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1 **Gut Hormones, Adipokines, and Pro- and Anti-inflammatory Cytokines/Markers in Loss**
2 **of Control Eating: A Scoping Review**

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Abstract

22
23 Loss of control (LOC) eating is the defining feature of binge-eating disorder, and it has particular
24 relevance for bariatric patients. The biomarkers of LOC eating are unclear; however, gut
25 hormones (i.e., ghrelin, cholecystokinin [CCK], peptide YY [PYY], glucagon-like peptide 1
26 [GLP-1], and pancreatic polypeptide [PP]), adipokines (i.e., leptin, adiponectin), and pro- and
27 anti-inflammatory cytokines/markers (e.g., high-sensitivity C-reactive protein [hsCRP]) are
28 candidates due to their involvement in the psychophysiological mechanisms of LOC eating. This
29 review aimed to synthesize research that has investigated these biomarkers with LOC eating.
30 Because LOC eating is commonly examined within the context of binge-eating disorder, is
31 sometimes used interchangeably with subclinical binge-eating, and is the latent construct
32 underlying disinhibition, uncontrolled eating, and food addiction, these eating behaviors were
33 included in the search. Only studies among individuals with overweight or obesity were included.
34 Among the identified 31 studies, 2 studies directly examined LOC eating and 4 studies were
35 conducted among bariatric patients. Most studies were case-control in design (n=16) and
36 comprised female-dominant (n=13) or female-only (n=13) samples. Studies generally excluded
37 fasting total ghrelin, fasting CCK, fasting PYY, and fasting PP as correlates of the examined
38 eating behaviors. However, there was evidence that the examined eating behaviors were
39 associated with lower levels of fasting acyl ghrelin (the active form of ghrelin) and adiponectin,
40 higher levels of leptin and hsCRP, and altered responses of postprandial ghrelin, CCK, and PYY.
41 The use of GLP-1 analog was able to decrease binge-eating. In conclusion, this review identified
42 potential biomarkers of LOC eating. Future studies would benefit from a direct focus on LOC
43 eating (especially in the bariatric population), using longitudinal designs, exploring potential
44 mediators and moderators, and increased inclusion of the male population.

45 **Key words**

46 Loss of control eating; binge-eating; gut hormones; adipokines; inflammation; cytokines

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47

Abbreviations that are not Standard in the Field

48 LOC eating¹

49

50 Loss of control eating

51

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52 **1. Introduction**

53 Binge-eating disorder (BED), a mental illness that affects 2.8% of women and 1.0% of
54 men worldwide¹, is characterized by recurrent binge-eating episodes in the absence of
55 compensatory behaviors². Binge-eating consists of two intercorrelated but dissociable features:
56 loss of control (LOC) eating and overeating^{2,3}. LOC eating is a subjective experience of being or
57 feeling out of control when eating, no matter the amount of food consumed. As individuals who
58 report LOC eating with or without overeating present similar psychosocial profiles^{3,4}, LOC
59 eating has been widely recognized as the most salient feature of BED. Notably, emerging
60 evidence has supported the validity of LOC eating as an independent construct³. For example,
61 independent of overeating, LOC eating has unique cross-sectional and prospective associations
62 with various physiological and psychosocial vulnerabilities, such as obesity⁵, metabolic
63 syndromes^{6,7}, general psychopathology (e.g., depression), and eating-related psychopathology
64 (e.g., body dissatisfaction)^{8,9}.

65 The clinical relevance of LOC eating is particularly prominent in patients with severe
66 obesity who seek or have undergone bariatric surgery. First, LOC eating is prevalent in this
67 population, affecting 6.6%¹⁰-78.6%¹¹ of the preoperative and 5.4¹²-50.7%¹³ of the postoperative
68 patients. Second, LOC eating at post-surgery predicts adverse surgical outcomes, including less
69 weight loss¹⁴, weight regain^{12,15}, and more surgical complications (e.g., dumping, vomiting)^{16,17}.
70 Finally, and most importantly, as bariatric surgery dramatically reduces gastric volume, alters the
71 gastrointestinal environment, and restricts eating capacity, overeating becomes physically
72 difficult or impossible after surgery (at least in the short-term). Therefore, the overeating feature
73 inherent in binge-eating may not apply to postoperative patients, making the importance of LOC
74 eating stand out.

75 With increasing emphasis being placed on LOC eating, many efforts have been made to
76 identify its physiological and psychosocial mechanisms. Physiologically, LOC eating involves a
77 disruption in the homeostatic and hedonic eating regulation systems. The homeostatic system
78 controls physiological appetite, hunger, and satiety through the hypothalamus, especially the
79 hypothalamic arcuate nucleus¹⁸. Individuals with LOC eating have imbalanced expressions of
80 orexigenic and anorexigenic neurons within the arcuate nucleus, reporting increased appetite,
81 increased hunger, and decreased satiety compared to healthy controls¹⁹. The hedonic system
82 regulates eating behavior mainly through the dopamine and opioid reward circuits, which
83 modulates the “wanting” (food craving) and “liking” (food enjoyment) components of food
84 reward, respectively. An accumulating body of behavioral and neuroimaging studies has shown
85 that individuals with LOC eating self-report an elevated food craving/enjoyment and exhibit
86 heightened brain reward responsivity for high-energy foods (e.g., fats, sweets)^{20,21}. Consequently,
87 they are more likely to consume high-energy foods than those without LOC eating^{22,23}.

88 The psychosocial mechanisms of LOC eating have been widely studied both within and
89 outside the bariatric population, and several types of risk factors have been identified. These
90 psychosocial risk factors include negative emotions or affects (e.g., anxiety, depression,
91 distress)^{24,25}, maladaptive emotion regulations strategies (e.g., rumination, suppression)^{26,27},
92 weight-related dysfunctional behaviors or perceptions (e.g., dietary restraint, weight suppression,
93 and body dissatisfaction)^{28,29}, and deficits in cognitive control, especially inhibitory control^{30,31}.

94 While the physiological and psychosocial mechanisms of LOC eating have been
95 somewhat uncovered, the biomarkers that are related to LOC eating are less studied in the
96 general or bariatric population. Gut hormones, adipokines, and pro- and anti-inflammatory
97 cytokines/markers are promising candidates because they are involved in homeostatic and

98 hedonic eating regulation. As a result, the alterations of these biomarkers may increase or
99 decrease individuals' appetite, hunger, satiety, and perceived food reward, thus ultimately
100 influencing their motivations to initiate or stop consuming food and contributing to the
101 experience of difficulty in stopping eating (LOC eating). Additionally, these biomarkers are
102 closely related to the psychosocial risk factors of LOC eating³²⁻³⁷.

103 Gut hormones include the "hunger" hormone, ghrelin, and "satiety" hormones such as
104 cholecystokinin (CCK), peptide YY (PYY), glucagon-like peptide 1 (GLP-1), and pancreatic
105 polypeptide (PP). Ghrelin activates the orexigenic and dopaminergic neurons, thus promoting
106 appetite, hunger, food craving, and food-seeking behaviors^{38,39}. Ghrelin exists in circulation in
107 two major forms: acylated and des-acylated ghrelin. While the majority of circulating ghrelin is
108 des-acylated, the acylated form is thought to be essential for ghrelin's biological activity in
109 appetite stimulation³⁸. In contrast, CCK, PYY, GLP-1, and PP inhibit orexigenic neurons, thus
110 promoting satiety and eating termination. Additionally, there is evidence that GLP-1 also
111 suppresses dopaminergic neurons⁴⁰. Besides participating in hemostatic and hedonic eating
112 regulations, gut hormones also regulate moods and cognitive functions that are related to LOC
113 eating. For example, research has repeatedly demonstrated the anti-depressant effect of ghrelin³²
114 and the cognitive-enhancing effect of GLP-1³³.

115 Leptin and adiponectin are adipokines that are mostly secreted by white adipose tissue.
116 Leptin acts as a negative feedback signal to control energy homeostasis at the hypothalamus^{41,42},
117 and it also suppresses dopamine signaling to reduce the craving or motivation to seek and
118 consume food⁴². Although there is no consensus, it has been suggested that adiponectin regulates
119 eating behavior in a glucose-dependent fashion. At low glucose conditions, adiponectin
120 downregulates orexigenic and upregulates anorexigenic neurons to attenuate appetite; at high

121 glucose levels, adiponectin downregulates both orexigenic and anorexigenic activities^{43,44}. In
122 addition to regulating eating behaviors, as the receptors of leptin and adiponectin are widely
123 distributed in hippocampus and neocortex, they are likely to be involved in emotion regulation
124 and cognitive control^{34,36}.

125 Pro-inflammatory cytokines, including interleukin-1 α (IL-1 α), interleukin-1 β (IL-1 β),
126 interleukin-6 (IL-6), and tumor necrosis factor- α (TNF- α) are involved in eating regulation
127 possibly through acting on hypothalamus^{45,46} and dopaminergic neurocircuits^{47,48}, and through
128 interacting with eating-regulation hormones (e.g., ghrelin, GLP-1, leptin)^{49,50}. Despite
129 inconclusive evidence, animal and human studies have shown that IL-1 β , IL-6, and TNF- α
130 suppress appetite^{51,52} and eating motivation^{53,54}, hence reducing meal size and meal duration^{55,56}.
131 Furthermore, pro-inflammatory cytokines are intertwined with emotions and moods, such as
132 depression, anxiety, and stress³⁷, which are all risk factors of LOC eating.

133 The pro-inflammatory cytokines are positively associated with inflammatory markers: C-
134 reactive protein (CRP) and erythrocyte sedimentation rate (ESR). In contrast, the synthesis of
135 pro-inflammatory cytokines is inhibited by anti-inflammatory cytokines, such as interleukin-10
136 (IL-10) and interleukin-13 (IL-13). Therefore, these inflammatory markers and anti-
137 inflammatory cytokines are potentially related to LOC eating, although their effect on eating
138 regulation is unclear.

139 Recognizing the role of gut hormones, adipokines, and inflammatory cytokines/markers
140 in eating regulation and psychosocial functioning, prior studies have reviewed their relationships
141 with eating disorders⁵⁷⁻⁵⁹. However, these reviews predominantly focused on bulimia nervosa
142 and anorexia nervosa, of which information cannot be readily applied to LOC eating due to the
143 distinctive disease symptoms among these eating behaviors⁵⁷⁻⁵⁹. To address this gap, the aim of

144 this review was to identify and synthesize quantitative research that has thus far investigated the
145 associations of gut hormones, adipokines, or pro- and anti-inflammatory cytokines/markers with
146 LOC eating in both children and adults. A greater understanding of the biomarkers related to
147 LOC eating could advance the identification, prevention, and treatment of this disordered eating
148 behavior. Additionally, as LOC eating is prevalent and negatively impacts weight loss and
149 metabolic outcomes among bariatric patients, such knowledge could facilitate the development
150 of interventions to promote optimal surgical outcomes.

151 **2. Methods**

152 **2.1 Study design**

153 A scoping review was conducted to examine the range and extent of existing evidence
154 available on gut hormones, adipokines, and inflammatory cytokines/markers related to LOC
155 eating. A scoping review instead of a systematic review was chosen because there are few
156 publications on this topic, and the goal of a scoping review is to examine emerging and unclear
157 evidence⁶⁰. A standard five-stage process of scoping review was followed: (1) identifying the
158 research question, (2) identifying relevant studies, (3) selecting studies, (4) charting the data, (5)
159 collating, summarizing, and reporting the results⁶⁰.

160 **2.2 Inclusion/exclusion criteria**

161 Although LOC eating has been increasingly accepted as an independent construct and
162 there is a growing call to study LOC eating and overeating separately^{3,61,62}, in most cases, it is
163 examined within the context of BED. Additionally, subclinical binge-eating is sometimes used
164 interchangeably with LOC eating. Therefore, both BED and subclinical binge-eating were
165 included in the search. Moreover, given studies have provided evidence that disinhibition,
166 uncontrolled eating (a loss of control on food intake in response to emotional or external stimuli),

167 and food addiction (a loss of control on food intake accompanied by other symptoms such as
168 tolerance and withdraw) share the latent construct of “loss of control”^{63,64}, these eating behaviors
169 were also included in the review. Studies that examined bulimia nervosa were excluded because
170 although it has the defining feature of binge-eating, the levels of hormones, adipokines, and
171 inflammatory cytokines/markers could be influenced by compensatory behaviors such as purging
172 behaviors and vomiting.

173 It is known that the levels and functions of gut hormones, adipokines, and inflammatory
174 cytokines/markers are influenced by body weight. For example, in individuals with obesity,
175 levels of ghrelin and adiponectin are generally lower while levels of leptin and pro-inflammatory
176 cytokines/markers are elevated^{65,66}. To eliminate the confounding effect of body weight and
177 increase the potential applicability to bariatric patients, only studies conducted among
178 individuals with overweight or obesity (body mass index [BMI]>25 kg/m²) were included.

179 In general, studies were included if they 1) reported associations of gut hormones,
180 adipokines, or inflammatory cytokines/markers with LOC eating, subclinical binge-eating, BED,
181 disinhibition, uncontrolled eating, or food addiction among individuals with overweight or
182 obesity, or 2) compared the biomarkers between individuals with these eating behaviors and
183 unaffected controls, where participants in both groups were overweight or obese, or 3)
184 manipulated the biomarkers and reported an effect on these eating behaviors among individuals
185 with overweight or obesity.

186 Studies published in non-English languages, and animal studies, abstracts, editorials, case
187 studies, book chapters, dissertation work, review papers, or kin studies were excluded.

188 **2.3 Search strategy**

189 Three electronic databases, including PubMed (1947–March 5, 2021), PsycINFO (1967–
190 March 5, 2021), and Embase (1974–March 5, 2021) were searched to obtain relevant studies.
191 The search included the combination of the following keywords: gut hormones, ghrelin, CCK,
192 PYY, GLP-1, PP, leptin, adiponectin, inflammation, IL-1 α , IL-1 β , IL-6, TNF- α , IL-10, IL-13,
193 CRP, ESR, LOC eating, binge-eating, disinhibition, uncontrolled eating, and food addiction. The
194 database search was complemented by a hand search of the reference lists obtained from the
195 identified articles.

