieu et al. unpublished results].
332 Moreover, habitual (self-reported) PA may differently impact food cravings depending on
333 exercise type and sex [100]. These studies suggest there may be differing effects of PA on
334 non-homeostatic appetite control according to an individual's body fat status, sex, and dose
335 and type of PA.

336 Therefore, we can propose that in addition to individuals with non-regulated appetite
337 having blunted satiety signalling, excess body fat in these individuals may amplify non-
338 homeostatic inputs favouring overconsumption. In contrast, individuals with regulated appetite
339 with higher levels of PA have enhanced postprandial sensitivity, allowing for EI to be better
340 matched to EE in response to hunger and satiety signals. This is demonstrated in an updated
341 perspective of the zones of appetite control in Figure 3.



342

343 **Figure 3** An updated perspective of appetite control along the spectrum of PA levels based
344 on the study by Mayer et al. [61] and Blundell [75]. Individuals with non-regulated appetite
345 have lower levels of PA, higher body fat, greater non-homeostatic influences favouring

346 overconsumption and weaker satiety response to food. Those with regulated appetite have
347 higher levels of PA, lower body fat, increased drive to eat and enhanced satiety response to
348 food.

349

350 **6. Interaction between physical activity and diet composition on energy intake and** 351 **energy balance**

352 While PA appears to affect several mechanisms of appetite control, as described
353 above, whether it renders individuals less susceptible to overconsumption in the current
354 obesogenic food environment has not been extensively examined. This is important to
355 consider, with headlines stating “You cannot outrun a bad diet” [101]. Only a few studies have
356 investigated the impact of PA and diet composition on EI and energy balance. A study by
357 Tremblay et al. [102] in males found that consumption of a high-fat diet over two days following
358 a 500-kcal exercise bout led to a positive energy balance, whereas consumption of a low-fat
359 diet was able to maintain the energy deficit produced by exercise. Along those lines,
360 Murgatroyd et al. [103] showed in males that increasing the dietary fat content (and energy
361 density) of an ad libitum diet in a day where exercise was imposed (~675 kcal) increased EI
362 and led to a positive energy balance (albeit not statistically significant). Moreover, consumption
363 of a high-fat diet while imposing inactivity resulted in a daily positive energy balance of
364 approximately 1000 kcal more than with imposed exercise, and 1200 kcal more than with
365 exercise on a low-fat diet. Other studies in males [104] and females [105] corroborated these
366 findings by demonstrating that the consumption of a high-fat meal following an exercise bout
367 resulted in significantly greater relative EI (after considering the EE of the exercise) compared
368 with a low-fat meal. Interestingly, palatability of both high-fat and low-fat meals increased after
369 exercise compared to rest in females, but not in males [105]. These studies highlight the
370 potency of the phenomenon of passive overconsumption. Therefore, the degree of
371 compensation observed in response to PAEE can readily be modulated simply by altering the
372 energy density of the diet and proposals about compensation need to be interpreted with care.

373 Whether being habitually physically active enhances the response to dietary
374 manipulations is also of interest. As discussed above, we have shown that physically active
375 individuals are also prone to acute passive overconsumption with an imposed high-fat meal
376 [9]. In addition, while individuals with higher levels of PA were found to be sensitive to the
377 acute nutritional manipulation of preloads varying in energy content by reducing EI at the
378 following meal, objectively-measured daily EI (including the preload) was greater after a high-
379 energy relative to a low-energy preload regardless of PA level, demonstrating an effect of
380 passive overconsumption [10]. Others have shown that active individuals may compensate
381 beyond the immediate meal following intake of a high-energy preload, attenuating the risk of
382 overconsumption, but this was measured with food records and daily EI including the preload
383 was not reported [90, 91, 93]. The long-term compensatory response to high energy density
384 food consumption in physically active individuals is unknown. The EE associated with PA may
385 be helpful in mitigating episodes of overconsumption and fluctuations in EI over time [106,
386 107]. However, given the available evidence, in the general population, higher levels of
387 habitual PA in conjunction with a diet lower in energy density appear to be optimal for appetite
388 control and energy balance.

