Regulation of Appetite, Satiation, and Body Weight by Enteroendocrine Cells. Part 2:
Therapeutic Potential of Enteroendocrine Cells in the Treatment of Obesity

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Key Words
Enteroendocrine cells · Satiation · Appetite · Obesity · Taste receptors

Abstract
Obesity is an epidemic and medical issue. Investigating the pathways regulating appetite, food intake, and body weight is crucial to find strategies for the prevention and treatment of obesity. In the context of therapeutic strategies, we focus here on the potential of enteroendocrine cells (EECs) and their secreted hormones in the regulation of body weight. We review the role of the enteroendocrine system during weight loss after lifestyle intervention or after bariatric surgery. We discuss the therapeutic potential of EECs and their hormones as targets for new treatment strategies. In fact, targeting nutrient receptors of EECs with a nutritional approach, pharmaceutical agents or prebiotics delivered to the lumen may provide a promising new approach.

Introduction
The current worldwide obesity epidemic is a growing medical issue. In fact, overweight and obesity are the fifth leading risk for global deaths as stated in the WHO fact sheet No. 311. Importantly, even a moderate weight loss has a significant health benefit. Dietary lifestyle interventions reducing energy intake effectively reduce weight but do not work well in the long term [1]. If lifestyle interventions fail to control body weight, we need other effective and safe treatments. To date, only bariatric surgery is efficient for long-term weight reduction and reduces overall mortality in severely obese patients [2]. The convincing effect of these surgical interventions is not only due to limiting food ingestion or malabsorption but also due to altered gut hormone release [e.g. peptide YY (PYY), glucagon-like peptide 1 (GLP-1), and ghrelin] [3] affecting appetite and satiety. Pharmacological approaches using gut hormones are often disappointing as these hormones have a short half-life in the blood (e.g. GLP-1) and adverse effects. Therefore, investigating gut hormone-producing enteroendocrine cells (EECs) as targets for weight management seems to be reasonable.

Targeting EEC nutrient receptors and thereby inducing the release of their hormones, e.g. GLP-1 via GPR119, is possible and might be an alternative treatment option also in humans [4]. In fact, nutrition alters the differentiation of EECs and, thus, the gut hormone profile, e.g. L-cell differentiation and the release of GLP-1 are promoted by non-digestible carbohydrates [5]. Finally, the microbiome and its products directly influence differentiation and function of EECs [6, 7]. On these grounds,
prebiotics may have beneficial therapeutic properties in the treatment of obesity [8]. Here, we review the therapeutic potential of EECs in the treatment of obesity.

EECs in the Context of Therapeutic Approaches towards Body Weight Control

Energy homeostasis, satiety and body weight are centrally regulated in a highly complex system comprising orexigenic centres with neurons expressing NPY and AgRP or anorexigenic centres with proopiomelanocortin (POMC) as well as cocaine and amphetamine-regulated transcript (CART)-expressing neurons in the hypothalamus. Peripheral regulators comprise hormones derived from the gastrointestinal (GI) tract, the pancreas, and the adipose tissue [9]. The gut-brain axis involves both neuronal and hormonal signals that are generated in the GI tract (fig. 1) [10]. The integration of peripheral signals is not limited to the hypothalamus, but includes other brain areas such as the limbic system, the cortex, the midbrain, and the brain stem. Signals induced by the hormones of EECs in response to the volume of food, the caloric content, and the quality of food are integrated in different brain areas, which participate in appetite regulation, such as the hypothalamus (fig. 1). Some of the key neurons are located in the arcuate nucleus (ARC), which is located outside the blood-brain barrier. Thus, the ARC is accessible by circulating hormones, e.g. ghrelin, which activates appetite-stimulating neuropeptides NPY and AgRP. The neuronal input of satiety from the periphery is mediated via the vagus nerve to the brain stem and the nucleus tractus solitarius. Gut hormones play an important role in the digestive process, regulating appetite and satiety via the gut-brain axis as reviewed in part 1 [11]. Ghrelin, PYY, GLP-1, and cholecystokinin (CCK) are released in the periphery in response to the presence or absence of nutrients in the gut and are able to act peripherally on the vagus nerve and centrally on target areas in the hypothalamus (fig. 1).