196 **2.4 Study selection**

197 A total of 31 studies were included in the final review. The study selection flow is
198 presented in Figure 1. Author 1 (YY) screened titles and abstracts, both author 1 and author 2
199 (WZ) reviewed full-text articles.

200 **2.5 Data extraction and synthesis**

201 Key components extracted from each paper included: 1) study characteristics of title, first
202 author, country, publication year, and study design; 2) participant characteristics of gender, age,
203 race, BMI, sample size, and inclusion/exclusion criteria; 3) biomarkers assessed; 4) eating
204 behaviors assessed; and 5) main study conclusion.

205 **3. Results**

206 **3.1 Study and participant characteristics**

207 The extracted data were organized according to individual biomarkers (Table 1). In terms
208 of study characteristics, the 31 studies were conducted in 13 different countries with Italy (n=9)
209 and US (n=8) accounted for the majority. Eleven studies were published in the last 5 years and 9
210 in the last 10 years. Most of the studies were case-control in design (n=16), followed by
211 longitudinal (n=6), cohort (n=2), cross-sectional (n=4), randomized controlled trial (RCT, n=2),

212 and randomized cross-over (n=1) designs. For the 16 case-control and 2 cohort studies, except
213 for 1 study⁶⁷, BMI was either comparable between groups⁶⁸⁻⁸⁰ or was adjusted as a covariate in
214 analysis⁸¹⁻⁸⁴. Regarding sample characteristics, 4 studies were conducted among pre- or post-
215 bariatric patients^{75,85-87}, and participants from the remaining studies were patients who were
216 seeking eating disorder (n=3) or behavioral weight loss treatment (n=6), were attending pediatric
217 (n=3), endocrinology (n=3), or psychiatric clinics (n=1), non-treatment seeking (n=5), a mix of
218 treatment-seeking and non-treatment seeking (n=2), and not specified (n=4). Six studies were
219 conducted among children or adolescents^{78,79,83,84,88,89}, and the remaining 25 were among adults.
220 Except for 1 study that did not report sex composition⁹⁰ and 4 studies that comprised comparable
221 proportions of males and females^{72,84,88,91}, studies were female-dominant (n=13) or female-only
222 (n=13). Two studies directly examined LOC eating^{78,83} and 6 studies examined subclinical binge-
223 eating^{72-74,84,90,92} (none of these were conducted among bariatric population), others examined
224 BED (n=13), disinhibition or uncontrolled eating (n=8), and food addiction (n=2).

225 **3.2 Biomarkers Related to LOC Eating, Subclinical Binge-Eating, BED, Disinhibition,** 226 **Uncontrolled Eating, or Food Addiction**

227 **3.2.1 Fasting ghrelin**

228 Ghrelin is an appetite-stimulating peptide that increases food intake. In a healthy
229 population, circulating levels of ghrelin increase during fasting to promote meal initiation and
230 decrease shortly after meal consumption to terminate an eating episode³⁸. Ghrelin has two major
231 molecular forms (acylated and des-acylated ghrelin)—only the acyl ghrelin is able to bind to the
232 ghrelin receptor and stimulate appetite⁹³.

233 Twelve studies have examined fasting ghrelin in relation to subclinical binge-eating,
234 BED, disinhibition, uncontrolled eating, or food addiction. The majority (n=9) did not support a
235 role of fasting total ghrelin in these eating behaviors. Specifically, 3 studies reported that fasting

236 total ghrelin did not differ between individuals with subclinical binge-eating⁸⁴, BED⁶⁸, or food
237 addiction⁶⁹ and unaffected controls. Next, 5 studies did not find significant cross-sectional or
238 longitudinal associations between fasting total ghrelin and disinhibition^{86,94,95} or uncontrolled
239 eating^{85,87} among patients who sought or had undergone surgical or behavioral weight loss
240 treatment. Finally, 1 longitudinal study⁷⁰ assessed fasting total ghrelin and BED before and after
241 a cognitive-behavioral therapy (CBT). This study found that fasting total ghrelin did not respond
242 to the intervention, and it failed to predict binge-eating behaviors at pre- or post-intervention.

243 In contrast to these null findings, an earlier study found that fasting total ghrelin was
244 lower in women with BED relative to non-BED controls, and this difference was normalized
245 after CBT⁷¹. However, results should be interpreted with caution because the ghrelin changes
246 from pre- to post-intervention were independent of the CBT treatment and were not related to the
247 self-reported binge-eating days⁷¹.

248 Unlike the above studies that examined total ghrelin, two recent studies^{72,92} specifically
249 examined its active form—acyl ghrelin—and reported significant findings. One study compared
250 the fasting acyl ghrelin between adults with and without subclinical binge-eating, which revealed
251 that participants with subclinical binge-eating had significantly lower fasting acyl ghrelin
252 concentrations than unaffected controls⁷². Similarly, a significant negative association was
253 observed between fasting acyl ghrelin and binge-eating behaviors in a cross-sectional study
254 conducted among 88 adults with overweight or obesity⁹².

255 Overall, studies appeared to be consistent in indicating that fasting total ghrelin did not
256 have a role in subclinical binge-eating, BED, disinhibition, uncontrolled eating, and food
257 addiction. However, some evidence supports a reduced level of acyl ghrelin in patients with
258 subclinical binge-eating that warrants future investigation.

259 3.2.2 Postprandial ghrelin

260 Postprandial ghrelin (total or acylated) has been assessed in 6 studies with conflicting
261 results. In one study⁷¹ conducted among women with BED (n=10) and controls (n=9), the
262 authors found that postprandial total ghrelin was lower and showed a blunted decline following
263 meal consumption in the BED group after adjusting for weight change. One recent study⁷² had
264 similar findings in that they documented that acyl ghrelin (but not des-acyl ghrelin) was
265 significantly lower and declined slower after a meal in adults with subclinical binge-eating (n=20)
266 than in unaffected controls (n=22). However, 3 other studies^{70,81,84} were unable to replicate this
267 finding, which reported levels and responses of total ghrelin following test meals were
268 comparable between adults or adolescents with and without BED. Additionally, a study that was
269 conducted among bariatric patients did not detect any significant associations between
270 postprandial total ghrelin and uncontrolled eating before or at 12 months following bariatric
271 surgery⁸⁷.

272 Two studies of the above 6 studies also tested whether postprandial ghrelin changed
273 following CBT. One study⁷¹ reported that the lower level and blunted response of postprandial
274 total ghrelin in the BED group were “normalized” after CBT, although the “normalization” could
275 not be conclusively attributed to the intervention. Conversely, the other study⁷⁰ did not find any
276 intervention effect on postprandial total ghrelin level or response, despite 50% of the patients
277 achieved BED abstinence at the end of the intervention.

278 The inconsistent results among these studies could be partially attributed to a lack of
279 differentiation the two forms (acylated and des-acylated) of ghrelin, which may bias the study
280 results. Furthermore, the small sample sizes (ranged from 6 to 20 in the binge-eating group),
281 different participant characteristics (e.g., 5 different countries: US, Italy, UK, Canada,

282 Switzerland; a wide age range: mean age ranged from 13 to 50), and methodological differences
283 across the studies could also contribute to the mixed findings. For example, the 6 studies used 6
284 different test meals with different total energy (ranged from 300 kcal to 797 kcal) and energy
285 compositions. In addition, they adjusted for different covariates (e.g., age^{72,81}, sex^{70,72,81,84},
286 BMI^{70,72,81,87}, weight change⁷¹, fat mass⁸⁴), were comprised of treatment-seeking^{81,84,87} or non-
287 treatment seeking samples⁷⁰, and adopted different outcome measures (e.g., Eating Disorder
288 Examination^{70,72,84}, Questionnaire on Eating and Weight Patterns^{71,72}, Three Factor Eating
289 Questionnaire⁸⁷).

290 In summary, study results are mixed in terms of the association between postprandial
291 ghrelin and LOC eating. However, there is some evidence that the level and response of
292 postprandial ghrelin, especially acyl ghrelin, to meal consumption may be lower and blunted in
293 patients with subclinical binge-eating or BED^{71,72}.

294 3.2.3 Fasting and postprandial CCK

295 CCK is a satiety hormone that reduces food intake⁹⁶. In the general population, the
296 circulating level of CCK increases rapidly in response to a meal to promote meal termination⁹⁶.
297 Three studies have examined fasting CCK related to subclinical binge-eating or BED. All 3 of
298 them reported that fasting CCK did not differ between adults with and without binge-eating^{70,72,73}.
299 One of these studies was longitudinal in design, in which 18 patients with BED received an 8-
300 week CBT. Neither the fasting CCK was affected by the intervention, nor it was related to the
301 binge-eating behaviors at post-intervention⁷⁰.

302 Postprandial CCK was assessed in 4 studies and findings were mixed. Two studies^{72,73}
303 did not find significant differences in postprandial CCK between adults with subclinical binge-
304 eating or BED and unaffected controls. However, in one study⁷⁰, patients with BED relative to

305 controls exhibited an augmented CCK secretion within 60 minutes following meal ingestion,
306 which was not corrected by an 8-week CBT. The authors interpreted the heightened CCK
307 stimulation as an initial effort of the central nervous system to prevent individuals with BED
308 from bingeing. In contrast, in another study conducted among a cohort of women with
309 overweight, the authors observed a positive association between blunted CCK secretion and high
310 disinhibition, but the association was significant only among women who had a high level of
311 dietary restraint⁹⁷. This result suggested potential interactions among dietary restraint, CCK
312 responses, and disinhibition.

313 The samples across the 4 studies that examined postprandial CCK tended to share similar
314 characteristics. For example, they were mostly female, middle-aged, from US, and non-treatment
315 seeking. However, all of the studies had small sample sizes (ranged from 11 to 20 in the binge-
316 eating group). Additionally, there were other variations among studies, including various test
317 meals (4 different meals), different covariates controlled in the analysis (e.g., age⁷², sex^{70,72},
318 BMI^{70,72}, and weight change⁹⁷), and varied outcome measurements (e.g., Questionnaire on Eating
319 and Weight Patterns, Eating Disorder Examination).

320 To sum up, despite the limited number of studies that have been done, fasting CCK has
321 been consistently found to be unrelated to subclinical binge-eating or BED. The association of
322 postprandial CCK with BED⁷⁰ or disinhibition⁹⁷ was found in two studies. However, the
323 responses of postprandial CCK were controversial between these studies^{70,97}. Additionally, there
324 is preliminary evidence suggesting the interactions among dietary restraint, disinhibition, and
325 CCK secretion⁹⁷.

326 **3.2.4 Fasting and postprandial PYY**

327 PYY is a hormone that increases satiety and suppresses food intake. In the general
328 population, PYY levels increase within 15 minutes in response to food intake and contribute to
329 termination of food intake⁹⁸. Eight studies have examined fasting PYY with subclinical binge-
330 eating, BED, uncontrolled eating, disinhibition, or food addiction, and all of them reported null
331 findings. In detail, 5 studies documented that fasting PYY did not differ between adults or
332 adolescents with subclinical binge-eating^{72,84}, BED^{70,71}, or food addiction⁶⁹ and unaffected
333 controls. Additionally, 2 of these studies did not find any intervention effect of CBT on fasting
334 PYY among patients with BED^{70,71}. Finally, 3 studies did not find significant cross-sectional or
335 longitudinal associations between fasting PYY and uncontrolled eating⁸⁸ or disinhibition^{94,95}
336 among adolescents or adults with obesity.

337 Postprandial PYY has been assessed in 4 studies, and 3 of them reported that the PYY
338 secretion in response to test meals was comparable between adolescents or adults with
339 subclinical binge-eating^{72,84} or BED⁷¹ and controls. Additionally, postprandial PYY did not
340 respond to a 6-week CBT among patients with BED⁷¹. In contrast, 1 study observed a higher
341 increase in PYY within the first 80 minutes following meal ingestion in patients with BED vs
342 non-BED controls⁷⁰. It was interpreted that the more intense stimulation of the PYY secretion
343 after food intake was an adaptive response from the central nervous system and the gut to
344 counteract the initiation of binge-eating.

345 In summary, current studies did not support fasting PYY as a significant correlate of
346 subclinical binge-eating, BED, uncontrolled eating, disinhibition, or food addiction. While most
347 studies did not detect postprandial PYY alterations in subclinical binge-eating or BED, one study
348 found an augmented response that deserves further investigation⁷⁰.

349 **3.2.5 Fasting and postprandial GLP-1**

350 GLP-1 suppresses appetite, promotes satiety, and reduces energy intake⁹⁹. In the general
351 population, GLP-1 increases rapidly after meal ingestion to prevent overeating⁹⁹. Four
352 observational studies have examined fasting GLP-1 with subclinical binge-eating, BED,
353 uncontrolled eating, and food addiction, and they consistently reported null findings. Among
354 these 4 studies, 3 studies did not find significant differences in fasting GLP-1 between adults
355 with subclinical binge-eating⁷², BED⁷¹, or food addiction⁶⁹ and controls, and the fasting GLP-1
356 did not change after BED treatment⁷¹. In the remaining study, fasting GLP-1 was not related to
357 uncontrolled eating among 12 patients at pre-, 2 and 12 months post-bariatric surgery⁸⁷. Three of
358 the above 4 studies also assessed postprandial GLP-1. They found that postprandial GLP-1 was
359 comparable between adults with and without subclinical binge-eating⁷² or BED⁷¹, was
360 nonresponsive to CBT among patients with BED⁷¹, and was not significantly related to
361 uncontrolled eating among bariatric patients⁸⁷.

