Loss of dorsomedial hypothalamic GLP-1 signaling reduces BAT thermogenesis and increases adiposity

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ABSTRACT

Objective: Glucagon-like peptide-1 (GLP-1) neurons in the hindbrain densely innervate the dorsomedial hypothalamus (DMH), a nucleus strongly implicated in body weight regulation and the sympathetic control of brown adipose tissue (BAT) thermogenesis. Therefore, DMH GLP-1 receptors (GLP-1R) are well placed to regulate energy balance by controlling sympathetic outflow and BAT function.

Methods: We investigate this possibility in adult male rats by using direct administration of GLP-1 (0.5 ug) into the DMH, knocking down DMH GLP-1R mRNA with viral-mediated RNA interference, and by examining the neurochemical phenotype of GLP-1R expressing cells in the DMH using in situ hybridization.

Results: GLP-1 administered into the DMH increased BAT thermogenesis and hepatic triglyceride (TG) mobilization. On the other hand, Glp1r knockdown (KD) in the DMH increased body weight gain and adiposity, with a concomitant reduction in energy expenditure (EE), BAT temperature, and uncoupling protein 1 (UCP1) expression. Moreover, DMH Glp1rKD induced hepatic steatosis, increased plasma TG, and elevated liver specific de-novo lipogenesis, effects that collectively contributed to insulin resistance. Interestingly, DMH Glp1rKD increased neuropeptide Y (NPY) mRNA expression in the DMH. GLP-1R mRNA in the DMH, however, was found in GABAergic not NPY neurons, consistent with a GLP-1R-dependent inhibition of NPY neurons that is mediated by local GABAergic neurons. Finally, DMH Glp1r KD attenuated the anorexigenic effects of the GLP-1 agonist exendin-4, highlighting an important role of DMH GLP-1R signaling in GLP-1-based therapies.

Conclusions: Collectively, our data show that DMH GLP-1R signaling plays a key role for BAT thermogenesis and adiposity.

Keywords Neuropeptide; Hypothalamus; Sympathetic nerve; Adipose tissue; Obesity

1. INTRODUCTION

The intestine and brain both produce glucagon-like peptide-1 (GLP-1), which plays an important role in the control of food intake and glycemia [1,2]. GLP-1 producing neurons are primarily located in the nucleus tractus solitarii (NTS) in the hindbrain, where they integrate neural, hormonal, and visceral sensory information arising from meal ingestion [3]. GLP-1 neurons send projections to numerous brain areas involved in the neuroendocrine and autonomic regulation of energy homeostasis [4–6]. Recent studies have highlighted the role of central GLP-1R in mediating the anorexigenic effects of GLP-1R agonists (e.g., exenatide and liraglutide), which are pharmacological treatment options for Type II diabetes and, more recently, obesity [7–11]. Intracerebroventricular injection of GLP-1 or GLP-1R agonists decreases food intake and increases energy expenditure (EE) [12–14], resulting in weight loss.

The recent emergence of GLP-1’s role in the control of the sympathetic nervous system (SNS) highlights novel mechanisms through which GLP-1 contributes to energy balance [15]. Stimulation of central GLP-1R increases brown adipose tissue (BAT) thermogenesis and causes weight loss in a way that is largely independent of its anorexigenic effects [14,16–19]. Central GLP-1 infusion also decreases de-novo lipogenesis in the liver and hepatic as well as adipose tissue triglyceride content [20,21]. It is unknown, however, whether central GLP-1R-mediated effects on sympathetic regulation of adipose tissue and liver are associated with the pathogenesis of obesity and insulin resistance. A recent study showed that GLP-1R expression is reduced in postmortem hypothalamic nuclei of obese diabetic humans [22], suggesting that reduced hypothalamic GLP-1R signaling may contribute to the development of the metabolic syndrome. The dorsomedial nucleus of the hypothalamus (DMH), a key part of the sympathetic control network [23], expresses GLP-1R and is densely

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innervated by GLP-1 fibers [4,5,24,25], suggesting a possible role of DMH GLP-1R signaling in SNS regulation. DMH neurons send mono-synaptic projections to the rostral raphe pallidus (rRPa), which contains sympathetic premotor neurons regulating BAT activity [26–29]. Dis-inhibiting DMH glutamatergic neurons stimulates BAT sympathetic nerve activity (SNA) and thermogenesis [30,31]. Moreover, the DMH mediates increased blood pressure and heart rate during psychogenic nerve activity (SNA) and thermogenesis [32–34]. DMH leptin receptor (LEPR) expressing neurons have been synaptic and functionally linked to BAT thermogenesis [34–36]. Blocking or ablating DMH LEPR reduced EE and the thermogenic effects of leptin [34,35]. Similar to LEPR in the DMH [37], GLP-1R are expressed in the anterior and ventral part of the DMH [38,39]. However, the neurochemical phenotype and function of GLP-1R expressing neurons are unknown. Given the changes in GLP-1R expression observed in the hypothalamus of obese diabetic humans [22], a decrease in DMH GLP-1R signaling may also play an important role in the pathogenesis of obesity via a SNS-dependent mechanism. Neuropeptide Y (NPY) expressing neurons in the DMH have long been implicated in controlling ingestive behavior and body weight (BW) [39–43]. Knockdown of DMH NPY expression decreases food intake, reduces fat mass, and ameliorates diet-induced obesity (DIO) and insulin resistance [44–46]. Interestingly, DMH NPY knockdown also increases BAT thermogenesis and white adipose tissue (WAT) browning. Figure 1. Schematic diagram of the experiment design. NPY expression decreases food intake, reduces fat mass, and ameliorates diet-induced obesity (DIO) and insulin resistance [44–46]. Interestingly, DMH NPY knockdown also increases BAT thermogenesis and white adipose tissue (WAT) browning. Suggesting that DMH NPY neurons contribute to the regulation of obesity, in part by decreasing sympathetic outflow [47]. However, the mechanisms through DMH NPY neurons reduce sympathetic outflow are poorly understood. Arcuate nucleus (ARC) NPY neurons are inhibited by local GLP-1-sensitive GABAergic neurons [9], raising the possibility that a similar mechanism may contribute to DMH NPY neuronal regulation. Here we investigate the role of DMH GLP-1R signaling in the sympathetic regulation of BAT in male rats by assessing 1) the short-term effects of DMH GLP-1R injections on BAT thermogenesis, 2) the effects of long-term DMH Glp1r knockdown (KD) on BW, EE, BAT thermogenesis, and glucose metabolism, and 3) the neurochemical identity of DMH GLP-1R expressing neurons. We found that one DMH GLP-1R stimulation increased BAT thermogenesis and hepatic fuel mobilization, whereas DMH Glp1r KD increased adiposity, decreased EE, impaired BAT function, and hepatic steatosis; effects which collectively contributed to increased insulin resistance and blunted the anorectic response to the long-lasting GLP-1R agonist, exendin-4 (Ex-4). In addition, DMH Glp1r KD locally increased NPY expression, implicating NPY as a downstream substrate for DMH neuronal regulation of BAT thermogenesis. Collectively, our results show that DMH GLP-1R signaling is a physiologically important component of the central GLP-1 system that controls BAT thermogenesis in a way that helps maintain energy balance.