In addition, a crosstalk between the adipose tissue and the gut may also be relevant in the context of the regulation of energy homeostasis, satiety, and body weight. Leptin is released continuously from adipose tissue into the circulation and acts mainly on the hypothalamus to regulate long-term energy storage (fig. 1). In addition, exocrine-secreted gastric leptin is proposed to ensure proper food processing and food intake in the short term independent of adipose-derived leptin [12]. Thus, gastric- and adipose-derived leptin may complement one another [13].

GI Hormones during Weight Changes

Lifestyle intervention is recommended as first-line therapy for adult as well as for childhood obesity and comprises diet, exercise, and behavioural interventions [14, 15]. Programmes for obese children, which target the parents and home settings, are more effective for weight loss or maintenance than programmes which only focus...
on the behaviour of the children [15, 16]. It is well known that a hypocaloric diet leads to promising weight loss during the first months.

So far, the reported changes of GI hormones in connection with weight loss during lifestyle intervention are inconclusive. In adults, weight loss leads to a significant reduction in circulating levels of leptin, PYY, CCK, and insulin [17]. This is accompanied by a sustained increase in ghrelin and gastric inhibitory peptide (GIP) levels in the circulation [17]. Obesity in children is associated with reduced serum levels of ghrelin and PYY, while GLP-1 levels are unchanged compared to lean children [18–20]. Some data in children have shown that weight loss leads to an increase in PYY serum levels, while ghrelin levels remain unchanged or increase and GLP-1 levels decrease [18–21]. The initial alterations of gut hormones in obese children seem, therefore, to be reversible after weight loss [9]. One study, however, showed no changes in fasting or postprandial GLP-1 and PYY levels in obese children after weight reduction [22].

After weight loss during lifestyle intervention programmes, the majority of individuals will regain weight within the following months [1]. The high rate of relapses may be at least partly explained by the long-lasting counter-regulatory effect of several appetite-regulating hormones produced by the EECs. In fact, a recent study in adults has demonstrated that changes in circulating GI hormones induced by weight loss persist even after 1 year [17]. Thus, they may contribute to the observed propensity to regain weight, and long-term strategies to counteract these changes may be needed to prevent obesity relapse.

**The Role of EECs in Bariatric Surgery**

Bariatric surgery is the only successful treatment option leading to long-term weight loss with clinical relevance resulting in an improvement of obesity-associated comorbidities including type 2 diabetes. The most frequently applied method is the Roux-en-Y gastric bypass (RYGB) operation.

Recent data on patients after bariatric surgery show that unlike the effects of lifestyle changes and hypocaloric diets, bariatric surgery results in meaningful increases in circulating levels of PYY and GLP-1 and a decrease in ghrelin level [23]. GI bypass operations create a connection of two separated segments of the gut, thereby bypassing the entire duodenum and part of the jejunum, and resulting in an altered flow of nutrients reaching the distal part of the small intestine more rapidly than usual. According to the ‘lower intestinal hypothesis’ [24] (also known as ‘distal’ or ‘foregut hypothesis’), the rapid delivery of nutrients to the lower intestine and the colon leads to an excessive stimulation of L cells resulting in an altered secretion profile of GI hormones. In addition, bile acids which reach the distal intestine and the colon in an undiluted state via the biliopancreatic loop lead to a stimulation of L cells by the farnesoid receptor X [25]. Indeed, most data support this theory, showing increased levels of GLP-1, PYY, and oxyntomodulin (OXM) after RYGB operation [26, 27]. However, in one study, a decrease in GLP-1 levels was observed after bariatric surgery of morbidly obese adults [28].

The secretion of GLP-1 and PYY is directly associated with the caloric content of the chyme reaching the distal intestine. There is a positive correlation between the postprandial PYY and GLP-1 levels and the amount of weight loss after a RYGB operation [29]. Along with an elevated GLP-1 level, a decrease in dipeptidyl peptidase-4 (DPP-4), which rapidly degrades GLP-1, was found in patients after bypass surgery [30]. Interestingly, experimental inhibition of PYY and GLP-1 secretion with the somatostatin analogue octreotide resulted in an increased appetite and weight gain after RYGB operation [29]. Thus, an increase in PYY and GLP-1 after RYGB operation may account for the beneficial effects on satiety.

Besides the effects of these satiety hormones on appetite and food intake after bariatric surgery, a second mechanism leads to the enormous benefits of bariatric surgery. Gastric bypass operations lead to successful diabetes resolution in up to 80% of patients lasting up to 14 years postoperatively [31]. It is now clear that rapid resolution of type 2 diabetes is achieved by bypass surgery before substantial weight loss is observed. This is partly due to an increase in GLP-1 secretion. GLP-1 is able to improve glucose homeostasis by stimulating insulin secretion and GIP secretion and by suppressing glucagon secretion [32].

