Long-acting glucagon-like peptide-1 receptor agonists have direct access to and effects on pro-opiomelanocortin/cocaine- and amphetamine-stimulated transcript neurons in the mouse hypothalamus

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INTRODUCTION
Glucagon-like peptide-1 (GLP-1) was first identified in 1983, cloned by Graham Bell as part of the pre-proglucagon sequence, with the incretin effect being published in 19871–3. As the other incretin, glucose-dependent insulinotropic polypeptide (GIP), has little effect on insulin secretion in patients with type 2 dia-
abetes, drugs giving pharmacological levels of GLP-1 receptor (GLP-1R) agonists have become successful therapies in the treatment of type 2 diabetes, providing glucose control as well as weight loss4–7. GLP-1 was first shown to lower food intake in rodent animal models in 1996, followed by a study in humans in 19984–7. The study in humans showed a reduction in energy intake, and also reported that the mechanism involved a reduction in appetite along with an increase in satiety and a reduction in feelings of hunger, as shown by use of a visual analog scale questionnaire7. The most recently approved GLP-1R agonists for...
management of type 2 diabetes are dulaglutide and albiglutide, which have been developed for once-weekly administration. However, as more compounds are approved and results from an increasing number of large, randomized, controlled, double-blinded trials have been published, treatment-associated weight loss in patients with type 2 diabetes differs between compounds. Liraglutide is a once-daily compound acylated with a fatty acid to facilitate non-covalent binding to albumin in vivo as the protraction mechanism. Although there were some differences in glycemic control, a consistent finding was that liraglutide resulted in greater weight loss than dulaglutide and albiglutide, which are much larger molecules modified with covalent addition of either a crystallizable fragment or an albumin molecule. Furthermore, of the five GLP-1R agonists now approved in treatment of type 2 diabetes in various regions of the world, liraglutide is the only one that has also been investigated and approved for weight management, as adjunct to diet and exercise. The present mini-review focuses on a novel pathway in the brain that might mediate the weight-lowering effect of GLP-1R agonists, and reviews these data in relation to the rather large existing literature for GLP-1R action in the brain as well as in the periphery.

**LIRAGLUTIDE DIRECTLY ACCESSES THE RODENT BRAIN**

Using the novel technique of single-plane illumination microscopy (SPIM), whereby the entire perfused brain is scanned after peripheral administration of fluorescently labeled liraglutide (liraglutide) in live mice, two-dimensional digital images of the entire brain were obtained and assembled into a three-dimensional image for optimal analysis of spatial distribution. The images were analyzed carefully, and the anatomical location of liraglutide described. Figure 1 shows a dorsoventral