2. MATERIALS AND METHODS

2.1. Animals
Male Sprague Dawley (SD) rats were purchased from Charles River Laboratories (Sulzfeld, Germany) or Harlan (Envigo US) and housed individually in a climate-controlled room (temperature: 23 ± 2 °C, humidity: 55 ± 5%). Rats were maintained on a 12 h/12 h light/dark cycle with ad libitum access to standard chow diet (No 3436, Provimi Kliba AG, Kaiseraugst, Switzerland) and tap water, except as noted. All procedures were approved by the Cantonal Veterinary Office of Zurich, or the University of Southern California Institutional Animal Care and Use Committee.

2.2. Stereotoxic surgery
SD rats (320–340 g; pre-surgical BW) were anesthetized by intra-peritoneal injection of 2 mg/kg Xylazine (Rompun 2%, Provet AG, Lysach, Switzerland) and 10 mg/kg BW Ketamin (Ketalar 50 mg/ml, Pfizer AG, Zurich, Switzerland), and positioned in the stereotoxic apparatus. Acute DMH GLP-1R activation study: Bilateral guided cannulas (Bilaney, Dusseldorf, Germany) were positioned immediately above the DMH (3.3 mm caudal to the bregma, 1 mm from center to center, 8 mm below pedestal). Sham injections started after 7 days of recovery (injectors fit to 8 mm guided cannula with 1.5 mm projections). On the experiment day, GLP-1 (0.5 ug in 0.5 ul volume, n = 8) or saline (0.5 ul, n = 8) was injected bilaterally 4 h after dark onset. The placement of the cannula was verified during cryosectioning the brains.

DMH Glp1r knockdown: AAV-GFP or AAV GLP-1R shRNA constructs were previously described in detail [48,49]. 200 nl of AAV-GFP or AAV GLP-1R shRNA (5.22 e12 GC/ml) was bilaterally injected into the DMH (3.3 mm caudal, 0.5 mm lateral to the bregma, and 8.4 mm ventral to dura) using a glass capillary micropipette connected with polyethylene (PE)-tubing to a microinjector (Picospritzer III, Parker Hannifin, Hollis, USA). All injected rats (n = 7 for AAV-GFP and n = 10 for AAV GLP-1R) were also implanted with intraperitoneal catheters for subsequent drug injections. Food intake and BW were measured daily. Behavioral and metabolic effects were assessed beginning approximately 3 weeks after surgery. Only the animals with bilateral GFP expression within the DMH were included in the final analyses.

2.3. Indirect calorimetry
Respiratory exchange ratio (RER) and energy expenditure (EE) measurements were conducted in an open circuit calorimetry Phenomaster system (TSE) after 3 days of habituation. TSE Indirect calorimetry estimates EE (kcal/h/kg) from the animals’ O2 consumption (mL/kg/h) and CO2 production (mL/kg/h) using the Weir equation [50,51]. RER: VCO2/VO2: EE: (3.941 × VO2 + 1.106 × VCO2)/1000. For the Glp1rKD study, RER and EE were measured at 8 weeks after surgery. For the acute study, RER and EE were measured before and after intra-DHM GLP-1 (Bachem, 0.5 µg in 0.5 µl saline, n = 8) or vehicle (n = 8) administration for comparison. Data are presented as the average RER and EE values in dark, light, and total diurnal cycle for 3 consecutive days.

2.4. BAT and core temperature measurements
Infra-red (IR) pictures were taken with an E60 camera (FLIR), mounted vertically over the shaved interscapular area (distance 30 cm). Rats were lightly restrained in a stretched position under the camera and 3 snapshots/time point were taken for further analysis. For the acute study, baseline IR pictures were taken prior to intra-DHM vehicle/GLP-1 injection (0.5 µg in 0.5 µl saline) and a second series of IR pictures was taken 4 h after the injection. Body core temperature was measured with a rectal thermometer (WD-35427-20, Oakton instruments) before and 4 h after the injection. For the Glp1rKD study, all rats were subjected to BAT recording 5 h after dark onset at 4 weeks. For the β-3 receptor agonist study, rats had ad lib access to food for 2 h after dark onset. After 1.5 h of fasting, rats received 1 µg/kg IP injection of β-3 receptor agonist (Sigma—Aldrich CL316243) or PBS. IR pictures were taken 2 h after the injection. Final analyses of the IR pictures were done by defining a standard rectangular area (60 × 20 px) over the interscapular region and averaging the mean area temperatures of two pictures (FLIR Tool software for PC).
2.5 Measurement of food intake and body composition

Daily food intake and meal patterns were recorded by an in-house automated system [52]. For the fast-refeeding study, the rats were fasted for 24 h prior to refeeding. Food intake was measured at 0.5, 1, 2, 4, and 6 h after dark phase onset. Analysis of body composition was conducted using a computerized tomography system (La Theta LCT-100, Aloka) with a previously validated method [53]. Briefly, rats (AAV GLP-1R n = 10 and AAV-GFP n = 7) were anesthetized with isoflurane and placed supine in a cylindrical holder (120 mm inner diameter). An initial whole-body sagittal image was obtained to ensure proper placement of the animal. Scans were done from vertebrae L1 to L6 with a 2 mm pitch size. The Aloka software detects volumes of adipose tissue (fat mass), bone, air, and the remainder (lean mass) based on their X-ray absorption. Fat mass and lean mass were computed using the density factors of 0.92 g/cm³ and 1.10 g/cm³, respectively. Fat ratio is defined as fat mass/(lean + fat mass).