**Is There an Alternative to Bariatric Surgery?**

Pharmacological treatment strategies imitating the hormonal and malabsorptive changes induced by bypass surgery could probably be as effective as bypass surgery. For example, fat malabsorption can be induced by gastric and pancreatic lipase inhibitors such as Orlistat or Cetili-
Moreover, combining gut hormones effects, e.g. sibutramine or rimonabant have been withdrawn from the market due to severe side effects, and obesity drugs working on the central nervous system have been found to be effective. In obese rodents, the insulinotropic effect in synergism with a loss of fat mass of GLP-1 and GIP showed an anti-hyperglycaemic and hypolipidaemic effect. The endogenous release of gut hormones may be induced by the acetylated analogue liraglutide with a considerably extended half-life up to 11–15 h. A recently discovered peptide derived from an intermixed sequence was solved by the acetylated analogue liraglutide with a considerable extended half-life up to 11–15 h. A recently discovered peptide derived from an intermixed sequence of GLP-1 and GIP showed an anti-hyperglycaemic and insulinotropic effect in synergism with a loss of fat mass in obese rodents.[39]

Taken together, in addition to lifestyle interventions, future multifactorial long-term medical treatment approaches for obesity might include gastric and pancreatic lipase inhibitors as well as dietary or pharmacological regulators of gut hormones, e.g. simulating GLP-1, GIP, or PYY releases.

**Appetite Regulation by Targeting EEC Nutrient Receptors**

New anti-obesity drugs often fail to achieve a significant weight loss greater than 5% and are not superior to educational programs. In addition, they lead to side effects, most commonly nausea, cardiovascular events, and psychological side effects.[40] Moreover, effective anti-obesity drugs working on the central nervous system have been withdrawn from the market due to severe side effects, e.g. sibutramine or rimonabant.[41] These drugs mainly affected the central nervous system. Thus, we need to consider the potential of EECs and their hormones to regulate food intake, digestion, and thereby body weight without the adverse effects caused by systemic administration or by affecting primarily a central target. Triggering the enteroendocrine network by administration of nutritive factors or drug compounds might increase the efficacy of weight loss and simulate the physiologic signals of satiation and appetite control.

**Nutritional Factors Influence the Plasticity of EECs**

The very high turnover of EECs due to their short life span of around 4–6 days allows an extensive plasticity in contrast to other non-diffuse endocrine organs.[44] EECs revolve continuously from pluripotent intestinal stem cells from the villous crypts.[11] Differentiation is controlled by basic helix-loop-helix (bHLH) transcription factors. Interestingly, high-fat feeding reduces the expression of bHLH transcription factors and thereby the number of EECs.[45] In addition, intestinal gut hormone levels show a subsequent decrease.[45], demonstrating that high-fat feeding influences EEC differentiation and may thereby influence satiety and the development of obesity (fig. 3). Indeed, a high-fat diet suppresses lipid-induced CCK satiation signalling in obese-prone rats.[46] Furthermore, high-energy/high-fat feeding results in reduced GLP-1 serum levels in these rats.[47] Other studies demonstrated that non-digestible carbohydrates promote L-cell differentiation in the proximal colon via upregulation of bHLH transcription factors neurogenin 3 and NeuroD resulting in higher GLP-1 production.[5] Dietary fibres such as oligofructose have been shown to reduce food intake (fig. 2).[48] Finally, changing the
quality of nutritional intake is accompanied by a modulation of the gut microbiota, which in turn probably affects EECs.

The Impact of Microbiota on EECs and Food Intake

The microbiome influences gut homeostasis. Bacterial metabolic products or bacterial factors are able to directly interact with the intestinal epithelium. Furthermore, microbes can activate EECs to secrete, for example, serotonin and thereby stimulate enteric nerves and regulate GI motility and secretion. There is evidence that the EEC metabolism is controlled by the microbiome. Among pathological conditions, the microbiome plays a role in the development of diabetes and obesity by triggering low-grade inflammation, gut barrier dysfunction, and metabolic endotoxaemia [8]. Gut microbiota-derived lipopolysaccharides (LPS) have been identified as a key factor in the early development of low-grade inflammation and metabolic diseases [8, 49]. In addition, fat feeding and a high-fat diet induce elevated plasma LPS levels, the so-called metabolic endotoxaemia [49] (fig. 3).