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**Figure 1** | Distribution of fluorescently labeled liraglutide in the mouse brain. Representative whole brain images viewed in the (a) dorsoventral or (b) sagittal plane from C57BL/6J mice given liraglutide (unspecific staining has been removed from the left side of the brain). The brain tissue was scanned at 620 nm and 710 nm, representing both autofluorescence from the tissue (gray) and specific signal (green). The (c, f, i) red regions are shown at (d, g, j) higher magnification, respectively. (d, e, g, h, j, k) High-magnification views of a single section from (d, g, j) C57BL/6J or (e, h, k) Glp1r−/− mice given liraglutide. Liraglutide was detectable in (c, d) paraventricular nucleus of the hypothalamus (PVN), (f, g) the median eminence (ME), the arcuate nucleus (ARC), and (i, j) area postrema (AP). (e, h, k) In mice lacking a functional glucagon-like peptide-1 receptor (GLP-1R), no liraglutide signal could be detected in any of these regions. Scale bars, 200 μm (a, b, c, f, i); 50 μm (d, e); 100 μm (g, h, j, k). ©American Society for Clinical Investigation and reproduced from Secher et al. with permission.
(Figure 1a), or sagittal (Figure 1b), three-dimensional plane view of liraglutide594 distribution in the mouse brain. As can be seen, liraglutide594 was distributed to discrete regions of the brain. Regions with clear uptake were determined to be the paraventricular nucleus of the hypothalamus (PVN; Figure 1c, d); the arcuate nucleus (ARC) and the median eminence (ME) of the hypothalamus (Figure 1f,g); and the area postrema (AP) in the hindbrain, whereas there was no uptake in the nucleus tractus solitarius in the hindbrain (Figure 1i,j). The signal in all sites of uptake were found to depend entirely on expression of the GLP-1R, as no uptake was detectable at these sites in GLP-1R−/− mice (Figure 1e,h,k). Uptake was also found in the organum vasculosum of the lamina terminalis, the subformical organ, the supraoptic nucleus and the supraoptic decussation, as well as the choroid plexus. Except for the choroid plexus, the signal in these regions also depends on the GLP-1R. The uptake patterns for liraglutide in the pancreas and the hypothalamus were compared with the patterns of the specific GLP-1R antagonist exendin(9-39; Figure 2). A different fluorophore was used for these experiments, enabling co-staining for insulin or cocaine- and amphetamine-stimulated transcript (CART), respectively. In the pancreas, liraglutide594 was found to co-localize entirely with pancreatic β-cells, as evidenced by an overlap with insulin staining. Also, the liraglutide594 signal was present within the cells as evidence of internalization with the GLP-1R (Figure 2i,j). In contrast, exendin(9-39)594 was retained at the plasma membrane of the β-cells (Figure 2k,l). In the brain, liraglutide594 also appeared to be present within the neurons (Figure 2m,n), whereas exendin(9-39)594 appeared to be retained at the membrane. However, in the brain, internalization is more difficult to assess because of the high density of neuronal fibers. To further examine the precise localization of liraglutide594 in the ARC, co-staining with the CART neuropeptide that labels CART/pro-opiomelanocortin (POMC) neurons was carried out (Figure 3). Liraglutide594 was detected specifically in the ARC in the cytoplasm of neurons positive for CART (Figure 3). Nearly all CART-positive cells were positive for liraglutide594. However, a few cells were only positive for liraglutide594, showing that another cell type might be targeted by liraglutide in the hypothalamus.

**DISCUSSION**

It has been suggested that physiologically the inhibitory effect of GLP-1 on gastric emptying might be more important than the incretin effect1-3. Delayed gastric emptying leading to prolonged gastric distention might induce short-term satiety, and could therefore be the relevant mechanism for the postprandial changes in appetite induced by physiological doses of GLP-1. However, such mechanisms are unlikely to cause the lasting effects on fasting and postprandial appetite seen with treatment with long-acting GLP-1R agonists in patients with and without type 2 diabetes14-16. With continued exposure beyond the duration of a normal postprandial period, the ability of GLP-1 to delay gastric emptying is much diminished17. Similarly, liraglutide and other long-acting GLP-1R agonists have little effect on gastric emptying, making it unlikely that this is the main weight loss mechanism18,19. The brain is the integrating site for appetite regulation, and numerous studies have documented the expression and importance of GLP-1Rs in the brain, and its importance in the physiological regulation of appetite20. GLP-1 is also produced in the hindbrain, and has been proposed as a physiological appetite reduction signal21. Peripherally-acting GLP-1R agonists might communicate with the brain through the nodose ganglion where GLP-1Rs are expressed, either in the form of GLP-1 released postprandially from the L cells in the gastrointestinal tract or in the form of an exogenously dosed GLP-1R agonist. Sisley et al.22 described two novel animal models where mice were genetically engineered to be knockdown models for brain or nodose ganglion GLP-1R expression, respectively. They found that the majority of liraglutide-induced weight loss required brain GLP-1R expression, whereas similar weight loss was obtained in normal and nodose ganglion GLP-1R expression knockdown mice.

Hindbrain GLP-1Rs expressed in the AP are accessible to peripherally-dosed liraglutide as shown in Figure 1j. While the exact role of AP GLP-1Rs have not been determined, numerous studies support the role of hindbrain GLP-1 and GLP-1Rs in appetite regulation where they communicate through projections to other sites of the brain, and take part in a complex system integrating homeostatic and likely also hedonic parts of appetite regulation23-25. Secher et al.12 showed that the AP was not required for the weight loss effect of liraglutide. However, as documented in numerous studies, the hindbrain might still be an important site for integration of physiological effects of GLP-1; it is just not required for mediating the effects of pharmacological doses of a GLP-1R agonist on bodyweight loss. Human studies are not available and would be difficult to design.