2.6 Intraperitoneal glucose tolerance test (IPGTT)

All rats (AAV GLP-1R n = 10 and AAV-GFP, n = 7) were fasted overnight and injected with 2 g/kg glucose (IP). Tail vein blood was collected at 0, 15, 30, 60, 90, and 120 min after the glucose bolus injection.

2.7 Systemic Exendin-4 effects on FI, BW, and EE

On the day of the experiment, all rats (AAV GLP-1R n = 10 and AAV-GFP n = 7) were fasted for 4 h and received either Ex-4 (1.0 μg/kg, IP) or saline at dark cycle onset. RER and EE measurements were conducted in an open circuit calorimeter Phenomaster system (TSE) for 24 h after drug injection. Food intake and BW were measured at 24 h after drug injection.

2.8 Tissue morphology

For stainings with hematoxylin and eosin (H&E) and oil red O (ORO) adipose and liver tissues were fixed in 4% PFA (Sigma—Aldrich) and processed in a STP 120 (Microm) according to the standard protocol. Paraffin-embedded samples were cut at 5 μm on a Hyrax M55 (Zeiss), deparaffinized and stained on a Vastarin 24-4 (Shandon) for adipose tissue or ORO (Sigma—Aldrich, cat # 00625) for liver [54]. Pictures were obtained with an Axioscope A.1 microscope (Zeiss). H&E and ORO images were quantified for lipid fraction and ORO area using ImageJ software.

2.9 Plasma analysis

Triglycerides, FFA, glycerol, and cholesterol levels were analyzed by Cobas MIRA analyzer [63] (Hoffman LaRoche, Basel, Switzerland). Insulin and leptin levels were measured by using a Multi-Array Mouse/Rat Insulin and leptin kit (Meso Scale Discovery, Gaithersburg, USA).

2.10 Gene expression analyses

For the gene expression analyses in the acute DMH-GLP-1R activation study, rats were decapitated 4 h after the DMH injections of GLP-1 (0.5 μg in 0.5 μl saline) or vehicle. Brains and BAT were collected, frozen in dry ice and stored at −80 °C for RNA extraction. For the KD study, brains, liver, BAT, and WAT were collected, frozen in dry ice and stored at −80 °C for RNA and protein extraction. DMH, PVH, and ARC were micropunched using anatomical landmarks and RNA was extracted using Trizol (Life Technologies) according to the manufacturer’s protocol. RT-qPCR was performed using SYBR green on a OneStep Plus Real Time PCR instrument (Applied Biosystems) and results were analyzed using the 2ddCt method (all primer sequences available upon request).

2.11 Western blot analyses

Western blots were performed using a protocol previously described in detail [64]. Briefly, proteins were separated by 8—10% SDS-polyacrylamide gel polyacrylamide electrophoresis and transferred onto nitrocellulose membranes (pore size 0.2 μm; Protran, Whatman). Membranes were then blocked in 5% skim milk in Tris-buffered saline containing 0.1% Tween-20 for 1 h before incubating with the primary antibodies overnight at 4 °C. On the next day, the membranes were washed 3 times with Tween-20, and incubated with secondary horseradish peroxidase-conjugated anti-rabbit or anti-mouse IgG, respectively, for 1 h at RT. Protein bands were visualized using a chemiluminescent substrate and the intensities of the bands were quantified by densitometry using Image J software (ImageJ 1.46b).

Primary antibodies: Uncoupling protein 1 (UCP1; Rabbit antibody PA1-24894 1:2000, ThermoFisher), ACC (Rabbit antibody 3662, 1:1000, Cell Signaling), pACC (Rabbit antibody 3661, 1:1000, Cell Signaling), FAS (Rabbit antibody 3180, 1:1000, Cell Signaling), β-actin (Mouse antibody AC-74, 1:5000, Sigma—Aldrich), γ-tubulin (Mouse antibody T6557, 1:5000, Sigma—Aldrich).

2.12 In situ hybridization

2.12.1 Animals for in situ hybridization

Five male Sprague—Dawley rats (BW 275—300 g) were anesthetized and perfused through the ascending aorta with ice-cold 4% paraformaldehyde in 0.1 M sodium tetraborate (pH 9.5) as previously described [55]. Brains were removed and post-fixed for 24 h in the borate fixative containing 12% sucrose, frozen in hexanes cooled with dry ice, and then stored at −70 °C. Five series of one in five, 20 μm thick, frozen coronal sections were cut through the hypothalamus in the same approximate plane as the Swanson Rat Brain Atlas [56] and mounted on Superfrost Plus slides the same day, stored overnight under vacuum and then desiccated at −70 °C. One series was stained with thionin for local cytoarchitectonics.

We used two in situ hybridization (ISH) procedures to assess how GLP-1 mRNA was related to other key neuronal mRNAs in the DMH. First, single 35S cRNA ISH was used to survey the rostrocaudal distribution of VGlut2, GAD 65, NPY, and GLP-1 mRNA. To determine the chemical identity of DMH neurons containing GLP-1 mRNA we then used two double ISH methods: a 35S-labeled and digoxigenin fluorescence ISH (FISH) modified from Watts & Sanchez-Watts [57], and Advanced Cell Diagnostics RNAscope® Multiplex FISH [58].