362 In contrast to the above observational studies that reported null findings, 3 intervention
363 studies that tested the effect of GLP-1 analog—liraglutide—on subclinical binge-eating, BED,
364 and uncontrolled eating reported significant findings. One randomized controlled trial⁹⁰ assigned
365 non-diabetic patients with subclinical binge-eating to either a 12-week lifestyle intervention
366 (n=21) or lifestyle intervention plus liraglutide (n=21), and results revealed that at the end of the
367 intervention, participants who received liraglutide showed significant improvement in binge-
368 eating. The other 2 studies reported similar results by demonstrating significant reductions of
369 BED among adults with diabetes⁹¹ and reductions of uncontrolled eating among women with
370 polycystic ovary syndrome¹⁰⁰ after 12-week treatment of liraglutide.

371 Taken together, while observational studies consistently reported null findings,
372 intervention studies suggested the possible relevance of GLP-1 to subclinical binge-eating, BED,
373 and uncontrolled eating.

374 **3.2.6 Fasting PP**

375 PP is a gut hormone that reduces food intake⁹⁹. In the general population, PP is released
376 in response to food ingestion to prevent overconsumption of food⁹⁹. Only 2 studies have
377 examined PP, with one study that reported fasting PP did not differ between adults with and
378 without food addiction⁶⁹ and the other study reported no significant association between fasting
379 PP and disinhibition among a cohort of women with obesity⁹⁴.

380 **3.2.7 Leptin**

381 Thirteen studies have examined leptin in relation to LOC eating, subclinical binge-eating,
382 BED, disinhibition, uncontrolled eating, and food addiction, and 5 studies identified significantly
383 higher levels of leptin in these eating behaviors. In detail, 4 studies found significantly higher
384 levels of leptin in adolescents with LOC eating⁷⁸, bariatric candidates or women with BED^{75,101},
385 and adolescents with food addiction⁸⁹ relative to controls. Among these 4 studies, 1 study also
386 reported that higher levels of leptin predicted higher odds of binge-eating behaviors among
387 women with BED¹⁰¹, and another study demonstrated a positive association between leptin and
388 disinhibition among bariatric candidates (regardless of BED diagnosis)⁷⁵. In line with these
389 findings, 1 study reported that leptin levels went up in parallel with the increase of uncontrolled
390 eating among bariatric candidates⁸⁷.

391 There was one study that reported LOC eating as an independent outcome. In that study
392 the authors explored the mediating or moderating effects of dietary restraint and/or sex in the
393 relationship between leptin and LOC eating. Results revealed that the positive relationship

394 between leptin and LOC eating was significant for females only. Moreover, the relationship was
395 partially mediated by higher dietary restraint after controlling for age, race, sex, socioeconomic,
396 fat mass, height, treatment-seeking status, and pubertal status⁷⁸.

397 Unlike the above studies that supported significantly higher levels of leptin in LOC
398 eating, BED, uncontrolled eating, and food addiction, 6 studies reported null findings. However,
399 4 of them found that although lacking statistical significance, the leptin levels were higher in
400 adults or adolescents with subclinical binge-eating⁷⁴, BED^{76,79}, or food addiction⁶⁹ compared to
401 unaffected controls. Importantly, it was found that leptin levels increased linearly along with the
402 increase of binge-eating severity among women with obesity⁷⁴. It should be noted that all these 4
403 studies had a relatively small sample size (ranged from 18 to 35 in the binge-eating or food
404 addiction group), which may be underpowered to detect any statistical significance. There were
405 other 2 studies that did not detect differences of leptin between women with BED⁷³ or
406 disinhibition⁹⁴ and controls; yet, statistics were not provided in these 2 studies.

407 Two studies found lower leptin levels in BED vs non-BED groups. Specifically, in 1
408 case-control study, leptin levels were found to be significantly lower in women with BED than
409 non-BED controls⁶⁷. However, it should be noted that BMI was significantly higher in the non-
410 BED group than the BED group and it was not controlled as a covariate in the analysis, which
411 may explain the contrasting finding in this study. The lower leptin levels in women with vs.
412 without BED were observed in another study, but the difference was minor (46.4 vs. 50.7 μg),
413 and the authors did not provide a level of statistical significance⁷⁷.

414 Overall, with some exceptions, studies generally supported higher leptins levels in LOC
415 eating, subclinical binge-eating, BED, disinhibition, uncontrolled eating, and food addiction

416 (with or without statistical significance). Additionally, there is preliminary evidence that the
417 relationship is mediated by dietary restraint and moderated by sex.

418 **3.2.8 Adiponectin**

419 Three studies have examined adiponectin in relation to BED, disinhibition, and food
420 addiction, and they consistently demonstrated lower adiponectin levels in individuals with these
421 eating behaviors. Specifically, 1 study⁶⁷ observed significantly lower adiponectin levels in
422 women with BED compared to non-BED women; a second study⁶⁹ reported lower adiponectin
423 levels (statistically nonsignificant) in adults with food addiction compared with age-, sex-, BMI-,
424 and physical activity-matched controls; and a third study⁹⁴ detected a negative association
425 between adiponectin and disinhibition among 67 women who participated in a 4-week lifestyle
426 intervention.

427 Taken together, despite the limited number of studies, it appeared that the adiponectin
428 levels were lower in individuals with BED, disinhibition, or food addiction.

429 **3.2.9 Pro- and anti-inflammatory cytokines/markers**

430 Five studies have examined pro- (i.e., IL-1 α , IL-1 β , IL-6, TNF- α , CRP, ESR) and anti-
431 inflammatory (i.e., IL-10) cytokines/markers with LOC eating, BED, disinhibition, and food
432 addiction. CRP has been assessed in 3 studies: 2 of them reported that high-sensitivity CRP
433 (hsCRP) was significantly higher in adolescents with LOC eating⁸³ or adults with BED⁸² than
434 unaffected controls; the remaining study did not find a significant association between CRP and
435 disinhibition among a cohort of women with obesity⁹⁴. Notably, the latter study examined CRP
436 instead of hsCRP and did not adjust for covariates in analysis, which may introduce bias and lack
437 sensitivity in capturing the differences in inflammatory status. ESR has been assessed in 1
438 study⁸², in which significantly higher ESR levels were observed in the BED vs non-BED group.

439 TNF- α has been examined in 2 studies: 1 study showed a significantly lower level of
440 TNF- α in individuals with food addiction compared to those without food addiction⁶⁹; and the
441 other study showed no difference in levels between adults with BED and non-BED controls
442 (statistics not provided)⁸⁰. The different findings may be due to the sample characteristics (e.g.,
443 Canadian⁶⁹ vs Italian⁸⁰, non-treatment seeking⁶⁹ vs treatment seeking⁸⁰), outcome measures (food
444 addiction⁶⁹ vs BED⁸⁰), and covariates controlled in analysis (non-specified⁶⁹ vs depressive
445 symptoms, sex, and age⁸⁰). The latter study also examined other pro- and anti-inflammatory
446 cytokines such as IL-1 α , IL-1 β , IL-6, and IL-10. While no significant differences were detected
447 in IL-1 α , IL-1 β , or IL-6 between the BED and control groups, IL-10 was found to be
448 significantly lower in the BED group⁸⁰.

449 In summary, limited studies have assessed the relationship between inflammatory
450 cytokines/markers and LOC eating. There may be potential associations of LOC eating with
451 elevated inflammatory status, including higher hsCRP, higher ESR, and lower IL-10.

452 **4. Discussion**

453 This scoping review synthesized current evidence regarding gut hormones, adipokines,
454 and pro- and anti-inflammatory cytokines/markers related to LOC eating, subclinical binge-
455 eating, BED, disinhibition, uncontrolled eating, and food addiction, among children or adults
456 with overweight or obesity. Results can be summarized as follows: 1) only 2 studies directly
457 examined LOC eating and 6 examined subclinical binge-eating, and only 4 studies were
458 conducted among bariatric patients; 2) fasting total ghrelin, fasting CCK, fasting PYY, and
459 fasting PP did not appear to be related to the eating behaviors as mentioned above; 3) although
460 studies were scarce and findings were inconsistent, there was evidence supporting lower levels of
461 fasting acyl ghrelin and adiponectin, higher levels of leptin and pro-inflammatory markers (e.g.,

462 hsCRP, ESR), and altered responses of postprandial ghrelin (blunted), CCK (blunted or
463 amplified), and PYY (amplified) in the aforementioned eating behaviors; and 3) using GLP-1
464 analog decreased binge-eating.

465 The lower fasting acyl ghrelin and blunted responses of postprandial ghrelin (total or
466 acylated) observed in individuals with subclinical binge-eating or BED suggested that ghrelin,
467 especially acyl ghrelin, is a potential correlate of LOC eating. Considering that the acyl ghrelin
468 acts to stimulate appetite and promote eating, the lower fasting acyl ghrelin is unlikely to cause
469 binge-eating or LOC eating; instead, it may represent a secondary change or an adaption aiming
470 to counteract repeated binge-eating or LOC eating. The blunted responses of postprandial ghrelin
471 imply that the normal suppression of hunger and food craving after meal consumption is
472 impaired, which may contribute to the initiation or maintenance of binge-eating or LOC eating.
473 Studies in patients with bulimia nervosa have similarly observed an attenuated decrease of
474 postprandial ghrelin¹⁰²⁻¹⁰⁴ following a meal, although it is not clear whether the attenuation is due
475 to the binge behavior or purging behavior. Furthermore, animal studies have provided additional
476 evidence supporting the involvement of ghrelin in binge-eating or LOC eating. For example,
477 there have been observations that ghrelin receptor-deficient mice failed to induce binge-
478 eating^{105,106} and central ghrelin infusion enhanced binge-like behaviors in palatable schedule fed
479 rats¹⁰⁷.

480 It is worth emphasizing that the lower levels of fasting ghrelin and blunted responses of
481 postprandial ghrelin observed in subclinical binge-eating or BED are not universal findings. In
482 addition to reasons like small sample sizes and methodological differences across studies, the
483 failure to distinguish acylated and des-acylated ghrelin may bias and undermine the validity of
484 findings based on total ghrelin concentration. Given recent evidence that des-acylated ghrelin

485 can impair the orexigenic actions of acyl ghrelin^{108,109}, future studies should examine the separate
486 forms rather than the total concentration of ghrelin.

487 Included studies consistently showed that fasting CCK, PYY, and PP did not differ
488 between individuals with and without subclinical binge-eating, BED, uncontrolled eating,
489 disinhibition, or food addiction, indicating that these fasting hormones may not be significant
490 correlates of LOC eating. However, 2 studies reported altered postprandial CCK (amplified⁷⁰ or
491 blunted⁹⁷) and PYY responses (amplified⁷⁰) in those with BED or disinhibition compared to
492 unaffected controls, suggesting that the alterations may be relevant to LOC eating. The blunted
493 CCK increase post-meal has also been observed in the BN population¹¹⁰⁻¹¹², and it is speculated
494 that the blunted response may play a role in the diminished satiety observed in BN and contribute
495 to the initiation, perpetuation, or relapse of this eating disorder. The finding that individuals with
496 BED had amplified increases of postprandial CCK and PYY as reported in 1 study⁷⁰ was not
497 replicated anywhere else, although the interpretation that the augmented responses reflect an
498 effort to prevent binge initiation is plausible. Given that current studies produced inconsistent
499 findings of whether and in which direction the postprandial CCK and PYY responses were
500 altered, further investigations are needed on this topic.

501 All of the observational studies did not support fasting or postprandial GLP-1 as a
502 significant indicator of subclinical binge-eating, BED, uncontrolled eating, and food addiction.
503 However, the interventional drug trials utilizing GLP-1 analog have demonstrated effectiveness
504 in reducing binge-eating and uncontrolled eating, suggesting there is potential GLP-1
505 involvement in LOC eating. Several studies conducted in the BN population have established a
506 rationale to consider GLP-1 as an indicator of LOC eating. With that said, these studies have
507 noted that among patients with BN, GLP-1 was positively associated with bingeing behaviors¹¹³

508 but not purging behaviors¹¹⁴, implying that GLP-1 may be a unique indicator of binge-eating or
509 LOC eating.

510 The leptin level was found to be higher in patients with LOC eating, subclinical binge-
511 eating, BED, disinhibition, or uncontrolled eating compared with unaffected controls, albeit a
512 few exceptions. Since participants included in this review were all overweight or obese, the
513 higher levels of leptin suggest a specific link between this adipokine and LOC eating that is not
514 simply explained by the enhanced fat stores or leptin resistance associated with extra weight.
515 Furthermore, the study that reported a strong association between leptin and LOC eating after
516 adjusting for adiposity provides direct evidence to consider leptin as a significant indicator of
517 LOC eating. Besides, studies conducted in the BN population have also detected higher leptin
518 levels¹¹⁵ and a positive association between leptin and bingeing behaviors^{75,116} among patients
519 with BN, which adds support to the possibility that higher leptin levels may play a role in the
520 development or maintenance of LOC eating.

521 Adiponectin is less investigated than leptin, and the few studies consistently reported
522 lower levels of adiponectin in individuals with BED, disinhibition, or food addiction compared
523 to unaffected controls. As mentioned, the effect of adiponectin on eating regulation is glucose-
524 dependent. However, one limitation of these few studies is that they did not consider glucose
525 levels, which preclude clear conclusions about the role that adiponectin plays in these eating
526 behaviors. Despite this limitation, the lower levels of adiponectin are synchronized with
527 the behavioral manifestations of LOC eating. For example, individuals with LOC eating show a
528 trend of eating faster and consuming more high-energy foods, both of which are negatively
529 associated with adiponectin levels¹¹⁷⁻¹¹⁹.