Identifying the underlying mechanism of low-grade inflammation and metabolic endotoxaemia is therefore important.

Indeed, a significant change in the microbiome of obese individuals is present with lower anti-inflammatory Bacteroides proportions compared to lean individuals [50]. Even a microbiome with higher levels of Staphylococcus aureus and lower levels of Bifidobacteria during childhood may predict overweight [51]. However, it is still unclear whether gut microbiota alterations are caused by the diet or by the pathology of obesity itself. Interestingly, faecal transplantation of gut microbiota from a lean healthy donor temporarily improves insulin sensitivity in patients with metabolic syndrome [52].

According to rodent models, modulation of the microbiome by using prebiotics may improve gut barrier function, metabolic endotoxaemia, and inflammation in obesity and type 2 diabetes [53]. Prebiotics are able to decrease LPS levels in the circulation and increase faecal concentrations of short-chain fatty acids (SCFAs) [54]. Fibre fermentation leads to SCFAs, which can induce the expression of proglucagon, the precursor of GLP-1 [6]. Thus, prebiotics may change food intake and GI hormone secretion and influence the microbiome (fig. 2). Indeed, prebiotics increase GLP-1 and PYY and decrease ghrelin secretion in humans [6, 7]. In addition, the positive effects of prebiotics are associated with an increase in the num-

Fig. 2. Effects of prebiotics and endovanilloid compounds on EEC behaviour and food intake. EECs (violet) sense the luminal content on the apical membrane with, for example, G-protein-coupled receptors (GPRs). The release of hormone-containing secretory vesicles (pink) is controlled by the intracellular Ca²⁺ content (pale blue). Prebiotics, non-digestible fiber compounds such as oligofructose, induce SCFAs and increase the EEC number and thereby alter peptide release. In addition, prebiotics change the gut microbiota composition with a reduction of LPS and thereby increase the integrity of the gut barrier and prevent bacterial metabolites from crossing the barrier, entering the circulation, and promoting systemic inflammation. Similar endovanilloid compounds and GPR119 agonists induce the expression of incretin hormones and thereby suppress food intake and weight gain.
The number of L cells in the intestine and GLP-2-mediated improvement of gut permeability [53, 55]. This effect is dose dependent and suggests that prebiotics may have a therapeutic potential for the treatment of obesity and type 2 diabetes [8].

**Conclusion**

There is an enormous demand on an effective obesity treatment, but there is concern about the safety of anti-obesity drugs. Medical combination therapies imitating the physiological changes observed after bypass surgery might be as effective as bariatric surgery. Unfortunately, effective anti-obesity drugs have been withdrawn from the market due to severe side effects. Thus, in addition to lifestyle intervention, targeting specifically EECs as key regulators in the peripheral control of food intake and obesity seems to be a promising strategy with less adverse events. Indeed, GI hormones from EECs play an important role in the sustained weight loss and metabolic improvements achieved by gastric bypass operations. Recent advances in our knowledge regarding the food-sensing skills of EECs and the interaction with different macronutrients or diets and the gut microbiota may lead to new therapeutic approaches, starting with dietary modifications and prebiotics as a considerable strategy to prevent and treat metabolic diseases. Investigating the EEC network as a therapeutic target in the context of body weight regulation and type 2 diabetes is promising.

**Disclosure Statement**

The authors declare that they have no conflicts of interest.

**References**


**Fig. 3.** Effects of high-fat diet on EEC behaviour and food intake. EECs (violet) sense the luminal content on the apical membrane with, for example, G-protein-coupled receptors (GPRs). The release of hormone-containing secretory vesicles (pink) is controlled by the intracellular Ca\(^{2+}\) content (pale blue). A high-fat diet alters the EEC number and peptide release, i.e. the lipid-mediated satiety signalling via CCK is reduced and thereby may promote food intake and obesity. In addition, a high-fat diet changes the gut microbiota composition and thus increases LPS together with a reduction of bacteria necessary for the integrity of the gut barrier, allowing LPS and other bacterial metabolites to cross the barrier, enter the circulation (green arrow), and promote systemic inflammation.
Gastrointestinal Satiety Signalling and Obesity


Hellstrom PM: Satiety signals and obesity. Horm Res Paediatr 2015;83:11–18

DOI: 10.1159/000369555