As also aforementioned, some studies have examined the importance of the vagus nerve in the appetite effect of GLP-1. Acute dosing studies have shown that the vagus nerve is involved in mediating a reduction in food intake in both rodent models and in humans14,26. However, studies in animals with chronic exposure have shown that the vagus nerve is not required12,26. No human studies have used a chronic dosing regime and addressed the importance of the vagus nerve.

The study by Secher et al.12 is the first to show that a peripherally-dosed GLP-1R agonist (liraglutide) can directly access and affect the hypothalamus in the rodent brain. Data in that study showed that liraglutide might have important effects on the CART/POMC neurons that it is shown to access directly. Using the native ligand, GLP-1, GLP-1Rs in CART/POMC neurons are shown to be activated. Using electrophysiological methods, GLP-1 was shown to cause dose-dependent membrane depolarization and an increased firing rate of spontaneous action potentials. Furthermore, the effect was shown to be post-synaptic, in agreement with a potential direct effect of peripherally-dosed liraglutide. Importantly, liraglutide was shown to increase the messenger ribonucleic acid levels of
CART in the ARC, and to keep levels of neuropeptide Y (NPY) and agouti-related peptide (AgRP) at the level of normal control animals, whereas reduced-feed ‘control animals’ weight-matched to those on liraglutide had increased levels of both NPY and AgRP. The effect on AgRP/NPY is suggested to be indirect inhibition of the AgRP/NPY neurons through a local
inhibitory gamma-aminobutyric acid neuron, as AgRP/NPY neurons do not have GLP-1Rs. The hypothalamus is a key region in the brain for integration of appetite signals. The main primary neurons in the ARC are the CART/POMC neurons and the AgRP/NPY neurons. Appetite-inhibiting neurons contain POMC peptides, such as α-melanocyte-stimulating hormone and CART. α-Melanocyte-stimulating hormone acts on melanocortin anoxicenic MC3 and MC4 receptors. Appetite-stimulating neurons in the ARC contain NPY, which acts on the orexigenic Y1 and Y5 receptors; and AgRP, which is an antagonist on MC3/4 receptors. NPY and AgRP are the most important regulatory peptides made in the same neurons in the ARC that both act to increase food intake; similarly important are neurons in ARC that co-express POMC and CART signals, which reduce food intake27–31. The direct access of liraglutide to this part of the brain, along with the uptake in CART/POMC neurons and the effects of increasing levels of CART, and maintenance of the low levels of NPY and AgRP, highlight a likely new pathway of how GLP-1R agonists can regulate appetite. Previously, mainly insulin and leptin have been shown to be transported into the brain, and to have direct effects in the hypothalamus; recent evidence proposes that specialized tanycytes are mediating the uptake through the ME28,32,33. Schaeffer et al.34 recently showed circulating acylated ghrelin to be transported to the ARC and to have direct effects there, much like what is now proposed for GLP-1R agonists, such as liraglutide.

It is likely that the vast distribution of GLP-1Rs in the brain, along with integration of peripheral signals to the hindbrain, makes up a complicated interplay between the hypothalamus and hindbrain structures, including the nucleus tractus solitarius where GLP-1 is produced and which receives vagal afferent inputs. Inputs from higher-order areas of the brain involved in reward and cognition are integrated as well, with resultant effects on meal size and frequency, gut handling of ingested food, and energy expenditure. However, the way native GLP-1 affects postprandial appetite under physiological conditions is different from the way pharmacological exogenous doses of long-acting GLP-1R agonists affect appetite. As such, it is important to discriminate between those studies where GLP-1 or GLP-1R agonists are dosed directly into localized places in the brain, and those where pharmacological doses are injected peripherally. With so many different GLP-1R sites of expression in the brain, it will be very complicated to evaluate the exact importance of each individual site in the total resulting appetite regulation. Use of genetically-engineered mice might be able to show some further details, but then the precise relationship to humans will need to be investigated.