530 The inflammation markers (hsCRP, ESR) have been found to be elevated in adolescents
531 with LOC eating and adults with BED relative to unaffected controls, indicating that LOC eating
532 may be associated with an elevated inflammatory status. Additional support for this association
533 comes from the Avon Longitudinal Study of Parents and Children (ALSPAC), which followed
534 3480 nationally representative children from birth to 18 years of age. Using the ALSPAC data,
535 one study found that children with higher levels of IL-6 and CRP had greater odds of binge-
536 eating in adolescence, although these associations were weak¹²⁰. It is worth mentioning that one
537 study included in this review reported opposite findings, in which pro-inflammatory cytokine
538 (TNF- α) decreased in individuals with food addiction. Beyond differences in sample
539 characteristics and methodologies, the discrepancy may be due to the complex interaction
540 between inflammation and gut hormones and adipokines. For example, research has shown that
541 the gut hormones (e.g., ghrelin, CCK) and adiponectin suppress the production of pro-
542 inflammatory cytokines such as TNF- α , IL-6 and IL-1 β ^{121,122}. In contrast, leptin acts in
543 opposition facilitating the secretion of these cytokines¹²³. Therefore, without evaluating these
544 interactions, it is difficult to conclude on the relationships between the inflammatory
545 cytokines/markers and LOC eating.

546 Regardless of biomarker types, 4 common limitations were identified across all of the
547 reviewed studies: studies that directly examine LOC eating are limited in the general population
548 and are absent in bariatric population; there is an insufficient number of longitudinal studies;
549 there is a lack of examination of potential mediators and moderators; and lastly, an
550 underrepresentation of males.

551 This review only identified 2 studies that directly examined LOC eating^{78,83} and 6 studies
552 that examined subclinical binge-eating^{72-74,84,90,92}, all of which were conducted among non-

553 bariatric children and adolescents. Although all other eating behaviors (e.g., BED, disinhibition)
554 chosen for review share the core feature of “loss of control”, they are broader constructs that
555 embrace other disordered eating behaviors. For example, BED additionally includes overeating,
556 and disinhibition overlaps with emotional eating and external eating. Even for the studies that
557 examined LOC eating or subclinical binge-eating, measurement limitations exist—they involved
558 only a dichotomous assessment, which failed to reflect current evidence that LOC eating should
559 be studied on a continuum of severity^{124,125}. As LOC eating has been increasingly acknowledged
560 as an independent construct³ and two specific, continuous scales (Eating Loss of Control Scale,
561 Loss of Control over Eating Scale) have been developed and validated in individuals with
562 obesity¹²⁵⁻¹²⁸, future research is needed that makes LOC eating a more explicit focus.
563 Additionally, studies on bariatric patients are warranted because LOC eating is of particular
564 importance for this population.

565 A few of the studies included in this review longitudinally examined the associations of
566 gut hormones and leptin with BED, disinhibition, or uncontrolled eating following CBT^{70,71},
567 behavioral weight loss intervention^{88,94,95}, or surgical weight loss intervention^{86,87}. However,
568 these studies did not include patients at various eating behavior stages throughout the follow-up,
569 including new-onset, maintenance, and remission. Consequently, it is impossible to conclude
570 whether the biomarker alterations occur first or after these disordered eating behaviors, and
571 whether the alterations represent state or trait markers. Bariatric surgery, in which LOC eating
572 has particular clinical relevance, offers a unique opportunity to clarify if the biomarker
573 alterations cause or are secondary to LOC eating. From before surgery, four longitudinal patterns
574 of LOC eating have been observed at 6-12 months after surgery (proportion of patients in each
575 pattern): new-onset (17-40%), maintenance (25-38%), remission (27-60%), and LOC eating-free

576 (40-70%)^{10,129,130}. Meanwhile, gut hormones, adipokines, and inflammatory cytokines/markers
577 also change as a result of surgery, although the direction and degree of change depends on the
578 surgical techniques. Taking sleeve gastrectomy as an example, extensive studies among adults
579 and adolescents have shown that fasting and postprandial ghrelin (total and acylated)¹³¹⁻¹³³,
580 leptin^{133,134}, and pro-inflammatory cytokines/markers (e.g., IL-1 α , IL-1 β , IL-6, TNF- α , CRP,
581 ESR) decrease, while adiponectin and anti-inflammatory cytokines increase after surgery¹³⁴. The
582 different patterns of LOC eating and the parallel changes of biomarkers offers great value to
583 delineation of the role of these biomarkers in LOC eating.

584 It has been widely acknowledged that beyond regulating homeostatic and hedonic food
585 consumption, gut hormones, adipokines, and pro- and anti-inflammatory cytokines are closely
586 tied to an individual's psychosocial functioning. For example, human and animal studies indicate
587 that ghrelin, GLP-1, and leptin have anti-depressant effects^{32,34,35,135}, while levels of CCK and
588 pro-inflammatory cytokines increase with elevated anxiety and depressive symptoms^{136,137}.
589 Furthermore, postprandial CCK and PYY are blunted^{138,139} but leptin levels are increased^{140,141} in
590 individuals with dietary restraint. The intrinsic link between biomarkers and psychosocial
591 functioning highlights the importance to examine the indirect associations (through the
592 moderating or mediating effect of psychosocial functioning) between biomarkers and LOC
593 eating. However, among the included studies, only two studies examined dietary restraint as a
594 moderator of relationship between postprandial CCK and disinhibition⁹⁷, or as a mediation of the
595 relationship between leptin and LOC eating⁷⁸, signaling an area that needs further research.

596 As commonly seen in eating disorder research, most studies included in this review
597 consisted of female-only or female-dominant samples. Unlike BED that has a higher prevalence
598 in women than men, the distribution of LOC eating is far less skewed, with reports of

599 comparable prevalence between males and females both in the non-bariatric¹⁴² and bariatric
600 population¹⁴³. Furthermore, given the knowledge that reproductive hormones (e.g., estradiol,
601 progesterone) and females' menstrual cycle influence the symptoms of binge-eating¹⁴⁴, the
602 relationship between biomarkers and LOC eating may differ for males and females. Therefore,
603 future studies should increase the presentation of males and examine the gender differences
604 when investigating biomarkers with LOC eating.

605 This scoping review has several limitations. First, because studies that directly report
606 LOC eating are too scarce in number, several overlapping but broader eating behaviors were also
607 included. The inclusion of these eating behaviors is necessary to provide richer and more
608 comprehensive information, but it introduces heterogeneity and creates difficulty to isolate the
609 construct of LOC eating. Second, only published full articles were searched; therefore, relevant
610 information presented in the gray literature (e.g., unpublished manuscripts) may have been
611 missed. Third, the literature search was limited to 3 databases. Additional literature may exist
612 that was not included within the scope of this search. Fourth, this review only focused on the
613 significance testing results when reporting the relationships between biomarkers and LOC eating.
614 The lack of an examination of the effect sizes precludes the understanding of the strength of
615 these relationships. Given that the sample size in most of the included studies was small and this
616 scoping review provides initial evidence that several gut hormones (e.g., postprandial ghrelin),
617 adipokines (e.g., leptin), and pro-inflammatory markers (e.g., hsCRP) may be related to LOC
618 eating, future studies would benefit from conducting a meta-analysis and pooling the effect sizes
619 across studies.

620 **5. Conclusion**

621 Despite the limited number of studies and conflicting results, there is evidence that
622 supports the associations of lower levels of fasting acyl ghrelin and adiponectin, higher levels of
623 leptin, hsCRP, and ESR, and altered responses of postprandial ghrelin (blunted), CCK (blunted
624 or amplified), and PYY (amplified) to meal ingestion with the eating behaviors including LOC
625 eating, subclinical binge-eating, BED, disinhibition, uncontrolled eating, and food addiction.
626 Additionally, using GLP-1 analog may reduce binge-eating. Future studies would benefit from a
627 direct examination of LOC eating especially in the bariatric patients, a greater focus on
628 longitudinal studies, an in-depth exploration of the interactions between biomarkers and
629 psychosocial functioning, and an investigation of gender differences that may shape the
630 relationship between biomarkers and LOC eating. Other considerations include examining acyl
631 ghrelin instead of total ghrelin, reporting glucose levels with adiponectin, and investigating the
632 interactions among inflammation, gut hormones, and adipokines.

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635 **Author Contributions**

636 Y.Y. designed the study, searched, screened, reviewed, and synthesized the studies, drafted the
637 manuscript. I.F. and M.Y. critically reviewed and edited the manuscript. W.Z. verified the
638 included studies and reviewed the manuscript. S.G. supervised the review process and critically
639 reviewed the manuscript. All authors have read, reviewed and approved the manuscript and hold
640 the responsibility for its final content.

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644 **Declarations of Interest**

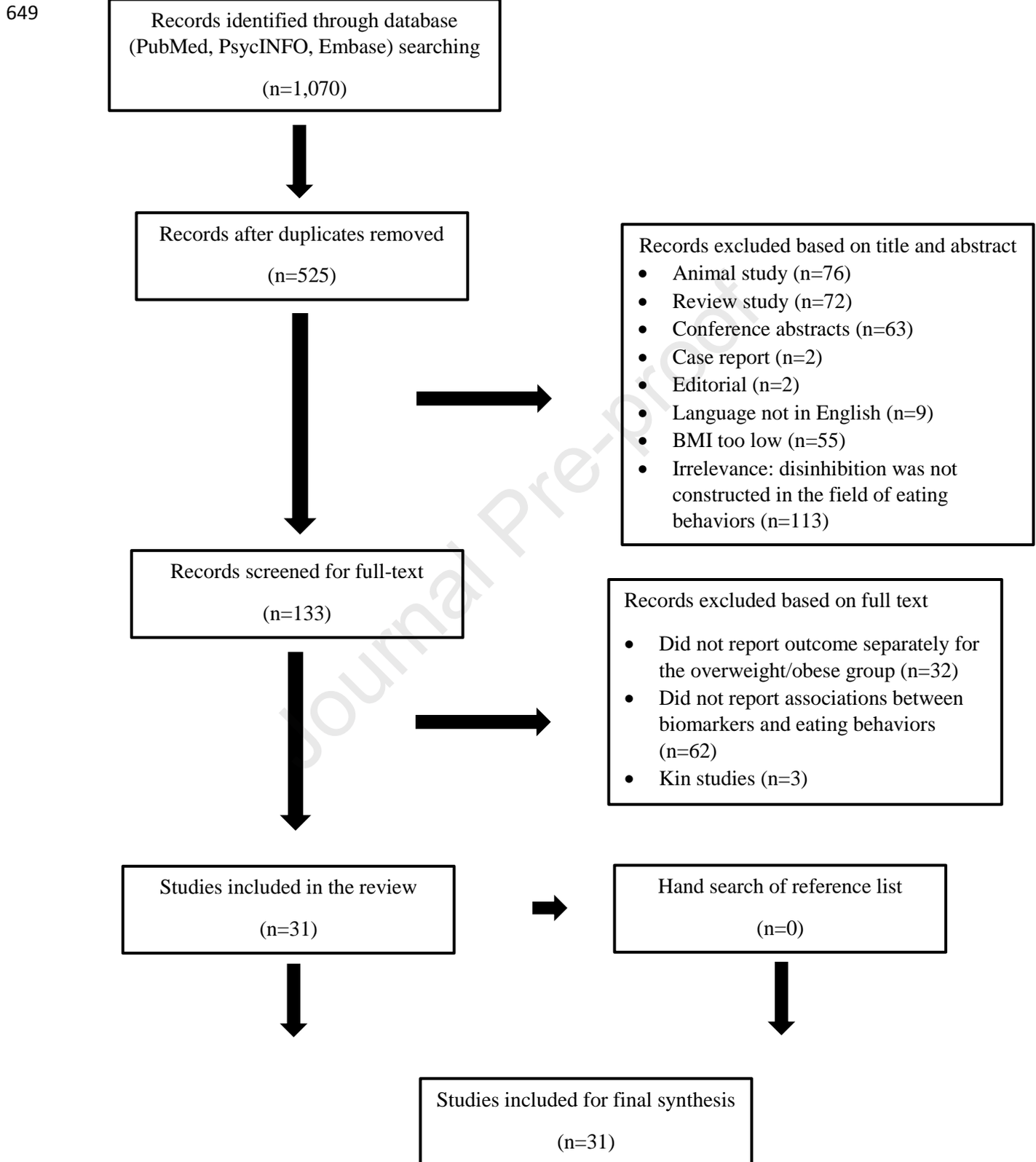
645 None.

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648 Figure 1 Study Selection Flow



650 Table 1 Description of included studies (N=31)

Study characteristics	Sample description	Biomarker	Eating behavior	Conclusion
Ghrelin				
<p>Circulating ghrelin is decreased in non-obese and obese women with binge eating disorder as well as in obese non-binge eating women, but not in patients with bulimia nervosa⁶⁸</p> <ul style="list-style-type: none"> Monteleone et al. 2004 Italy Case-control 	<ul style="list-style-type: none"> 56 women with BN: age: 23.4±4.3; BMI: 21.9±3.8 34 women with BED and obesity: age: 33.6±9.1; BMI: 39.8±4.9; binge frequency: 10.9±8.1 episodes/week 13 women with BED but without obesity: age: 26.9±8.0; BMI: 25.8±2.5; binge frequency: 9.5±6.5 episodes/week 28 women with obesity but without BED: age: 38.3±14.1; BMI: 38.1±6.3 51 healthy control women: age: 22.6±3.1; BMI: 21.7±2.3 Recruited from the Eating Disorder Center None of women with BED had a history of AN or BN, and they are free from drugs for at least 8 weeks 	<ul style="list-style-type: none"> Fasting ghrelin Covariates included BMI and body mass fat 	<ul style="list-style-type: none"> BED assessed by Structured Clinical Interview for DSM-IV 	<ul style="list-style-type: none"> Ghrelin in patients with BED and obesity did not significantly differ from ghrelin in non-BED patients with obesity Ghrelin was not significantly correlated to the binge episodes among women with BED
<p>Impact of laparoscopic adjustable gastric banding on plasma ghrelin, eating behaviour and body weight⁸⁶</p> <ul style="list-style-type: none"> Schindler et al 2004 Australia Longitudinal 	<ul style="list-style-type: none"> 23 candidates for bariatric surgery (gastric banding): age: 35.6±2.12; BMI: 44.8±1.0; female: 86.9% Recruited from Department of Surgery Exclusion: type 2 diabetes, myocardial infarction, any malignancy, chronic kidney or liver disease, seizure, obesity caused by an endocrine disorder, psychiatric disorders, current pregnancy or breastfeeding. 	<ul style="list-style-type: none"> Fasting ghrelin Assessment timepoints: before and 6 months after surgery 	<ul style="list-style-type: none"> Disinhibition subscale of Three Factor Eating Questionnaire 	<ul style="list-style-type: none"> Before surgery, ghrelin was not correlated with disinhibition After surgery, the change of ghrelin was not correlated with the change of disinhibition
<p>The acute ghrelin response to a psychological stress challenge does not predict the post-stress urge to eat⁸¹</p> <ul style="list-style-type: none"> Rouach et al 	<ul style="list-style-type: none"> 8 adults with BED: age: 49.7±4.8; BMI: 36.7±1.5; female: 75% 8 non-BED adults: age: 50.2±5.7; BMI: 35.2±1.4; female: 62.5% 8 adults with normal weight: age: 32.6±3.5; BMI: 21.2±0.7; female: 62.5% Recruited from an obesity clinic 	<ul style="list-style-type: none"> Postprandial ghrelin Subjects were instructed to eat a light breakfast around 7am, and tests began at 9 am Covariates included age, gender and BMI 	<ul style="list-style-type: none"> BED measure not specified 	<ul style="list-style-type: none"> Ghrelin levels in patients with BED were higher than healthy obese subjects, but this difference did not attain statistical significance.

