Glucagon-like peptide 1 receptor (GLP-1R) signaling in the CNS has been linked to reduced food intake, lower body weight, improved glucose homeostasis, and activation of CNS stress axes. GLP-1 is produced by cells that express proglucagon (GCG); however, the stimuli that activate GCG+ neurons are not well known, which has made understanding the role of this neuronal population in the CNS a challenge. In this issue of the *JCI*, Gaykema et al. use designer receptors exclusively activated by designer drugs (DREADD) technology to specifically activate GCG+ neurons in mouse models. While activation of GCG+ neurons did reduce food intake, and variably decreased hepatic glucose production, other GLP-1–associated effects were not observed — e.g., activation of stress axes or stimulation of insulin secretion — in response to GCG+ neuron activation. The authors have provided a valuable model to study this set of neurons in vivo, and their results provide new insights into the function of GCG+ neural activity in the brain and raise questions that will move research on this clinically relevant neural system forward.
DREADDing proglucagon neurons: a fresh look at metabolic regulation by the brain

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GLP-1 signaling in the CNS

The development of drugs based on the glucagon-like peptide 1 (GLP-1) signaling system has been the major advance in diabetes therapeutics over the past two decades (1). GLP-1 receptor (GLP-1R) agonists and dipeptidyl peptidase 4 inhibitors (DPP4i) are now commonly used to control hyperglycemia and lower glycated hemoglobin (HbA1c) levels in patients with type 2 diabetes (T2D). The efficacy of both GLP-1R agonists and DPP4i has generally been attributed to their actions in the pancreatic islet, where these drugs enhance insulin secretion and suppress glucagon levels. It is notable that GLP-1R agonists, which raise circulating GLP-1 activity to supraphysiological levels, also decrease food intake and cause modest body weight loss, whereas DPP4i increase physiological GLP-1 activity and have no impact on food intake or body weight. In mammals, GLP-1 is synthesized only in the three cell types that express the proglucagon-encoding gene (Gcg): enteroendocrine L cells in the intestinal mucosa; α cells in the islet; and a discrete population of neurons in the hindbrain, primarily the nucleus of the solitary tract (or nucleus tractus solitarius [NTS]). Most circulating GLP-1 is thought to originate in the gut, and nutrient-stimulated secretion supports the conventional endocrine model of GLP-1 action. The role of locally produced GLP-1 in the islet is a matter of some debate, although arguments for paracrine actions are plausible (2). The physiological importance of CNS-derived proglucagon products has been difficult to ascertain, although the distribution of the GLP-1R has been mapped across the CNS (3, 4), and pharmacologic studies to activate and antagonize these receptors have demonstrated a number of reproducible, if rather diffuse effects of brain GLP-1 (reviewed in refs. 2, 5).

Given the high concentration of GLP-1R in the hypothalamus, it is not surprising that many of the responses attributed to GLP-1 in the CNS are related to energy and nutrient metabolism. GLP-1R activation in specific areas of the brain potently decrease food intake, either as a consequence of engaging neuronal pathways that increase satiety or by activating neurons that lead to aversion or malaise. Chronic administration of GLP-1R agonists directly to the CNS decreases body weight in animal models in conjunction with decreased caloric consumption (5). In addition to the effects on body weight, there is evidence that GLP-1 regulates glucose homeostasis via neuronal pathways that enhance insulin secretion, decrease hepatic glucose output, and increase glucose disposal into skeletal muscle (6, 7). Finally, GLP-1 stimulates two different stress axes in the CNS — the hypothalamic-pituitary-adrenal (HPA) axis and the sympathetic nervous system (SNS) — thereby causing elevations in circulating corticosterone and epinephrine.