The hedonic pathways of appetite regulation are likely important for GLP-1R agonists. In fact, the only striking difference in GLP-1R expression between rodents and primates in the brain is that there seems to be much more GLP-1R expression in the primate brain in areas involved in hedonic aspects of food intake, such as the amygdala and the bed nucleus of the stria terminalis35. Although it is unclear how all these signals integrate, GLP-1 has been shown to reduce both basal and induced reward, and to change food preference in rodent animal models36–38. Raun et al.36 showed that when rats dosed with liraglutide for 3 months were given a choice between different kinds of candy and chocolate vs normal chow, they selectively chose to eat less chocolate and candy, and more chow. GLP-1R agonists have been shown to lead to taste aversion in rodents, and to nausea in humans; however, in rodents these effects are very short-lived (typically only 1–2 days), so the effect shown by Raun et al.36 is unlikely to be a taste aversion effect. Hansen et al.37 compared liraglutide with sibutramine in a model where

Figure 3 | Neuronal accumulation and activity after glucagon-like peptide-1 receptor stimulation. (a–c) Hypothalamic sections from rats injected with liraglutide594 (red) and stained with Hoechst nuclear stain (blue) and cocaine- and amphetamine-stimulated transcript (green). (b, c) High-magnification confocal images showed accumulation of fluoro liraglutide in the cytoplasm of cocaine- and amphetamine-stimulated transcript-positive cells (arrows). (b) Cocaine- and amphetamine-stimulated transcript and liraglutide594-positive cells. (c) The same image as in (b) with only liraglutide594 signal. Scale bars, 25 μm (b, c); 100 μm (a). © American Society for Clinical Investigation and reproduced from Secher et al.12 with permission.
rats were offered a choice between a chocolate and hazelnut spread (Nutella®; Ferrero SpA, Pino Torinese, Italy)/peanut butter and chow paste/regular chow; both compounds led to a reduction in Nutella/peanut butter paste, but only liraglutide led to an increase in chow intake, as in Raun et al. Nausea is a relatively common side-effect with GLP-1R agonist treatment in humans with and without type 2 diabetes, but also transient and dose-dependent – although it does last longer than in rodents, it is not believed to be the mediator of chronic weight loss. No human studies have yet investigated whether long-term GLP-1R agonist treatment in humans leads to changes in food preferences and/or eating behavior, but studies using functional magnetic resonance imaging scanning have shown an involvement of parts of the brain known for involvement in hedonic aspects of food intake.

Most of the aforementioned areas in the brain, to which liraglutide has direct access, and further areas described in Secher et al., are so-called circumventricular organs; that is, areas of the brain that have no classical blood–brain barrier. The AP, ME, the subfornical organ and the organum vasculosum of the lamina terminalis are circumventricular organs. Although it is logical that a peptide, such as GLP-1, and other drug-like GLP-1R agonists can bind to GLP-1Rs in circumventricular organs, it was perhaps less expected that pharmacological doses of a GLP-1R agonist, such as liraglutide, could reach areas in the brain that are protected by the blood–brain barrier. The PVN and the ARC are well-described areas for GLP-1R action in the brain, but direct access from the periphery has not been described before. GLP-1 and liraglutide have been described to cross the blood–brain barrier, but in light of the very specific binding in select regions as shown here, perhaps those older kinds of studies need to be re-thought as a methodology where the entire brain parenchyma is extracted seems less likely as a method – although it does last longer than in rodents, it is not believed to be the mediator of chronic weight loss. No human studies have yet investigated whether long-term GLP-1R agonist treatment in humans leads to changes in food preferences and/or eating behavior, but studies using functional magnetic resonance imaging scanning have shown an involvement of parts of the brain known for involvement in hedonic aspects of food intake.

In conclusion, new evidence from animal models has highlighted that pharmacological levels of peripherally-dosed GLP-1R agonists can access the hypothalamus directly and have local effects on key primary neurons in ARC, leading to an increase in satiety signals and a decrease in hunger signals. These data highlight a potential new important pathway that likely integrates with several other brain pathways where GLP-1Rs are expressed. Human mechanistic data showed that liraglutide increased feelings of satiety and decreased hunger, leading to an overall reduction of appetite and energy intake. More studies are required to further understand the differences in weight loss efficacy between structurally different GLP-1R agonists.

DISCLOSURE
All authors are full-time employees of, and minor employee-based shareholders in, Novo Nordisk, the company that developed the liraglutide compound.

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