<ul style="list-style-type: none"> • 2007 • Italy • Case-control 	<ul style="list-style-type: none"> • Exclusion: anorexia or bulimia nervosa, any psychiatric co-morbidities, taking medication affecting the central nervous system, receiving medication known to interfere with cortisol measurements 			
<p>Appetite-related gut peptides, ghrelin, PYY, and GLP-1 in obese women with and without binge eating disorder (BED)⁷¹</p> <ul style="list-style-type: none"> • Geliebter et al. • 2008 • US • Longitudinal 	<ul style="list-style-type: none"> • 10 women with BED: age: 29.9±8.2; BMI: 36.5±6.6; frequency of binge eating days: 3.2±0.8d • 9 women without BED: age: 30.3±8.4; BMI: 35.8±5.5 • All women were premenopausal and in otherwise good health • Eight BED and 8 non-BED participants were randomly assigned for 6 weeks to either (a) individual weekly treatment with nutritional counseling and cognitive behavior therapy or (b) a non-treatment control 	<ul style="list-style-type: none"> • Fasting and postprandial ghrelin • Test meal: 1254 kJ liquid meal • Blood draw: -15, and 0, 5, 15, 30, 60, 90, and 120 min after test meal • Covariates included weight change 	<ul style="list-style-type: none"> • BED assessed by Questionnaire on Eating and Weight Patterns followed by a clinical interview 	<ul style="list-style-type: none"> • Fasting ghrelin was lower and declined less postprandially in the BED group. • After intervention, ghrelin normalized and there were no differences in fasting and postprandial ghrelin between groups. However, the changes in ghrelin pre to post intervention were not related to treatment condition or change in binge eating days.
<p>Lifestyle intervention discloses an association of the Eating Inventory-51 factors with cardiometabolic health risks⁹⁴</p> <ul style="list-style-type: none"> • Hainerova et al • 2013 • Czech Republic • Longitudinal 	<ul style="list-style-type: none"> • 67 women with obesity: age: 48.7±12.2; BMI: 32.4±4.4 • Women participated in a 4-week lifestyle obesity management (e.g., low-calorie diet, physical activity, and cognitive behavioral therapy) 	<ul style="list-style-type: none"> • Fasting ghrelin • Assessment timepoints: before and after a 3-week weight management • Covariates not specified 	<ul style="list-style-type: none"> • Disinhibition subscale of Eating Inventory 	<ul style="list-style-type: none"> • No significant associations between fasting ghrelin and disinhibition at baseline
<p>Relationships among tonic and episodic aspects of motivation to eat, gut peptides, and</p>	<ul style="list-style-type: none"> • 12 candidates for bariatric surgery (gastric bypass): age: 36±2; BMI: 45.3±1.9; female: 75% • Exclusion criteria not specified 	<ul style="list-style-type: none"> • Fasting and postprandial ghrelin • Test meal: liquid meal (200 ml, 300 kcal) 	<ul style="list-style-type: none"> • Uncontrolled eating subscale of the Three Factor Eating 	<ul style="list-style-type: none"> • Fasting and postprandial ghrelin was not associated with uncontrolled eating at

weight before and after bariatric surgery ⁸⁷ <ul style="list-style-type: none"> • Bryant et al • 2013 • UK • Longitudinal 		<ul style="list-style-type: none"> • Blood draw: -10, and 0, 10, 20, 30, 60, 90, 120, and 180 minutes after consumption • Assessment timepoints: pre-surgery, and 2 months, and 1 year post-surgery • Covariates included BMI 	Questionnaire	before and 12 months following surgery
Ghrelin and peptide YY increase with weight loss during a 12-month intervention to reduce dietary energy density in obese women ⁹⁵ <ul style="list-style-type: none"> • Hill et al. • 2013 • US • Longitudinal 	<ul style="list-style-type: none"> • 71 women with obesity: age: 46.7±1.0; BMI: 33.3±0.3 • Participants were attending a 12-month intervention focused on reducing energy density • Inclusion: women ages 20 – 60 years, BMI between 30–40 kg/m². • Exclusion: high blood pressure, serum triacylglycerols, and total cholesterol, major medical conditions, pregnancy/lactation, taking selective serotonin reuptake inhibitors, symptoms of depression or disordered eating, currently participating in a weight loss program. 	<ul style="list-style-type: none"> • Fasting ghrelin • Assessment timepoints: 0, 3, 6 and 12 months after intervention • Covariates included energy density, body weight, hunger, dietary restraint 	<ul style="list-style-type: none"> • Disinhibition subscale of Eating Inventory 	<ul style="list-style-type: none"> • There was no significant longitudinal association between ghrelin and disinhibition
Hormonal and dietary characteristics in obese human subjects with and without food addiction ⁶⁹ <ul style="list-style-type: none"> • Pedram et al • 2014 • Canada • Case-control 	<ul style="list-style-type: none"> • 29 adults with food addiction: age: 42.5±9.6; BMI: 32.5±6; female: 82.7% • 29 non-food-addicted subjects matched for age, sex, BMI and physical activity: age: 42±8.9; BMI: 32±4.4; female: 82.7% • Recruited from non-clinical setting • Inclusion: age >19 years, without serious metabolic, cardiovascular or endocrine diseases, and not pregnant at the time of the study. 	<ul style="list-style-type: none"> • Fasting ghrelin • Covariates not specified 	<ul style="list-style-type: none"> • Food addiction assessed by Yale Food Addiction Scale 	<ul style="list-style-type: none"> • Ghrelin did not differ between the two groups
Appetite sensations, appetite signaling proteins, and glucose in obese adolescents with subclinical binge eating	<ul style="list-style-type: none"> • 6 adolescents with binge eating: age: 13.7±0.7; BMI: 35.9±2.2; female: 33.3% • 9 adolescents without binge eating: age: 14.5±0.8; BMI: 38.5±3.0; female: 83.3% 	<ul style="list-style-type: none"> • Fasting and postprandial ghrelin • Test meal: 571 kcal standardized breakfast (50%) 	<ul style="list-style-type: none"> • Binge eating assessed by Eating Disorder Diagnostic Scale 	<ul style="list-style-type: none"> • Overall concentrations of ghrelin across the monitoring period did not significantly differ between the two groups

disorder ⁸⁴ <ul style="list-style-type: none"> • Adamo et al • 2014 • Canada • Case-control 	<ul style="list-style-type: none"> • Recruited from pediatric endocrinology clinic • Exclusion: had type 2 diabetes or were taking medications that could influence body composition or appetite. 	carbohydrate, 35% fat, and 15% protein) <ul style="list-style-type: none"> • Fasting, and postprandially at 15, 30, 60, 90, 120, and 240 minutes • Covariates included sex and fat mass 		<ul style="list-style-type: none"> • Hunger and satiety were not significantly correlated with ghrelin
Plasma ghrelin levels and weight regain after Roux-en-Y Gastric Bypass Surgery ⁸⁵ <ul style="list-style-type: none"> • Abu Dayyeh et al • 2017 • US • Cross-sectional 	<ul style="list-style-type: none"> • 36 patients who have undergone bariatric surgery (gastric bypass): age: 47±10; BMI: 38±7.7; female: 94%; length post-surgery: 5±4 years; race: (white: 68%, black: 18%, Hispanic: 14%) • Inclusion: at least 1 year post-surgery • Exclusion: gastrogastic fistula(e) 	<ul style="list-style-type: none"> • Fasting ghrelin 	<ul style="list-style-type: none"> • Uncontrolled subscale of Three Factor Eating Questionnaire 	<ul style="list-style-type: none"> • There was no association between ghrelin and uncontrolled eating
CCK, ghrelin, and PYY responses in individuals with binge eating disorder before and after a cognitive behavioral treatment (CBT) ⁷⁰ <ul style="list-style-type: none"> • Munsch et al. • 2019 • Switzerland • Longitudinal 	<ul style="list-style-type: none"> • 18 adults with BED: age: 50.2±9.5; BMI: 32.4±5.4; female: 77% • 20 age- and BMI-matched healthy controls: age: 48.6±9.7; BMI: 34.3±7.6; female: 95% • Recruited from non-clinical setting. • Inclusion: between 18 and 70 years old, BMI between 27 and 40, and meet full criteria for BED. • Exclusion: unstable medical conditions, mental disorders warranting immediate treatment, pregnancy, participation weight loss treatment • Patients with BED received a 8-week CBT 	<ul style="list-style-type: none"> • Fasting and postprandial ghrelin • Test meal: 797 kcal standard breakfast (62% carbohydrate, 10% protein, and 28% fat) • -20 min, -5 min, and 15, 30, 45, 60, 90,120, and 180 min after the meal • Covariates included baseline BMI and sex 	<ul style="list-style-type: none"> • BED assessed by Eating Disorder Examination 	<ul style="list-style-type: none"> • Before and short-term after intervention, fasting ghrelin did not differ between groups • Before intervention, postprandial ghrelin levels did not differ between BED participants and controls • In the BED group, the fasting and postprandial ghrelin did not change before and after intervention • Ghrelin were not related to objective binge eating post-intervention
Meal-related acyl and des-acyl ghrelin and other appetite-related	<ul style="list-style-type: none"> • 20 adults with binge eating (10 BED and 10 subthreshold binge eating): age: 37.8±8.4; BMI: 36.2±5.5; female: 60% 	<ul style="list-style-type: none"> • Fasting and postprandial ghrelin (des-acyl ghrelin, acyl ghrelin 	<ul style="list-style-type: none"> • Binge eating assessed by Eating Disorder 	<ul style="list-style-type: none"> • Significantly lower acyl ghrelin concentrations for the binge eating group

<p>hormones in people with obesity and binge eating⁷²</p> <ul style="list-style-type: none"> Hernandez et al 2019 US Case-control 	<ul style="list-style-type: none"> 22 adults without binge eating: age: 34.0±8.2; BMI: 36.4±4.9; female: 31.8% Race in the total group: Black/African American (61%), Hispanic (19.5%), non-Hispanic White (14.6%), and other (4.8%). Recruited from non-clinical settings Exclusion: individuals with other eating disorders 	<ul style="list-style-type: none"> Test meal: 1254 KJ liquid meal (55% carbohydrate, 24% protein, 21% fat) Blood draw: -15, and 0, 10, 30, and 60 minutes after meal The two groups did not differ in age, BMI, and sex 	<p>Examination and the Questionnaire on Eating and Weight Patterns</p>	<p>compared with the non-binge eating group</p> <ul style="list-style-type: none"> Decrease of postprandial AG was significantly smaller for the binge eating group at 30 and 60 minutes than the decreases in the non-binge eating group Decrease of postprandial des-acyl ghrelin did not significantly differ between groups
<p>Association between des-acyl ghrelin at fasting and predictive index of muscle derangement, metabolic markers and eating disorders: a cross-sectional study in overweight and obese adults⁹²</p> <ul style="list-style-type: none"> Perna et al. 2020 Italy Cross-sectional 	<ul style="list-style-type: none"> 88 adults: age: 43.3±9.3; BMI: 30.2±3.3; female: 72.7%; Exclusion: hepatic or renal disease, diabetes, cardiovascular disease, hypertension, diagnosed bulimia, cancer, surgery for weight loss, weight loss medication, depressive disorder, female patients were excluded if they were pregnant or lactating or had entered menopause 	<ul style="list-style-type: none"> Fasting ghrelin (des-acyl ghrelin, acyl ghrelin) Covariates not specified 	<ul style="list-style-type: none"> Binge eating behavior measured by Binge Eating Scale 	<ul style="list-style-type: none"> Acyl ghrelin was negatively associated with binge eating, while des-acyl ghrelin was not significantly associated with binge eating
CCK				
<p>Gastric capacity, test meal intake, and appetitive hormones in binge eating disorder⁷³</p> <ul style="list-style-type: none"> Geliebter et al. 2004 US Case-control 	<ul style="list-style-type: none"> 11 women with BED: age: 29±8; BMI: 36.6±6.2 13 women with subclinical binge eating: age: 29±7; BMI: 35.8±5.5 13 non-binge eating women: age: 32±9; BMI: 35.1±5.3 Inclusion: premenopausal, nonsmokers, not taking illegal drugs or medications affecting 	<ul style="list-style-type: none"> Fasting and postprandial CCK Test meal: 600 ml liquid meal (with an energy density of 4.2 J/g) Blood draw: -15, and 0, 5, 15, 30, 60, 90, and 120 min after meal 	<ul style="list-style-type: none"> Binge eating assessed by Questionnaire on Eating and Weight Patterns and clinical interview 	<ul style="list-style-type: none"> Fasting and postprandial CCK did not differ between groups