Antisense cRNA probes for VGlut2 [59], GAD 65 [60], NPY [61], and GLP-1R [62] mRNAs were transcribed using either 35S-UTP or a Digoxigenin RNA labeling mix (Roche) as previously described with minor modifications [57].

**Combined 35S- and digoxigenin-labeled (ISH/FISH):** An optimized mixture of either 35S-labeled NPY or GAD65 cRNA probe was cotraded with the Dig-labeled GLP-1R cRNA probe and used for ISH as previously described with minor modifications [57]. Slide-mounted sections were exposed to Agfa Mamaray HDR-C film for 4 days to estimate the exposure time required for emulsion autoradiography for the 35S hybridization. To visualize the digoxigenin-RNA the same slides were processed with the HNPP Fluorescent Detection Kit (Roche) modified from the manufacturer’s protocol. Slides were incubated in blocking buffer containing 0.05% Triton-100/2% Normal Sheep Serum/2X SSC for 2.5 h at 30 °C, rinsed in Digoxigenin Buffer 1.
(100 mM Tri-HCl/150 mM NaCl pH 7.5) and then incubated overnight at 4 °C in Anti-Digoxigenin-AP Fab fragments diluted 1:3 K with Digoxigenin Buffer 1/0.3% Triton-100/1% Normal Sheep Serum, and then washed in Digoxigenin Buffer 3 (Washing Buffer: 0.05% Tween 20/Digoxigenin Buffer 1) followed by Digoxigenin Buffer 4 (Detection Buffer: 100 mM Tris- HCl/100 mM NaCl/10 mM MgCl2 pH 8). To reveal a specific fluorescent signal, a Fast Red TR solution was prepared first at a concentration of 25 mg/ml in redistilled water. The final mixture contained 10 μl Fast Red solution and 10 μl HPDP per ml of Digoxigenin Buffer 4 (Detection Buffer pH 8). The HPDP/Fast Red TR mixture was filtered through a 0.2 μm nylon syringe filter immediately before use. 10 μl of filtered HPDP/Fast Red TR mix was applied to each slide, coverslipped and incubated in the dark at 25 °C for 30 min then washed in Digoxigenin Buffer 3. These two steps were repeated 3 times before slides were rinsed in distilled water and air dried in the dark.

To visualize 35S-labeled cRNA hybridization, slides were dipped in Kodak autoradiography emulsion type NTB (diluted 1:1 with distilled water), exposed for 4 days (GAD65) and 12 days (NPY), developed with Kodak autoradiography emulsion type NTB (diluted 1:1 with distilled water), and then coverslipped with DABCO.

To visualize fluorescent (RNAscope®) in situ hybridization (FISH) for GLP-1R, NPY, GAD65, and VGlut2 mRNAs: Slide-mounted sections from the same animals used for the combined 35S- and digoxigenin-labeled (ISH/FISH) were processed to detect the mRNAs for GLP-1R (ACD 315221), NPY (ACD 435801-C2), and VGlut2 (ACD 317011-C2) using RNAscope® Multiplex FISH, modified from Hsu et al. [58].

2.12.3. Image analysis
Sections were photographed with a Zeiss Axioimager Z1 epifluorescence microscope through a 5 × or 10 × plan-apochromat objective lenses. Dark field illumination to visualize silver grains was provided by a Microvid lateral illumination system (Micro Video Inc., Avon MA). Fluorescence and dark field images of the same field were captured with a Hamamatsu Orca ER monochromatic digital camera controlled with Velocity 6.1 software (Perkin Elmer, Waltham, MA). All photo-images were then imported into Adobe Photoshop (CS3; Adobe Systems, Inc., San Jose, CA), and the fluorescence and dark field images were merged using separate Photoshop layers. All figure panels were assembled in Adobe Illustrator (CS3) to identify double-labeled neurons.

2.13. Statistical analyses
Data were analyzed by a Student’s t-test for unpaired normally distributed values of equal variance using GraphPad Prism (version 6.05 for Windows). Where the dependent variable was affected by two factors, the data were analyzed with a 2-way ANOVA. When the main effect—or interaction terms were significant, post-hoc analyses using the Bonferroni correction were performed. Data are presented as means ± SEM. P-values < 0.05 were considered significant. All graphs were generated using GraphPad Prism (version 6.05 for Windows).

3. RESULTS
3.1. Acute DMH GLP-1R stimulation increases BAT thermogenesis
We injected GLP-1 (0.5 μg) bilaterally into the DMH and measured BAT temperature under a special feeding schedule to standardize food intake (Figure 1A), thereby isolating the effects of GLP-1R activation on BAT thermogenesis. DMH GLP-1R stimulation increased interscapular BAT temperature (Figure 1B,C, p < 0.001) and core body temperature (Figure 1D, p < 0.05) at 4 h after injection. In support of increased BAT temperature, mRNA expression of β3-adrenergic receptor (ADRB3, p < 0.05), peroxisome proliferator-activated receptor gamma coactivator 1-alpha (PGC1α, p < 0.05), peroxisome proliferator-activated receptor gamma (PPARγ, p < 0.05), and cell death-inducing DFFA-like effector A (CIDEA, p < 0.05) was up-regulated in the BAT at 4 h after GLP-1 injection (Figure 1E). There was, however, no significant change in uncoupling protein 1 (UCP1) mRNA expression in the BAT. DMH GLP-1 injection also increased respiratory exchange ratio (RER, Figure 1F, p < 0.05) but did not affect EE (S1A). BAT thermogenesis requires fatty acids as fuel for heat generation [65]. Interestingly, acute DMH GLP-1R stimulation increased TG levels in the plasma (Figure 1G, p < 0.05) as well as liver fatty acid synthase (FAS) gene expression (Figure 1H, p < 0.05). Free fatty acids (FFA), glycerol, and β-hydroxybutyrate (BHB) levels were unchanged in the plasma of GLP-1 injected rats (S1B). DMH GLP-1R stimulation led to a decrease in NPY (p < 0.05) mRNA expression, but did not significantly affect GLP-1R and coin- amphetamine-regulated transcript (CART) mRNA expression (Figure 1I), raising the possibility that inhibition of NPY gene expression by GLP-1 is involved in the sympathetic control of BAT thermogenesis.