389

390 **7. Implications and future directions**

391 The impact of PA on the mechanisms of appetite control has implications for individuals
392 wishing to lose fat mass through exercise as large variability in the individual response to
393 exercise interventions have been observed [89, 108]. These varying responses in fat loss to
394 exercise training suggest that some individuals compensate for the increase in PA (and EE)
395 through greater food intake or other mechanisms impacting on energy balance, minimizing
396 the effect of exercise on fat loss. In both those susceptible and resistant to exercise-induced
397 weight loss, hunger and the strength of satiety were enhanced with exercise training, showing
398 a robust effect of the dual-process action of PA on appetite control; however, the increase in

399 hunger was greater in those resistant to weight loss [89]. The compensatory adaptations in
400 appetite control and eating behaviour following exercise-induced weight loss are beyond the
401 scope of this review and have been reviewed elsewhere [20, 109, 110]. Nevertheless, it should
402 be acknowledged that certain baseline (pre-intervention) characteristics of appetite may
403 predict the susceptibility to exercise-induced weight loss such as the hedonic response to
404 acute exercise [111] and the peptide response to food consumption [112], which is of interest
405 for future research to help personalise interventions to promote successful fat loss with
406 exercise.

407 The role of PAEE in driving EI is important for future research to clarify as it can make
408 up a significant proportion of TDEE in physically active individuals (see Figure 1) [113]. While
409 the influence of PA on some processes of appetite appear to be independent of body fat, more
410 research is required to understand the role of body composition and body fat status in the
411 relationship between PA level and appetite control. We have reported above that PA (or
412 factors associated with PA) may moderate the relationship between fat mass and EI, which is
413 an interesting avenue for future research. Other potential moderators of the relationship
414 between PA and appetite that remain to be examined further include sex [114] and age [93].
415 Additionally, very little is known on how the type, dose, intensity and timing of habitual PA and
416 exercise affect homeostatic and non-homeostatic appetite. The mechanisms responsible for
417 the apparent enhancement in the satiety response to food consumption in physically active
418 individuals also remain to be fully elucidated. Finally, in light of the research on the interaction
419 between PA and dietary manipulations, it is important for future research to take an energy
420 balance perspective [115] to increase our understanding of the complex relationships and
421 interactions among PA, diet composition, body composition and appetite control along the
422 spectrum of PA levels.

423

424 **8. Summary**

425 Food intake is modulated by several homeostatic and non-homeostatic mechanisms
426 controlling appetite. Evidence is accumulating to support the view that EI along the spectrum
427 of PA is J-shaped, with individuals with low levels of PA being in a non-regulated zone of
428 appetite whereas those with higher levels of PA operating in a regulated zone with more
429 sensitive appetite control. Body fat also varies along the spectrum of PA and may impact the
430 sensitivity of satiety signals and non-homeostatic inputs (food hedonics and behavioural traits)
431 favouring overconsumption at lower levels of PA, but this remains to be fully understood. PA
432 affects the homeostatic mechanisms of appetite via a proposed dual-process action of
433 increased drive to eat from greater EE, but also by enhanced satiety response to food, likely
434 through more sensitive postprandial signalling. An important tenet of our current position is
435 that PAEE is a determinant of EI (although with greater variability than RMR). These
436 processes generate a better adjustment of EI to EE in response to hunger and satiety signals
437 at higher levels of PA. However, special attention needs to be given to diet composition, with
438 a high-fat energy-dense diet leading to acute passive overconsumption of energy along the
439 entire spectrum of PA. Importantly, the strength of the various mechanisms and determinants
440 of appetite will vary between individuals along the spectrum of PA, highlighting the need to
441 recognise that the impact of PA on appetite control is not a case of 'one size fits all'.

442

443 **Conflict of interest**

444 The authors declare no conflicts of interest.

445

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