	weight, and weight stable within past 3 months.	<ul style="list-style-type: none"> • Covariates not specified 		
<p>Glycemic index, cholecystokinin, satiety and disinhibition: is there an unappreciated paradox for overweight women?⁹⁷</p> <ul style="list-style-type: none"> • Burton-Freeman et al. • 2008 • US • Randomized cross-over 	<ul style="list-style-type: none"> • 21 women with overweight: age: 31±8; BMI: 27±1 • Recruited from non-clinical setting • Exclusion: cardiovascular or metabolic disorders, pregnant or had been pregnant within 18 months before the study, were taking medications to manage body weight or control appetite. 	<ul style="list-style-type: none"> • Postprandial CCK • Test meal: breakfast with high glycemic index (54.2% carbohydrate; 15% protein; 30.8% fat); breakfast with low glycemic index (54.6% carbohydrate; 14.7% protein; 30.7% fat) • Fasting, and 30, 60, 90, 120, 150, 210, 270, 360 and 480 min following the breakfast • Women were instructed to maintain a stable weight 	<ul style="list-style-type: none"> • Disinhibition assessed by Three Factor Eating Questionnaire 	<ul style="list-style-type: none"> • Among participants with dietary restraint, those with higher disinhibition scores had a blunted CCK response to both high glycemic index and low glycemic index meals
<p>CCK, ghrelin, and PYY responses in individuals with binge eating disorder before and after a cognitive behavioral treatment (CBT)⁷⁰</p> <ul style="list-style-type: none"> • Munsch et al. • 2019 • Switzerland • Longitudinal 	<ul style="list-style-type: none"> • 18 adults with BED: age: 50.2±9.5; BMI: 32.4±5.4; female: 77% • 20 age- and BMI-matched healthy controls: age: 48.6±9.7; BMI: 34.3±7.6; female: 95% • Recruited from non-clinical setting. • Inclusion: between 18 and 70 years old, BMI between 27 and 40, and meet full criteria for BED. • Exclusion: unstable medical conditions, mental disorders warranting immediate treatment, pregnancy, participation weight loss treatment • Patients with BED received a 8-week CBT 	<ul style="list-style-type: none"> • Fasting and postprandial CCK • Test meal: 797 kcal standard breakfast (62% carbohydrate, 10% protein, and 28% fat) • -20 min, -5 min, and 15, 30, 45, 60, 90,120, and 180 min after the meal • Covariates included baseline BMI and sex 	<ul style="list-style-type: none"> • BED assessed by Eating Disorder Examination 	<ul style="list-style-type: none"> • Before and short-term after intervention, fasting CCK did not differ between groups • Before intervention, BED participants revealed a higher meal-induced increase and stronger decline thereafter in CCK compared to controls • In the BED group, the fasting and postprandial CCK did not change before and after intervention • CCK was not related to objective binge eating post-intervention
<p>Meal-related acyl and des-acyl ghrelin and other appetite-related</p>	<ul style="list-style-type: none"> • 20 adults with binge eating (10 BED and 10 subthreshold binge eating): age: 37.8±8.4; BMI: 36.2±5.5; female: 60% 	<ul style="list-style-type: none"> • Fasting and postprandial CCK • Test meal: 1254 KJ liquid 	<ul style="list-style-type: none"> • Binge eating assessed by Eating Disorder 	<ul style="list-style-type: none"> • CCK did not differ between groups

<p>hormones in people with obesity and binge eating⁷²</p> <ul style="list-style-type: none"> Hernandez et al 2019 US Case-control 	<ul style="list-style-type: none"> 22 adults without binge eating: age: 34.0±8.2; BMI: 36.4±4.9; female: 31.8% Race in the total group: Black/African American (61%), Hispanic (19.5%), non-Hispanic White (14.6%), and other (4.8%). Recruited from non-clinical settings Exclusion: individuals with other eating disorders 	<p>meal (55% carbohydrate, 24% protein, 21% fat)</p> <ul style="list-style-type: none"> Blood draw: -15, and 0, 10, 30, and 60 minutes after meal The two groups did not differ in age, BMI, and sex 	<p>Examination and the Questionnaire on Eating and Weight Patterns</p>	
PYY				
<p>Appetite-related gut peptides, ghrelin, PYY, and GLP-1 in obese women with and without binge eating disorder (BED)⁷¹</p> <ul style="list-style-type: none"> Geliebter et al. 2008 US Longitudinal 	<ul style="list-style-type: none"> 10 women with BED: age: 29.9±8.2; BMI: 36.5±6.6; frequency of binge eating days: 3.2±0.8d 9 women without BED: age: 30.3±8.4; BMI: 35.8±5.5 All women were premenopausal and in otherwise good health Eight BED and 8 non-BED participants were randomly assigned for 6 weeks to either (a) individual weekly treatment with nutritional counseling and cognitive behavior therapy or (b) a non-treatment control 	<ul style="list-style-type: none"> Fasting and postprandial PYY Test meal: 1254 kJ liquid meal Blood draw: -15, and 0, 5, 15, 30, 60, 90, and 120 min after test meal Covariates included weight change 	<ul style="list-style-type: none"> BED assessed by Questionnaire on Eating and Weight Patterns followed by a clinical interview 	<ul style="list-style-type: none"> Fasting and postprandial PYY did not differ between groups. Fasting and postprandial PYY did not change after intervention
<p>Ghrelin and peptide YY increase with weight loss during a 12-month intervention to reduce dietary energy density in obese women⁹⁵</p> <ul style="list-style-type: none"> Hill et al. 2013 US Longitudinal 	<ul style="list-style-type: none"> 71 women with obesity: age: 46.7±1.0; BMI: 33.3±0.3 Participants were attending a 12-month intervention focused on reducing energy density Inclusion: women ages 20 – 60 years, BMI between 30–40 kg/m². Exclusion: high blood pressure, serum triacylglycerols, and total cholesterol, major medical conditions, pregnancy/lactation, taking selective serotonin reuptake inhibitors, symptoms of depression or disordered eating, currently participating in a weight loss program. 	<ul style="list-style-type: none"> Fasting PYY Assessment timepoints: 0, 3, 6 and 12 months after intervention Covariates included energy density, body weight, hunger, dietary restraint 	<ul style="list-style-type: none"> Disinhibition subscale of Eating Inventory 	<ul style="list-style-type: none"> There was no significant longitudinal association between PYY and disinhibition

<p>Lifestyle intervention discloses an association of the Eating Inventory-51 factors with cardiometabolic health risks⁹⁴</p> <ul style="list-style-type: none"> • Hainerova´ et al • 2013 • Czech Republic • Longitudinal 	<ul style="list-style-type: none"> • 67 women with obesity: age: 48.7±12.2; BMI: 32.4±4.4 • Women participated in a 4-week lifestyle obesity management (e.g., low-calorie diet, physical activity, and cognitive behavioral therapy) 	<ul style="list-style-type: none"> • Fasting PYY • Assessment timepoints: before and after a 3-week weight management • Covariates not specified 	<ul style="list-style-type: none"> • Disinhibition subscale of Eating Inventory 	<ul style="list-style-type: none"> • No significant associations between PYY and disinhibition at baseline
<p>Hormonal and dietary characteristics in obese human subjects with and without food addiction⁶⁹</p> <ul style="list-style-type: none"> • Pedram et al • 2014 • Canada • Case-control 	<ul style="list-style-type: none"> • 29 adults with food addiction: age: 42.5±9.6; BMI: 32.5±6; female: 82.7% • 29 non-food-addicted subjects matched for age, sex, BMI and physical activity: age: 42±8.9; BMI: 32±4.4; female: 82.7% • Recruited from non-clinical setting • Inclusion: age >19 years, without serious metabolic, cardiovascular or endocrine diseases, and not pregnant at the time of the study. 	<ul style="list-style-type: none"> • Fasting PYY • Covariates not specified 	<ul style="list-style-type: none"> • Food addiction assessed by Yale Food Addiction Scale 	<ul style="list-style-type: none"> • The 2 groups did not differ in terms of PYY
<p>Appetite sensations, appetite signaling proteins, and glucose in obese adolescents with subclinical binge eating disorder⁸⁴</p> <ul style="list-style-type: none"> • Adamo et al • 2014 • Canada • Case-control 	<ul style="list-style-type: none"> • 6 adolescents with binge eating: age: 13.7±0.7; BMI: 35.9±2.2; female: 33.3% • 9 adolescents without binge eating: age: 14.5±0.8; BMI: 38.5±3.0; female: 83.3% • Recruited from pediatric endocrinology clinic • Exclusion: had type 2 diabetes or were taking medications that could influence body composition or appetite. 	<ul style="list-style-type: none"> • Fasting and postprandial PYY • Test meal: 571 kcal standardized breakfast (50% carbohydrate, 35% fat, and 15% protein) • Fasting, and postprandially at 15, 30, 60, 90, 120, and 240 minutes • Covariates included sex and fat mass 	<ul style="list-style-type: none"> • Binge eating assessed by Eating Disorder Diagnostic Scale 	<ul style="list-style-type: none"> • Overall concentrations of PYY across the monitoring period did not significantly differ between the two groups • Hunger and satiety were not significantly correlated with PYY
<p>CCK, ghrelin, and PYY responses in individuals with binge eating disorder before and after</p>	<ul style="list-style-type: none"> • 18 adults with BED: age: 50.2±9.5; BMI: 32.4±5.4; female: 77% • 20 age- and BMI-matched healthy controls: 	<ul style="list-style-type: none"> • Fasting and postprandial CCK • Test meal: 797 kcal standard 	<ul style="list-style-type: none"> • BED assessed by Eating Disorder Examination 	<ul style="list-style-type: none"> • Before and short-term after intervention, fasting PYY did not differ

<p>a cognitive behavioral treatment (CBT)⁷⁰</p> <ul style="list-style-type: none"> • Munsch et al. • 2019 • Switzerland • Longitudinal 	<p>age: 48.6±9.7; BMI: 34.3±7.6; female: 95%</p> <ul style="list-style-type: none"> • Recruited from non-clinical setting. • Inclusion: between 18 and 70 years old, BMI between 27 and 40, and meet full criteria for BED. • Exclusion: unstable medical conditions, mental disorders warranting immediate treatment, pregnancy, participation weight loss treatment • Patients with BED received a 8-week CBT 	<p>breakfast (62% carbohydrate, 10% protein, and 28% fat)</p> <ul style="list-style-type: none"> • -20 min, -5 min, and 15, 30, 45, 60, 90,120, and 180 min after the meal • Covariates included baseline BMI and sex 		<p>between groups</p> <ul style="list-style-type: none"> • Before intervention, BED participants revealed a higher meal-induced increase and stronger decline thereafter in PYY compared to controls • In the BED group, the fasting and postprandial PYY did not change before and after intervention • PYY was not related to objective binge eating post-intervention
<p>Meal-related acyl and des-acyl ghrelin and other appetite-related hormones in people with obesity and binge eating⁷²</p> <ul style="list-style-type: none"> • Hernandez et al • 2019 • US • Case-control 	<ul style="list-style-type: none"> • 20 adults with binge eating (10 BED and 10 subthreshold binge eating): age: 37.8±8.4; BMI: 36.2±5.5; female: 60% • 22 adults without binge eating: age: 34.0±8.2; BMI: 36.4±4.9; female: 31.8% • Race in the total group: Black/African American (61%), Hispanic (19.5%), non-Hispanic White (14.6%), and other (4.8%). • Recruited from non-clinical settings • Exclusion: individuals with other eating disorders 	<ul style="list-style-type: none"> • Fasting and postprandial PYY • Test meal: 1254 KJ liquid meal (55% carbohydrate, 24% protein, 21% fat) • Blood draw: -15, and 0, 10, 30, and 60 minutes after meal • The two groups did not differ in age, BMI, and sex 	<ul style="list-style-type: none"> • Binge eating assessed by Eating Disorder Examination and the Questionnaire on Eating and Weight Patterns 	<ul style="list-style-type: none"> • PYY did not differ between groups
<p>Association between eating behavior, anthropometric and biochemical measurements, and peptide YY (PYY) hormone levels in obese adolescents in outpatient care⁸⁸</p> <ul style="list-style-type: none"> • Fernandes et al. 	<ul style="list-style-type: none"> • 51 adolescents with obesity receiving outpatient treatment (nutrition and physical activity intervention): age: 12.0±0.9; BMI: 29.6±4.4; female: 56.9% • Recruited from obesity outpatient • Inclusion: at least 10 years old and had a pubertal stage greater than I • Exclusion criteria not specified 	<ul style="list-style-type: none"> • Fasting PYY • Assessment time points: before treatment, and 24 and 48 weeks after treatment • Covariates not specified 	<ul style="list-style-type: none"> • Uncontrolled eating assessed by Three Factor Eating Questionnaire 	<ul style="list-style-type: none"> • PYY was not significantly related to uncontrolled eating at any assessment point