3.2. Long term DMH Glp1r KD increases adiposity
To further examine the role of DMH GLP-1R in BAT homeostasis, we generated a rat model of Glp1r KD specifically in the DMH using viral-mediated RNA interference (Figure 2A). Rats injected with AAV-GLP-1R shRNA showed a significant reduction (51%) of GLP-1R mRNA expression (Figure 2B, p < 0.001) in the DMH compared to rats injected with AAV-Control shRNA expressing GFP only. GLP-1R mRNA expression in the ARC and PVH was not affected by viral injection (Figure 2B). We also found a partial reduction of GLP-1R protein expression in the DMH (S2A, p = 0.05). Starting at 7 weeks after viral infection, AAV-GLP-1R rats displayed a small, but significant increase in BW gain compared to controls (Figure 2C, p < 0.05, BW in S2B). This BW gain was accompanied by an increase in fat mass (Figure 2D, p < 0.01) and an upward trend in fasting leptin levels (Figure 2E, p = 0.09). Food intake was not different between the two groups during the first 4 weeks post injection (Figure 2F). Despite the lack of a change in initial daily food intake, AAV-GLP-1R rats displayed an abnormal meal pattern, i.e., a decrease in meal size and an increase in meal frequency during the dark phase (S2C). Furthermore, AAV-GLP-1R rats reduced refeeding response (Figure 2G, p < 0.001), and BW change (Figure 2H, p < 0.05) to 24 h fasting. In addition, DMH Glp1r KD reduced adrenal gland weight, an indicator of altered sympathetic activity (S3A).

3.3. Long term DMH Glp1r KD decreases EE, BAT temperature, and UCP1 expression
Consistent with increased adiposity, DMH Glp1r KD rats exhibited reduced EE during the dark phase, but not during the light phase (Figure 3A,B, p < 0.001). On the other hand, RER was not different between AAV-GLP-1R and AAV-Control rats (Figure 3C,D). BAT temperature was significantly lower in AAV-GLP-1R rats compared to the controls (Figure 3E, p < 0.05), indicating low thermogenic capacity as a result of long term Glp1r KD. Core temperature was however not different between the two groups (Figure 3F). β-3 adrenergic receptor stimulation by CL- 316243 (0.1 mg/kg, SC) increased BAT temperature in the controls (Figure 3G, p < 0.005), but not in AAV-GLP-1R rats. Consistently, adipocytes were bigger in the BAT of AAV GLP-1R rats than in controls (Figure 3H, p < 0.05), indicating lipid accumulation in the cells. We hypothesized that DMH Glp1r KD blunts the stimulatory effect of β-3 adrenergic receptor agonist due to chronic reduction in
SNS outflow to brown adipocytes supressing the cellular machineries critical for therogenesis. In fact, AAV-GLP-1R rats had lower levels of UCP1 mRNA (Figure 3I, P < 0.005) and protein (Figure 3J, P < 0.01) in BAT, together with lower expression levels of other thermogenic genes including PGC1α (p < 0.05) and PPARγ (p < 0.05) in the BAT(Figure 3I). Together, these data show that reduced EE and BAT thermogenesis contributed to increased adiposity in DMH Glp1r KD rats.

3.4. Long term DMH Glp1r KD impairs lipid and glucose metabolism

Consistent with increased adiposity, plasma triglyceride levels were higher in AAV-GLP-1R rats compared to controls (Figure 4A, p < 0.005), but cholesterol, free fatty acid, and glycerol levels were comparable between the two groups (S3B–D). Oil red O staining revealed lipid accumulation in the livers of AAV-GLP-1R rats (Figure 4B, p < 0.001), further supported by the activation of de novo lipogenesis genes (Figure 4C). DMH Glp1r KD resulted in increased fatty acid synthase (FAS, p < 0.05) and a tendency of increased acetyl-CoA carboxylase (ACC, p = 0.06) gene expression. Similarly, FAS and ACC protein levels were higher in AAV-GLP-1R rats (Figure 4D, p < 0.05). pACC protein expression and pACC/ACC ratio were however unchanged. On the other hand, DMH Glp1r KD decreased glucose 6-phosphatase (G6Pase, p < 0.005) and phosphoenolpyruvate carboxykinase 1 (PEPCK1, p < 0.01), indicating decreased gluconeogenesis, presumably due to increased insulin production (Figure 4C). The expression of fibroblast growth factor 21 (FGF21) and PGC1α in the liver was similar in the two groups (Figure 4C). Unlike its expression in liver, FAS mRNA expression was down-regulated in the subcutaneous white adipose tissue (scWAT, Figure 4 E, p < 0.05). There was no change in the mRNA expression of lipolytic enzymes, adipose triglyceride lipase (ATGL) and hormone-sensitive lipase (HSL), in the scWAT of AAV-GLP-1R rats. Although DMH Glp1r KD did not affect the glucose profile during an intraperitoneal glucose tolerance test (IPGTT) at 7 weeks post viral injection (Figure 4 F), insulin secretion in response to the glucose challenge was greater in AAV-GLP-1R rats than in controls (Figure 4G, p < 0.05), suggesting that AAV-GLP-1R rats are more insulin resistant. Because BW gain was higher in AAV-GLP-1R rats, this decrease in insulin sensitivity may be secondary to
the change in BW gain rather than a direct effect of KD. In addition, the fasting insulin level was higher in AAV-GLP-1R rats compared to the controls (Figure 4H, p < 0.05). Together, these results indicate that Glp1r KD in the DMH alters lipid metabolism in liver and adipose tissue, which is associated with insulin resistance.