Activating Gcg-expressing neurons with DREADDs

The stimuli that activate hindbrain Gcg-expressing neurons are not well established and have been difficult to ascertain in controlled experiments. Thus, understanding the physiology of GCG-positive neurons has depended entirely on specific gain- and loss-of-function studies of GLP-1R signaling in the CNS. In this issue (8), Gaykema and colleagues present a method that they devised for interrogating the brain GLP-1 system. Their approach involves using designer receptors exclusively activated by designer drugs (DREADD) technology to activate Gcg-expressing neurons. Stereotaxic administration of an adeno-associated virus (AAV) into the caudal medulla delivered a Gqα-coupled construct that is expressed in the presence of Cre recombinase. Expression of the Gqα-coupled DREADD (hM3Dq)
in Gcg-Cre mice resulted in selective expression of this receptor in GCG+ neurons, which could then be activated with clozapine N-oxide (CNO). This system proved to be both specific and efficacious for activating GCG+ neurons with CNO, as evidenced by both colocalization of Fos and GLP-1, as well as electrophysiological recordings of neuronal firing following CNO administration. Remarkably, activation of GCG+ neurons failed to engage many of the previously defined actions of GLP-1R agonism in the CNS (Figure 1). Acute treatment with CNO did not impact insulin levels or skeletal muscle glucose uptake. And while CNO modestly decreased gluconeogenesis in mice on a standard rodent diet, this effect did not translate into a measurable impact on glucose tolerance. Furthermore, CNO administration did not engage the HPA axis, elevate SNS activity, or change anxiety behaviors, all of which have been noted in previous studies of GLP-1R stimulation in the CNS. The most apparent effect on GCG+ neuron activation was a robust decrease in food intake, which was attributed to an increase in satiety rather than malaise. Chronic GCG+ neuron activation failed to induce changes in body weight in mice on a standard diet. Interestingly, decreased food intake was magnified when the experiment was repeated in mice on a high-fat diet (HFD), to the extent that chronic CNO treatment caused a decrease in body weight. This interaction of diet and GCG+ neuron activation failed to elicit previously demonstrated effects of CNS GLP-1 leaves open the question as to which neurotransmitter these cells express. It still remains to be seen whether these discrepancies arise due to differences in methodology (i.e., physiology versus pharmacology) or if activation of GCG+ neurons provides more precise activation of biological events (i.e., only food intake). OXM, oxyntomodulin; Ach, acetylcholine; NE, norepinephrine.
normal neural function. On the other, in order to fully interpret these findings more details and validation of the Gcg-DREADD model are needed, including whether activation by CNO achieves the full range of physiologic neural activation and what factors are released from GCG+ nerves. In fact, based on the data presented, it can only be assumed that the effect of CNO to reduce food intake involves GLP-1, or Gcg-associated products. Combining the Gcg-DREADD model with CNS administration of GLP-1R or GCG receptor antagonists could provide clarity here. Moreover, it remains possible that food intake is the most sensitive response to GCG+ neural activation, with stress and illness responses occurring only when the system is firing at near maximal rates. Nonetheless, the Gcg-DREADD model provides a firm base from which to address a tractable neural system that likely plays a significant role in systemic metabolism.

**Future considerations**

Some of the findings of Gaykema et al. (8) support their contention that targeting GCG+ neurons might have distinct potential for therapeutic development. For example, it would be very advantageous if activation of these nerves can cause satiety without malaise or visceral illness, since nausea is currently the limiting side effect of GLP-1R agonists in clinical use. It is also interesting to consider whether targeting GCG+ neurons might truly alleviate activation of stress responses seen with GLP-1R agonists. This could provide greater or more lasting weight loss, as CNS administration of GLP-1 to animals (9) and peripheral GLP-1 administration to humans (10) increase circulating glucocorticoids as well as satiety, responses with opposing impacts on body weight. While these examples are speculative at present, they provide an impetus for a better understanding of GCG+ neurons and their physiologic and pharmacologic activation.

One of the most interesting findings in these studies is the enhanced effect of GCG+ neuron activation on anorexia and weight loss in obese mice. These findings are consistent with previous work showing potentiation of GLP-1 effects on feeding by leptin (11), a peptide increased in obesity. This interaction suggests a role for GLP-1 to mitigate the effects of metabolic stress in the CNS, potentially as a mechanism to restore homeostasis. A similar hypothesis has been proposed for understanding the role of GLP-1 produced in other tissues. α Cell production of GLP-1 is minimal in healthy rodent and human islets yet increases dramatically in the face of stressors such as proinflammatory cytokines (12) or streptozotocin (13). Hence, GLP-1 produced in α cells is situated to respond to metabolic stress by stimulating β cell function and promoting glucose homeostasis. In the gut, both IL-6 and bacterial endotoxin stimulate GLP-1 from L cells, and GLP-1Rs in the intestine are located on intraepithelial lymphocytes (14), suggesting a mucosal GLP-1 system that is responsive to environmental stress. This broad framing of GLP-1 actions may provide some predictive utility in understanding Gcg-encoded peptides in the three regions where they are produced. The contribution of Gaykema and colleagues presented here provides a tangible means to move ahead in the study of the least-understood of these regions, the brain.

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