<ul style="list-style-type: none"> • 2020 • Brazil • Longitudinal 				
GLP-1				
<p>Appetite-related gut peptides, ghrelin, PYY, and GLP-1 in obese women with and without binge eating disorder (BED)⁷¹</p> <ul style="list-style-type: none"> • Geliebter et al. • 2008 • US • Longitudinal 	<ul style="list-style-type: none"> • 10 women with BED: age: 29.9±8.2; BMI: 36.5±6.6; frequency of binge eating days: 3.2±0.8d • 9 women without BED: age: 30.3±8.4; BMI: 35.8±5.5 • All women were premenopausal and in otherwise good health • Eight BED and 8 non-BED participants were randomly assigned for 6 weeks to either (a) individual weekly treatment with nutritional counseling and cognitive behavior therapy or (b) a non-treatment control 	<ul style="list-style-type: none"> • Fasting and postprandial GLP-1 • Test meal: 1254 kJ liquid meal • Blood draw: -15, and 0, 5, 15, 30, 60, 90, and 120 min after test meal • Covariates included weight change 	<ul style="list-style-type: none"> • BED assessed by Questionnaire on Eating and Weight Patterns followed by a clinical interview 	<ul style="list-style-type: none"> • Fasting and postprandial GLP-1 did not differ between groups.
<p>Relationships among tonic and episodic aspects of motivation to eat, gut peptides, and weight before and after bariatric surgery⁸⁷</p> <ul style="list-style-type: none"> • Bryant et al • 2013 • UK • Longitudinal 	<ul style="list-style-type: none"> • 12 candidates for bariatric surgery (gastric bypass): age: 36±2; BMI: 45.3±1.9; female: 75% • Exclusion criteria not specified 	<ul style="list-style-type: none"> • Fasting and postprandial GLP-1 • Test meal: liquid meal (200 ml, 300 kcal) • Blood draw: -10, and 0, 10, 20, 30, 60, 90, 120, and 180 minutes after consumption • Assessment timepoints: pre-surgery, and 2 months, and 1 year post-surgery • Covariates included BMI 	<ul style="list-style-type: none"> • Uncontrolled eating subscale of the Three Factor Eating Questionnaire 	<ul style="list-style-type: none"> • Fasting and postprandial GLP-1 were not related to uncontrolled eating at before or after surgery
<p>Hormonal and dietary characteristics in obese human subjects with and without food addiction⁶⁹</p> <ul style="list-style-type: none"> • Pedram et al • 2014 • Canada 	<ul style="list-style-type: none"> • 29 adults with food addiction: age: 42.5±9.6; BMI: 32.5±6; female: 82.7% • 29 non-food-addicted subjects matched for age, sex, BMI and physical activity: age: 42±8.9; BMI: 32±4.4; female: 82.7% • Recruited from non-clinical setting • Inclusion: age >19 years, without serious 	<ul style="list-style-type: none"> • Fasting GLP-1 • Covariates not specified 	<ul style="list-style-type: none"> • Food addiction assessed by Yale Food Addiction Scale 	<ul style="list-style-type: none"> • The 2 groups did not differ in terms of GLP-1

<ul style="list-style-type: none"> Case-control 	<p>metabolic, cardiovascular or endocrine diseases, and not pregnant at the time of the study.</p>			
<p>Short-term intervention with liraglutide improved eating behavior in obese women with polycystic ovary syndrome¹⁰⁰</p> <ul style="list-style-type: none"> Jensterle et al 2014 Slovenia Pre-post 	<ul style="list-style-type: none"> 36 women with polycystic ovary syndrome: age: 31.2±7.8; BMI: 38.7±0.1 Recruited from outpatients Department of Endocrinology Inclusion: weight stable in the last 6 months, more than 18 years old, premenopausal, BMI >30 kg/m², had been taking metformin 2000 mg for at least 6 months Exclusion: type 1 or type 2 diabetes, history of carcinoma, cardiovascular, kidney or liver disease and the use of medications other than metformin known to affect metabolic functions within past 90 days 	<ul style="list-style-type: none"> GLP-1 analog 12-week liraglutide at a dose of 0.6 mg/day, which increased to 1.2 mg/day after 1 week 	<ul style="list-style-type: none"> Uncontrolled eating subscale of Three Factor Eating Questionnaire 	<ul style="list-style-type: none"> After treatment with liraglutide, uncontrolled eating significantly decreased
<p>Improvement in binge eating in non-diabetic obese individuals after 3 months of treatment with liraglutide — A pilot study⁹⁰</p> <ul style="list-style-type: none"> Robert et al 2015 Malaysia RCT 	<ul style="list-style-type: none"> 42 adults with binge eating were randomly assigned to the liraglutide or control group 21 adults in the liraglutide group: BMI: 36.1±3.8 21 in the control group: BMI: 35.7±4.5 Total group: age: 34±9 Exclusion: a history of taking medications that may affect weight and appetite, contraindications to liraglutide, and any chronic illnesses such as diabetes mellitus and cardiovascular diseases 	<ul style="list-style-type: none"> GLP-1 analog 12-week liraglutide treatment 	<ul style="list-style-type: none"> Binge eating assessed by Binge Eating Scale 	<ul style="list-style-type: none"> Participants who received liraglutide had significant reductions in binge eating at 12 weeks 81% of those receiving liraglutide had binge eating remission
<p>Dulaglutide reduces binge episodes in type 2 diabetic patients with binge eating disorder: A pilot study⁹¹</p> <ul style="list-style-type: none"> Porto et al 2020 Italy 	<ul style="list-style-type: none"> 60 adults with BED and diabetes were randomly assigned to the liraglutide or gliclazide group 30 adults in the dulaglutide group: age: 54.2±8.9; BMI: 34.7±5.6; female: 46.7% 30 adults in the gliclazide group: age: 55.1±6.4; BMI: 34.1±6.7; female: 60% 	<ul style="list-style-type: none"> GLP-1 analog 12-week liraglutide treatment 	<ul style="list-style-type: none"> Binge eating assessed by Binge Eating Scale 	<ul style="list-style-type: none"> Participants who received liraglutide had significant reductions in binge eating at 12 weeks

<ul style="list-style-type: none"> RCT 	<ul style="list-style-type: none"> Recruited from diabetes outpatient clinic Inclusion: diagnosis of type 2 diabetes being treated only with metformin, with suboptimal metabolic control, age below 65 years, with diagnosis of BED 			
<p>Meal-related acyl and des-acyl ghrelin and other appetite-related hormones in people with obesity and binge eating⁷²</p> <ul style="list-style-type: none"> Hernandez et al 2019 US Case-control 	<ul style="list-style-type: none"> 20 adults with binge eating (10 BED and 10 subthreshold binge eating): age: 37.8±8.4; BMI: 36.2±5.5; female: 60% 22 adults without binge eating: age: 34.0±8.2; BMI: 36.4±4.9; female: 31.8% Race in the total group: Black/African American (61%), Hispanic (19.5%), non-Hispanic White (14.6%), and other (4.8%). Recruited from non-clinical settings Exclusion: individuals with other eating disorders 	<ul style="list-style-type: none"> Fasting and postprandial GLP-1 Test meal: 1254 KJ liquid meal (55% carbohydrate, 24% protein, 21% fat) Blood draw: -15, and 0, 10, 30, and 60 minutes after meal The two groups did not differ in age, BMI, and sex 	<ul style="list-style-type: none"> Binge eating assessed by Eating Disorder Examination and the Questionnaire on Eating and Weight Patterns 	<ul style="list-style-type: none"> GLP-1 did not differ between groups
PP				
<p>Lifestyle intervention discloses an association of the Eating Inventory-51 factors with cardiometabolic health risks⁹⁴</p> <ul style="list-style-type: none"> Hainerova' et al 2013 Czech Republic Longitudinal 	<ul style="list-style-type: none"> 67 women with obesity: age: 48.7±12.2; BMI: 32.4±4.4 Women participated in a 4-week lifestyle obesity management (e.g., low-calorie diet, physical activity, and cognitive behavioral therapy) 	<ul style="list-style-type: none"> Fasting PP Assessment timepoints: before and after a 3-week weight management Covariates not specified 	<ul style="list-style-type: none"> Disinhibition subscale of Eating Inventory 	<ul style="list-style-type: none"> No significant associations between PP and disinhibition at baseline
<p>Hormonal and dietary characteristics in obese human subjects with and without food addiction⁶⁹</p> <ul style="list-style-type: none"> Pedram et al 2014 Canada 	<ul style="list-style-type: none"> 29 adults with food addiction: age: 42.5±9.6; BMI: 32.5±6; female: 82.7% 29 non-food-addicted subjects matched for age, sex, BMI and physical activity: age: 42±8.9; BMI: 32±4.4; female: 82.7% Recruited from non-clinical setting Inclusion: age >19 years, without serious 	<ul style="list-style-type: none"> Fasting ghrelin Covariates not specified 	<ul style="list-style-type: none"> Food addiction assessed by Yale Food Addiction Scale 	<ul style="list-style-type: none"> The 2 groups did not differ in terms of PP

<ul style="list-style-type: none"> Case-control 	<p>metabolic, cardiovascular or endocrine diseases, and not pregnant at the time of the study.</p>			
Leptin & Adiponectin				
<p>Relationship between dietary restraint, binge eating, and leptin in obese women⁷⁴</p> <ul style="list-style-type: none"> d'Amore et al 2001 Italy Case-control 	<ul style="list-style-type: none"> 8 women with severe binge eating: age: 46.4±5.9; BMI: 35.4±4.5 11 women with moderate binge eating: age: 46.0±14.5; BMI: 35.0±2.2 23 women without binge eating: age: 48.4±10.1; BMI: 33.9±2.9 Women were recruited before the beginning of a residential weight-reduction program None of the subjects was currently dieting or had markedly lost weight during the three months preceding the start of the study. Exclusion: type 1 diabetes; cancer; anaemia; any cardiac abnormalities; any past and present cerebrovascular, kidney, liver or thyroid disease; any significant psychiatric illness; smoking and use of medications known to affect weight or energy expenditure. 	<ul style="list-style-type: none"> Leptin Covariates not specified 	<ul style="list-style-type: none"> Binge eating measured by Binge Eating Scale 	<ul style="list-style-type: none"> Leptin was not significantly associated with binge eating However, severe binge-eaters showed higher leptin levels compared to non-binge-eaters, although the difference did not reach statistical significance.
<p>Serum leptin concentration in obese patients with binge eating disorder⁷⁵</p> <ul style="list-style-type: none"> Adami et al 2002 Italy Case-control 	<ul style="list-style-type: none"> 30 candidates for bariatric surgery 14 women with BED: BMI: 46.4±9.9 16 non-BED women: BMI: 46.5±7.8 All patients had stable weight in the past 6 months, were in completely good health, and did not take any medication known to influence metabolic parameters 	<ul style="list-style-type: none"> Leptin Covariates not specified 	<ul style="list-style-type: none"> BED assessed by Eating Disorder Inventory Disinhibition assessed by Eating Inventory 	<ul style="list-style-type: none"> Higher serum leptin concentrations were found in the BED patients In all patients, there was a significant positive associations between serum leptin and disinhibition
<p>Opposite modifications in circulating leptin and soluble leptin receptor across the eating disorder spectrum⁷⁶</p> <ul style="list-style-type: none"> Monteleone et al. 	<ul style="list-style-type: none"> 22 women with AN: age: 22.5±6.4; BMI: 15.9±1.5 45 women with BN-purging: age: 23.9±3.8; BMI: 20.9±1.9 18 women with BED: age: 29.0±9.8; BMI: 	<ul style="list-style-type: none"> Leptin Covariates included age 	<ul style="list-style-type: none"> BED assessed by Structured Clinical Interview for DSM-IV 	<ul style="list-style-type: none"> Women with BED had enhanced blood concentrations of leptin than non-BED women with obesity, but this difference did not reach

<ul style="list-style-type: none"> • 2002 • Italy • Case-control 	<p>35.9±8.9; binge frequency 2.7±1.1 episodes per day</p> <ul style="list-style-type: none"> • 12 non-BED women with obesity: age: 37.4±15.4; BMI: 37.0±2.8 • 33 healthy women: age: 24.8±4.1; BMI: 21.6±1.8 • Recruited from outpatient unit at eating disorder center • Exclusion: taking oral contraceptives, had a history of alcohol or drug abuse, present and past psychiatric disorders, endocrine diseases known to cause obesity, psychotropic drugs for more than 6 weeks 			<p>statistical significance</p>
<p>Gastric capacity, test meal intake, and appetitive hormones in binge eating disorder⁷³</p> <ul style="list-style-type: none"> • Geliebter et al. • 2004 • US • Case-control 	<ul style="list-style-type: none"> • 11 women with BED: age: 29±8; BMI: 36.6±6.2 • 13 women with subclinical binge eating: age: 29±7; BMI: 35.8±5.5 • 13 non-binge eating women: age: 32±9; BMI: 35.1±5.3 • Inclusion: premenopausal, nonsmokers, not taking illegal drugs or medications affecting weight, and weight stable within past 3 months. 	<ul style="list-style-type: none"> • Leptin • Covariates not specified 	<ul style="list-style-type: none"> • Binge eating assessed by Questionnaire on Eating and Weight Patterns and clinical interview 	<ul style="list-style-type: none"> • Leptin did not differ between groups (statistical data not provided)
<p>Leptin/adiponectin ratio in obese women with and without binge eating disorder⁶⁷</p> <ul style="list-style-type: none"> • Brandão et al • 2010 • Brazil • Case-control 	<ul style="list-style-type: none"> • 8 women with BED: age: 42.1±6.3; BMI: 32.3±2.1 • 7 non-BED women with obesity: age: 38.3±5.5; BMI: 34.9±3.9 • 8 normal weight women without BED: age: 44.9±6.6; BMI: 23.6±1.0 • Recruited from Piquet Carneiro Polyclinic • Women were not taking any medications and had no evidence of disease other than obesity and BED • Exclusion: a diagnosis of hypothyroidism, hyperthyroidism, diabetes, hypertension, 	<ul style="list-style-type: none"> • Leptin and adiponectin • Covariates not specified 	<ul style="list-style-type: none"> • Binge eating assessed by Binge Eating Scale 	<ul style="list-style-type: none"> • Women with BED had a lower leptin concentration than non-BED women with obesity • Adiponectin was lower in women with BED compared to the obese non-BED group