3.5. Long term DMH Glp1r KD increases DMH NPY mRNA expression

In contrast to DMH GLP-1R stimulation, DMH Glp1r KD increased NPY mRNA expression in the DMH (Figure 5A, p < 0.001), providing further support for the regulation of NPY transcription by GLP-1R signaling. The leptin receptor (LEPR) in the DMH also regulates BAT thermogenesis [36]. We did, however, not detect a change in either LEPR, the cholecystokinin receptor (CCKR), or CART expression in the DMH of AAV-GLP-1R rats. Because DMH NPY neurons send major projections to the PVH [28], it is possible that NPY regulates BAT thermogenesis via modulating gene expression in the PVH [66]. While there were no statistically significant changes in thyrotropin releasing hormone (TRH) and CART mRNA levels after knocking down DMH GLP-1R mRNA, there were trends towards changes in corticosterone releasing hormone (CRH) and tyrosine hydroxylase (TH, p = 0.07) mRNAs (Figure 5B). Interpreting these results conservatively therefore leaves open the possibility that DMH NPY projections to the PVH can in some way influence the autonomic and neuroendocrine regulation of energy balance. Levels of proopiomelanocortin (POMC), agouti-related protein (AgRP), NPY, and CART mRNAs were not different between the two groups in the ARC (Figure 5C).

3.6. DMH-GLP-1 neurons express GAD65 but not NPY mRNAs

To determine the neurotransmitter identity of the DMH neurons containing GLP-1R mRNA, we used two in situ hybridization techniques that can detect multiple mRNAs simultaneously. GLP-1R mRNA-containing neurons were found throughout the rostrocaudal extent of the DMH [54].
They were however conspicuously absent from the compact subdivision of DMH (DMHp), which contains the majority of DMH NPY neurons (S4). We found less than 5% of GLP-1R mRNA expressing neurons contained NPY mRNA in the DMH (Figure 5D, S5A), strongly supporting an indirect way ANOVA; \( *p < 0.0001 \). (C) RER over 24 h in AAV-control and AAV-GLP-1R rats. (D) Average RER in dark and light phases. (E) BAT temperature during the dark phase (n = 7/8; Student t-test; \( *p < 0.01 \)). (F) Rectal temperature during the dark phase. (G) BAT temperature change 2 h after \( \beta \)-3 receptor agonist CL316243 injection (1 \( \mu g/kg \) i.p.; n = 3–5; Two-way ANOVA; \( *p < 0.005 \)). (H) Representative pictures of H&E staining (Scale bar: 100 \( \mu m \)) and lipid area fraction in BAT of AAV-control and AAV-GLP-1R rats (n = 6/5; Student t-test; \( *p < 0.05 \)). (J) Representative pictures of H&E staining (Scale bar: 100 \( \mu m \)).

3.7. DMH Gp1r KD attenuates the anorexigenic effects of peripheral GLP-1R agonist treatment

The anorexigenic effects of peripherally administered agonists are thought to be mediated, in part, by GLP-1R expressing neurons in the brain [8,9,11]. To test whether reduced DMH GLP-1R expression influences the anorexigenic effects of a peripheral GLP-1R agonist, we injected Exendin-4 (Ex-4, 1 \( \mu g/kg \), IP) and measured BW, food intake, EE and RER. Ex-4 decreased 24 h food intake in the controls (\( p < 0.05 \), but this effect was attenuated in AAV-GLP-1R rats (Figure 6A). It is important to mention that the baseline food intake was reduced in AAV-GLP-1R rats after 7 weeks, likely compromising Ex-4’s ability to further suppress food intake. Similarly, Ex-4-induced 24 h BW loss was attenuated in AAV-GLP-1R rats compared to the controls (Figure 6B, \( p < 0.005 \)). In contrast to the stimulatory effects of central GLP-1 or GLP-1 analogs on EE [67,68], peripheral GLP-1/Ex-4 administration decreases RER and diet-induced thermogenesis in rodents and humans [12,69], suggesting that the activation of peripheral or central GLP-1R has opposite effects on EE. Whereas Ex-4 treatment at dark phase onset decreased EE in the controls during dark phase (\( p < 0.005 \)), it did not further decrease EE in AAV-GLP-1R rats that are in a metabolically compromised state (Figure 6C,D; S6A). Ex-4 treatment also decreased RER in the controls during the dark phase (\( p < 0.001 \), but this effect was attenuated in AAV-GLP-1R rats (Figure 6E,F; S6B). Our data demonstrate that DMH GLP-1R expression is necessary for normal EE, and that peripheral Ex-4 treatment fails to decrease food intake and BW in the state of low metabolic efficiency.
4. DISCUSSION

GLP-1 producing neurons send extensive projections to brain areas involved in the regulation of energy balance [6]. GLP-1 neurons affect food intake and EE primarily by modulating forebrain neural circuitries [16,70–72]. The DMH, a major hypothalamic site receiving GLP-1 projections, is well known for modulating pre-autonomic neurons involved in BAT thermogenesis, HPA-axis activation, and blood pressure [32,73,74]. Here we show that acute GLP-1R stimulation in the DMH increases BAT thermogenesis, whereas viral-mediated Glp1r KD in the
DMH increases adiposity, hepatic steatosis and insulin resistance. In support of obese phenotypes, DMH Glp1r KD reduced EE and BAT thermogenesis, with a concomitant increase in DMH NPY mRNA expression. Furthermore, the GLP-1R agonist Ex-4 failed to elicit its full anorexigenic and metabolic effects in DMH Glp1r KD rats, raising the possibility that increased NPY in the DMH and low sympathetic tone reduces the efficacy of GLP-1 drug treatment in the obese diabetic state.