	and polycystic ovary syndrome, pregnant, breastfeeding or menopausal			
<p>Cardiovascular stress reactivity and recovery in bulimia nervosa and binge eating disorder⁷⁷</p> <ul style="list-style-type: none"> • Messerli-Bürgy et al • 2010 • UK • Case-control 	<ul style="list-style-type: none"> • 12 women with BN: age: 24.4±5.7; BMI: 23.1±5.6 • 13 women with BED: age: 33.9±7.0; BMI: 37.9±6.4 • 13 non-BED women: age: 41.1±8.9; BMI: 35.9±5.0 • Recruited from the psychiatric outpatient clinic • Participants should have stabilized electrolyte conditions, unchanged weight levels during the past 10 to 12 weeks • Exclusion: medical history of cardiovascular disease, metabolic disease, other disease or medication that could influence autonomic functioning 	<ul style="list-style-type: none"> • Leptin • Covariates not specified 	<ul style="list-style-type: none"> • Binge eating assessed by Eating Disorder Inventory 	<ul style="list-style-type: none"> • Leptin levels were lower in the BED group than non-BED group, but its statistical significance is unclear
<p>Lifestyle intervention discloses an association of the Eating Inventory-51 factors with cardiometabolic health risks⁹⁴</p> <ul style="list-style-type: none"> • Hainerova et al • 2013 • Czech Republic • Longitudinal 	<ul style="list-style-type: none"> • 67 women with obesity: age: 48.7±12.2; BMI: 32.4±4.4 • Women participated in a 4-week lifestyle obesity management (e.g., low-calorie diet, physical activity, and cognitive behavioral therapy) 	<ul style="list-style-type: none"> • Leptin and adiponectin • Assessment timepoints: before and after a 3-week weight management • Covariates not specified 	<ul style="list-style-type: none"> • Disinhibition subscale of Eating Inventory 	<ul style="list-style-type: none"> • Leptin was not associated with disinhibition at before and after intervention • Adiponectin was negative related to disinhibition after weight loss
<p>Relationships among tonic and episodic aspects of motivation to eat, gut peptides, and weight before and after bariatric surgery⁸⁷</p> <ul style="list-style-type: none"> • Bryant et al • 2013 	<ul style="list-style-type: none"> • 12 candidates for bariatric surgery (RYGB): age: 36±2; BMI: 45.3±1.9; female: 75% • Exclusion criteria not specified 	<ul style="list-style-type: none"> • Leptin • Assessment timepoints: pre-surgery, and 2 months, and 1 year post-surgery • Covariates included BMI 	<ul style="list-style-type: none"> • Uncontrolled eating subscale of the Three Factor Eating Questionnaire 	<ul style="list-style-type: none"> • Before surgery, there was a marginal, positive, association between leptin and uncontrolled eating

<ul style="list-style-type: none"> • UK • Longitudinal 				
<p>Hormonal and dietary characteristics in obese human subjects with and without food addiction⁶⁹</p> <ul style="list-style-type: none"> • Pedram et al • 2014 • Canada • Case-control 	<ul style="list-style-type: none"> • 29 adults with food addiction: age: 42.5±9.6; BMI: 32.5±6; female: 82.7% • 29 non-food-addicted subjects matched for age, sex, BMI and physical activity: age: 42±8.9; BMI: 32±4.4; female: 82.7% • Recruited from non-clinical setting • Inclusion: age >19 years, without serious metabolic, cardiovascular or endocrine diseases, and not pregnant at the time of the study. 	<ul style="list-style-type: none"> • Leptin and adiponectin • Covariates not specified 	<ul style="list-style-type: none"> • Food addiction assessed by Yale Food Addiction Scale 	<ul style="list-style-type: none"> • The 2 groups did not differ in terms adiponectin • Leptin was higher in the food addiction group, but not statistically significant
<p>Serum leptin and loss of control eating in children and adolescents⁷⁸</p> <ul style="list-style-type: none"> • Miller et al • 2014 • US • Case-control 	<ul style="list-style-type: none"> • 196 children and adolescents with LOC eating: age: 14.1±2.5; BMI: 29.1±8.9; female: 82.0% • 311 children and adolescents without LOC eating: age: 13.6±2.5; BMI: 29.8±11.9; female: 58.0% • Participants were either treatment-seeking or non-treatment seeking • Exclusion: major medical issues, major psychiatric conditions, pregnancy, medications known to affect weight, had lost more than 5% of their body weight in the 3 months prior to assessment, were currently involved in weight loss treatment programs 	<ul style="list-style-type: none"> • Leptin • Covariates included age, race, sex, socioeconomic, fat mass (kg), height (cm), dietary restraint, treatment-seeking status, and pubertal status 	<ul style="list-style-type: none"> • LOC eating assessed by Eating Disorder Examination: LOC eating was defined by the presence of one or more objective binge episodes and/or subjective binge episodes in the previous month. 	<ul style="list-style-type: none"> • Those reporting at least one episode of LOC in the past month had significantly higher leptin compared to those reporting no LOC episodes • This relationship was partially mediated by increased dietary restraint
<p>The association of serum leptin levels with food addiction is moderated by weight status in adolescent psychiatric inpatients⁸⁹</p> <ul style="list-style-type: none"> • Peters et al • 2018 	<ul style="list-style-type: none"> • 45 adolescents of underweight • 39 adolescents with overweight • 121 adolescents of normal weight • Total group: age: 16.1±1.1; BMI: 22.9±6.2; female: 60.5% • All of the participants had psychiatric disorders, including mood disorder, substance abuse. 	<ul style="list-style-type: none"> • Leptin • Covariates included sex, body fat percentage 	<ul style="list-style-type: none"> • Food addiction assessed by Yale Food Addiction Scale 	<ul style="list-style-type: none"> • In adolescents with overweight, there was a positive association between leptin and food addiction

<ul style="list-style-type: none"> Germany Case-control 	<ul style="list-style-type: none"> Recruited from inpatient units for adolescents Exclusion: intake of any psychopharmacological drug, any known endocrinological disorder, and intellectual disability 			
<p>Comparison of endocannabinoids levels, FAAH gene polymorphisms, and appetite regulatory substances in women with and without binge eating disorder: a cross-sectional study¹⁰¹</p> <ul style="list-style-type: none"> Yagin et al 2020 Iran Cross-sectional 	<ul style="list-style-type: none"> 180 premenopausal women: age: 34.2±8.3; BMI: 32.5±3.7; 41.6% of the subjects were diagnosed with BED Exclusion: chronic or metabolic disorders history, pregnancy or lactating, any appetite- affecting medicine usage, and significant weight loss over the last 3 months Participants were recruited from non-clinical settings 	<ul style="list-style-type: none"> Leptin Covariates included BMI 	<ul style="list-style-type: none"> Binge eating assessed by Binge Eating Scale 	<ul style="list-style-type: none"> Women with BED exhibited significantly higher levels of leptin compared to non-BED women Higher leptin was associated with higher odds of BED, adjusting for BMI
<p>Altered regional grey matter volume and appetite- related hormone levels in adolescent obesity with or without binge- eating disorder⁷⁹</p> <ul style="list-style-type: none"> Turan et al Turkey 2021 Case-control 	<ul style="list-style-type: none"> 25 adolescents with BED: age: 15.0±1.8; BMI-Z score: 2.9±0.4; female: 68% 25 non-BED adolescents: age: 14.6±1.7; BMI-Z score: 2.9±0.4; female: 68% 27 population-based healthy controls (matched with age, gender, and education): age: 14.6±1.4; BMI-Z score: -0.3±0.9; female: 70.4% Recruited from outpatient unit at the Department of Pediatrics and Department of Pediatric Endocrinology and Metabolism Exclusion criteria: history of psychiatric illness, intellectual disability, psychotic symptoms, drug/alcohol abuse, bipolar disorder, neurological disorders, claustrophobia, ADHD, left-handedness, a 	<ul style="list-style-type: none"> Fasting leptin 	<ul style="list-style-type: none"> BED assessed by Eating Disorder Examination Questionnaire 	<ul style="list-style-type: none"> Leptin was higher in the BED group than the non-BED group, but the difference was not statistically significant.

	history of AN or BN, and antipsychotic medication			
Inflammatory markers				
Lifestyle intervention discloses an association of the Eating Inventory-51 factors with cardiometabolic health risks ⁹⁴ <ul style="list-style-type: none"> Hainerova' et al 2013 Czech Republic Longitudinal 	<ul style="list-style-type: none"> 67 women with obesity: age: 48.7±12.2; BMI: 32.4±4.4 Women participated in a 4-week lifestyle obesity management (e.g., low-calorie diet, physical activity, and cognitive behavioral therapy) 	<ul style="list-style-type: none"> CRP Assessment timepoints: before and after a 3-week weight management Covariates not specified 	<ul style="list-style-type: none"> Disinhibition subscale of Eating Inventory 	<ul style="list-style-type: none"> No significant associations between CRP and disinhibition at baseline
Hormonal and dietary characteristics in obese human subjects with and without food addiction ⁶⁹ <ul style="list-style-type: none"> Pedram et al 2014 Canada Case-control 	<ul style="list-style-type: none"> 29 adults with food addiction: age: 42.5±9.6; BMI: 32.5±6; female: 82.7% 29 non-food-addicted subjects matched for age, sex, BMI and physical activity: age: 42±8.9; BMI: 32±4.4; female: 82.7% Recruited from non-clinical settings Inclusion: age >19 years, without serious metabolic, cardiovascular or endocrine diseases, and not pregnant at the time of the study. 	<ul style="list-style-type: none"> TNF-α Covariates not specified 	<ul style="list-style-type: none"> Food addiction assessed by Yale Food Addiction Scale 	<ul style="list-style-type: none"> The food addiction group had a significantly lower level of TNF-α as compared to the non-food addiction group
Obese patients with a binge eating disorder have an unfavorable metabolic and inflammatory profile ⁸² <ul style="list-style-type: none"> Succurro et al 2015 Italy Case-control 	<ul style="list-style-type: none"> 30 adults with BED: age: 36.8±12.7; BMI: 43.7±6.8; female: 73.3% 85 non-BED adults: age: 41.8±12.8; BMI: 37.2±6.2; female: 62.3% Patients were seeking weight reduction therapy Inclusion: aged between 20 and 65 years, BMI >30 kg/m² Exclusion: pregnancy or having recently given birth, previous diagnosis of diabetes mellitus, known inflammatory disease, a 	<ul style="list-style-type: none"> hsCRP, erythrocyte sedimentation rate, Covariates included age, sex, and BMI 	<ul style="list-style-type: none"> BED assessed by Binge Eating Scale and Clinical Interview 	<ul style="list-style-type: none"> BED-obese group had significantly higher ESR and hsCRP

	history of malignant disease or pathologies, or drugs able to modify glucose metabolism			
<p>Pediatric loss of control eating and high-sensitivity C-Reactive Protein concentrations⁸³</p> <ul style="list-style-type: none"> Shank et al 2017 US Case-control 	<ul style="list-style-type: none"> 75 children and adolescents with LOC eating: age: 13.5±2.3; BMI-z score: 1.9±0.8; female: 73.3%; race (non-Hispanic white: 37.3%, non-Hispanic black: 42.7%, Hispanic: 8%) 119 without LOC eating: age: 14.7±1.8; BMI-z score: 1.5±1.2; female: 58%; race (non-Hispanic white: 39.5%, non-Hispanic black: 48.7%, Hispanic: 2.5%) Inclusion: children or adolescents with overweight or obesity Exclusion: pregnancy, major medical or psychiatric illnesses, and use of medication known to affect weight or eating behavior 	<ul style="list-style-type: none"> hsCRP Covariates: sex, fat mass (kg), height (cm) 	<ul style="list-style-type: none"> LOC eating assessed by the child version of Eating Disorder Examination: LOC eating was deemed present if participants endorsed at least one episode of LOC eating in the past 28 days 	<ul style="list-style-type: none"> Presence of LOC eating was significantly associated with hsCRP concentration The number of LOC eating episodes was significantly associated with hsCRP concentration These associations were not mediated by depressive symptoms or eating-related psychopathology
<p>Brain-behavior-immune interaction: serum cytokines and growth factors in patients with eating disorders at extremes of the body mass index (BMI) Spectrum⁸⁰</p> <ul style="list-style-type: none"> Caroleo et al 2019 Italy Case-control 	<ul style="list-style-type: none"> 14 adults with AN: age: 25.1±11.6; BMI: 16.6±1.1 27 adults with BED: age: 41.0±11.8; BMI: 38.4±7.9 28 adults with obesity: age: 41.9±11.4; BMI: 42.2±10.5 21 healthy controls: age: 31.8±12.3; BMI: 21.3±2.7 Total sample: female (69%) Outpatients seeking treatment for eating disorder Exclusion: aged under 18 or over 65 years, patients with BN; normal-weight persons with any psychiatric comorbidity; diabetes mellitus, neurological or other medical conditions; hormonal and pharmacological treatment; smokers; pregnancy or childbirth over the previous 12 months 	<ul style="list-style-type: none"> Pro- and anti-inflammatory cytokines (IL-1α, IL-1β, IL-6, IL-10, TNF-α) Controlling for depressive symptoms, sex, and age 	<ul style="list-style-type: none"> BED assessed by Eating Disorder Examination 	<ul style="list-style-type: none"> IL-1α was not significant different between BED and obesity group IL-10 in the BED group was higher than obesity group (statistical difference unknown) There were no significant differences in IL-1β, IL-6, IL-8, TNF-α between BED and obesity group.

651 Abbreviations:

652 BED: binge-eating disorder; BN: bulimia nervosa; AN: anorexia nervosa; BMI: body mass index; DSM: Diagnostic and Statistical Manual of
653 Mental Disorders; CBT: cognitive behavioral therapy; CCK: cholecystinin; PYY: peptide YY; GLP-1: glucagon-like peptide 1; PP: pancreatic
654 polypeptide; IL-1 α : interleukin-1 α ; IL-1 β : interleukin-1 β ; IL-6: interleukin-6; TNF- α : tumor necrosis factor- α ; CRP: C-reactive protein; ESR:
655 erythrocyte sedimentation rate; IL-10: interleukin-10; IL-13: interleukin-13.

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Ethical Statement

This manuscript was a review; therefore, it did not require ethics approval.

Journal Pre-proof