A recent report attempted to dissect the role of various hypothalamic GLP-1R expressing neurons by deleting GLP-1R using a cre-lox recombination technology [10]. Surprisingly, their partial knockdown of GLP-1R mRNA in whole hypothalamus, PVH, or specifically in POMC neurons produced inconsistent and very modest effects on BW and EE. Therefore, to avoid developmental compensation of embryonic gene deletion, more time-controlled and region-specific studies targeting different GLP-1R populations in normally-developed adult models are clearly needed. To this end, we used acute pharmacological stimulation and chronic viral-mediated reduction of GLP-1R mRNA in rats to reveal a novel physiological role for GLP-1R signaling effects on DMH neurons known to regulate BAT sympathetic activity.

Central GLP-1R agonism increases EE and BAT thermogenesis [16,18]. In particular, administration of the GLP-1R agonist liraglutide into the ventromedial hypothalamus stimulates BAT thermogenesis and adipocyte browning via hypothalamic AMPK activity [16]. In the same study, DMH GLP-1R stimulation by liraglutide did not alter food intake, BW, or BAT UCP1 expression. In contrast, we demonstrate that native GLP-1 injection into the DMH increases BAT activity and core temperature, as well as BAT thermogenic gene expression. This discrepancy may reflect signaling differences between native GLP-1 and liraglutide, causing differential cellular and behavioral responses. Some evidence suggests that GLP-1 analogs can differentially influence pathway specificity of G-protein coupled receptors [75,76]. Consistent with the effects of acute GLP-1R stimulation, DMH Glp1r KD decreased EE, BAT temperature and thermogenic markers. Therefore, it is reasonable to conclude that DMH GLP-1R signaling profoundly affects BAT thermogenesis, presumably by modulating sympathetic nerve activity.

The role of BAT in the regulation of BW is controversial in both humans and animal models [77]. There are many confounding factors including ambient temperature, diet, and body composition, making it difficult to isolate the role of BAT in BW regulation. In fact, UCP1 deficient mice are resistant to obesity unless they are housed in thermoneutrality [78]. Because we housed our animals at a lower temperature, we speculate that the small BW difference developed over several weeks after the KD, and this BW phenotype may be secondary to chronic reduction in BAT function/fuel utilization rather than a direct effect of DMH Glp1r KD. Future studies will be necessary to compare the BW effect of DMH Glp1r KD in thermoneutrality.

The hypothalamus plays an important role in TG metabolism via sympathetic regulation of BAT, liver, and WAT [15]. Evidence shows that GLP-1 can in part affect TG metabolism via a central mechanism [20,21]. Intriguingly, central GLP-1R agonists increase TG and glucose uptake in the BAT and WAT [18]. Our data show that the DMH is a critical site for GLP-1R-mediated stimulation of BAT thermogenesis, together with hepatic TG production and secretion, which provide additional fuel for BAT heat generation [65]. Central GLP-1R stimulation may also affect hepatic TG production via increased insulin secretion.

Figure 5: DMH GLP-1R signaling regulates NPY gene expression indirectly. (A) Relative mRNA expression in the DMH in AAV-control and AAV-GLP-1R rats: n = 6/9; Student t-test; *p < 0.0005 for NPY. ns: CART, LEPR, and CCKR. (B) Relative mRNA expression in the PVH. ns: CRH, TRH (p = 0.07), and CART. (C) Relative mRNA expression in the ARC. ns: NPY, CART, POMC, and AgRP. (D) ISH/FISH for GLP-1R and NPY mRNA. Top (DMH) and bottom (ARC) panels: Left-GLP-1R mRNA (red), Middle-GLP-1R (red) and NPY (green), Right-co-localization in yellow. (E) ISH/FISH for GLP-1R and GAD-65 mRNA. Top (DMH) and bottom (ARC) panels: Left-GLP-1R mRNA (red), Middle-GLP-1R (red) and GAD-65 (green), Right-co-localization in yellow. Scale bar for all panels = 200 µm. Data are mean ± SEM.
Insulin increases hepatic de novo lipogenesis, suppresses fatty acid oxidation, and promotes TG esterification and secretion [79]. We however did not measure plasma insulin after intra DMH GLP-1 injection: therefore, this possibility requires further testing. Intriguingly, long-term DMH Glp1r KD also increased hepatic fatty acid synthesis and plasma TG levels, which was associated with increased lipid accumulation in the BAT and hyperinsulinemia. It is widely accepted that defective adaptive thermogenesis contributes to the metabolic syndrome including hepatic steatosis and insulin resistance [81]. Therefore, reduced BAT thermogenesis following long-term DMH Glp1r KD likely contributes to the high levels of plasma TG and fat accumulation in the liver.

Recent studies have revealed different DMH neuronal populations linked to BAT thermogenesis [36,44,82–84]. Leptin’s thermogenic function awaits investigation and could be crucial for understanding thermogenic effects in DMH Glp1r KD. Further studies are needed to determine the role of leptin in regulating thermogenesis in DMH Glp1r KD and whether leptin regulates the expression of thermogenic genes in the BAT.

Figure 6: Long term DMH Glp1r KD attenuates the anorexigenic response to Ex-4. (A) 24 h food intake after PBS or Ex-4 (1.0 μg/kg, i.p.) treatment in AAV-control and AAV-GLP-1R rats (n = 7/9; Two-way ANOVA; drug effect * p < 0.001, group effect #p < 0.05). (B) 24 h BW change after PBS or Ex-4 treatment in AAV-control and AAV-GLP-1R rats (n = 7/9; Two-way ANOVA; drug effect * p < 0.0005). (C) EE over 24 h after PBS or Ex-4 treatment in AAV-control and AAV-GLP-1R rats. (D) Average EE in dark phase after PBS or Ex-4 treatment in AAV-control and AAV-GLP-1R rats. (E) Average RER in dark phase after PBS or Ex-4 treatment in AAV-control and AAV-GLP-1R rats (n = 7/9; Two-way ANOVA; drug effect p < 0.01, group effect p < 0.0005, *p < 0.005). Data are mean ± SEM.
while cholinergic DMH neurons decrease BAT activity by modulating serotonergic neurons in the rostral raphe pallidus (rRPa) [83]. We now show that DMH NPY mRNA levels are inversely correlated with GLP-1R signaling, which makes NPY a potential downstream substrate for sympathetic regulation of BAT thermogenesis. DMH NPY neurons project to the PVH, where they inhibit the sympathetic drive of BAT [28,85]. More specifically, NPY release in the PVH modulates presynaptic neurotransmitter release and, hence, strongly inhibits the sympathetic stimulation of BAT [86]. Reducing DMH-NPY expression increased UCP1 expression in BAT and promoted the browning of WAT [44]. On the other hand, over-expressing NPY mRNA in the DMH decreased BAT UCP1 expression and caused insulin resistance [87]. Given the role of DMH NPY neurons in BAT thermogenesis, elevated NPY is well placed to be a key contributor to the low BAT activity seen in DMH Glp1r KD rats.

Another recent study demonstrated that DMH NPY mRNA knockdown suppresses hepatic glucose production and increases insulin sensitivity via hepatic vagal innervation [45]. Our data also support the role of DMH NPY expression in hepatic glucose metabolism. DMH NPY overexpression induced by Glp1r KD was associated with the development of hepatic steatosis and whole body insulin resistance. However, PEPCk and G6Pase gene expression was down-regulated in our DMH Glp1r KD model, suggesting a differential mechanism contributing to insulin resistance. Although we cannot rule out the possibility that Glp1r KD directly affected hepatic glucose metabolism via an NPY-dependent pathway, defective BAT function may be the primary underlying cause of energy imbalance and insulin resistance as its normal function is important for the clearance of glucose and triglycerides from the circulation [88,89]. Further studies using BAT or hepatic sympathetic denervation techniques are necessary to implicate the role of DMH GLP-1R signaling in hepatic glucose and lipid metabolism.

The mechanism by which DMH GLP-1R signaling regulates NPY expression in the DMH remains unclear. Previous studies have shown that peripheral Ex-4 treatment decreases ARC NPY expression [90,91]. Moreover, GLP-1 treatment inhibited the electrical activity of ARC NPY neurons, which required GABAergic presynaptic input [9]. This is further supported by our observation that GLP-1R mRNA is not found in ARC NPY neurons, and by the lack of labeled GLP-1 uptake by NPY neurons in the ARC [9]. Although ARC and DMH NPY expression is differentially regulated by fasting and diet-induced obesity [92,93], a similar mechanism may play a role in regulating NPY expression in the DMH. Indeed, using two ISH techniques, we unequivocally demonstrate that DMH GLP-1R neurons do not express NPY, which means that GLP-1R signaling must regulate NPY expression indirectly. The fact that many DMH GLP-1R neurons are GABAergic supports a local inhibitory mechanism used by GLP-1 signaling to suppress NPY expression. However, based on anatomical evidence, we cannot exclude the possibility that a combined action of GABAergic and glutamatergic GLP-1R expressing neurons in the DMH modulates NPY expression and consequently sympathetic output to BAT by way of the PVH or the rRPa. Further studies are needed to dissect the DMH GLP-1R signaling influences on the neural substrates and pathways that lead to BAT thermogenesis. Finally, we note that our methods were not designed to account for all possible chemical phenotypes that express GLP-1R mRNA. Other studies have shown that DMH neurons express a range of neuropeptides and metabolically important receptors, including the leptin receptor [36], orexin [94], CART [95], and prolactin—releasing peptide [82]. This means that GLP-1R mRNA is found in a diverse but as yet uncharacterized set of DMH neurons.

Liraglutide, a GLP-1R agonist, has recently been approved for obesity treatment [96], and the mechanisms of its anorexigenic effects are still under active investigation. Because central GLP-1R expressing neurons are the key mediators for the anorexigenic effects of peripheral GLP-1R agonists [8,9,11], reduced hypothalamic GLP-1R expression and activity could explain the decreased sensitivity to GLP-1R agonists seen in obese diabetic conditions [97,98]. While ARC GLP-1R expressing cells are thought to mediate the anorexigenic effects of liraglutide [9], bilateral GLP-1 infusion into the ARC did not affect food intake [99]. Moreover, a study employing a POMC specific GLP-1R KO showed normal BW and food intake phenotypes during standard chow feeding [10]. Therefore, the exact role of ARC GLP-1R in food intake is still unclear. Our data showed that altered DMH GLP-1R expression does not affect daily food intake and the reduced refeeding response to overnight fasting presumably reflects a lower metabolic efficiency due to compromised BAT function. Therefore, our Glp1r KD model provides a valuable tool for understanding the clinical significance of reduced DMH GLP-1R signaling that appears to be critical for BAT regulation. BAT depots are present in adult humans where their thermogenic properties are metabolically beneficial [100,101]. Based on our findings following DMH Glp1r KD in rats, the attenuated anorexigenic response to peripheral Ex-4 treatment is correlated with reduced BAT thermogenic capacity. This means that obese diabetic patients with reduced BAT function may exhibit a central GLP-1 resistance that limits the effectiveness of GLP-1R agonist treatment as a weight loss strategy, although this hypothesis requires further testing.

5. CONCLUSIONS

Central GLP-1 relays peripheral nutrient and hormonal signals to the hypothalamus and other energy balance-relevant nuclei distributed across the neuraxis [3,6,25]. It is therefore imperative to clarify how hypothalamic GLP-1R signaling regulates energy balance and how its impairment contributes to the pathophysiology of metabolic syndrome and the efficacy of GLP-1 therapy. Our results point to an important link between hindbrain GLP-1 projections into the DMH and NPY-mediated sympathetic regulation of BAT thermogenesis and energy balance. In light of our findings, manipulating central GLP-1 production and receptor activation in a site-specific manner may hold the key to developing future therapeutic strategies for metabolic syndrome.

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CONFLICT OF INTEREST

None declared.
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