CART neurons in the arcuate nucleus and lateral hypothalamic area exert differential controls on energy homeostasis

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ABSTRACT

Objective: The cocaine- and amphetamine-regulated transcript (CART) codes for a pivotal neuropeptide important in the control of appetite and energy homeostasis. However, limited understanding exists for the defined effector sites underlying CART function, as discrepant effects of central CART administration have been reported.

Methods: By combining Cart-cre knock-in mice with a Cart adeno-associated viral vector designed using the flip-excision switch (AAV-FLEX) technology, specific reintroduction or overexpression of CART selectively in CART neurons in the arcuate nucleus (Arc) and lateral hypothalamic area (LHA), respectively, was achieved. The effects on energy homeostasis control were investigated.

Results: Here we show that CART neuron-specific reintroduction of CART into the Arc and LHA leads to distinct effects on energy homeostasis control. Specifically, CART reintroduction into the Arc of otherwise CART-deficient Cart e−/− mice markedly decreased fat mass and body weight, whereas CART reintroduction into the LHA caused significant fat mass gain and lean mass loss, but overall unaltered body weight. The reduced adiposity in Arc CART; Cart e−/− mice was associated with an increase in both energy expenditure and physical activity, along with significantly decreased Npy mRNA levels in the Arc but with no change in food consumption. Distinctively, the elevated fat mass in LHA CART; Cart e−/− mice was accompanied by diminished insulin responsiveness and glucose tolerance, greater spontaneous food intake, and reduced energy expenditure, which is consistent with the observed decrease of brown adipose tissue temperature. This is also in line with significantly reduced tyrosine hydroxylase (Th) and notably increased corticotropin-releasing hormone (Crh) mRNA expressions in the paraventricular nucleus (PVN).

Conclusions: Taken together, these results identify catabolic and anabolic effects of CART in the Arc and LHA, respectively, demonstrating for the first time the distinct and region-specific functions of CART in controlling feeding and energy homeostasis.

Keywords CART; Energy homeostasis; AAV-FLEX; Arcuate nucleus; Lateral hypothalamic area

1. INTRODUCTION

The cocaine- and amphetamine-regulated transcript (CART) is a major neuropeptide involved in the regulation of diverse biological processes, including appetite control, maintenance of body weight, reward and addiction, psychostimulant effects, and neuroendocrine functions [1]. Among the wide and abundant central distribution of the peptide associated with multiple neurocircuits in mammals [2–4], CART expression shows predominance in neuroendocrine neurons [5], particularly in the hypothalamus at feeding-related regions [6,7]. Extensive research has focused on the role of CART in modulating feeding behavior and energy homeostasis [1], thereby exploring therapeutic potentials in the treatment of obesity and other metabolic disorders. However, identification of the hypothalamic sites of action and the mechanisms underlying CART function in homeostatic regulation remains challenging, mainly owing to the lack of information on the elusive CART receptor(s) that remain(s) unidentified [1]. In addition to the ubiquitous expression and broad connection of the peptide with various neuronal networks, the biosynthesis of CART involves complex post-translational processing, including the formation of various disulfide bonds [5,8], further rendering pharmacological intervention using any synthetic ex-vivo produced CART analogs difficult.

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Received October 10, 2017 • Revision received October 27, 2017 • Accepted October 30, 2017 • Available online 8 November 2017

https://doi.org/10.1016/j.molmet.2017.10.015
The functional importance of CART is highlighted by the strong evolutionary conservation across species [1,9]. Moreover, CART expression patterns are very similar between rodents and human, with high density of expression in hypothalamic areas important for energy homeostasis regulation, including the arcuate nucleus (Arc), lateral hypothalamic area (LHA), paraventricular nucleus (PVN), and dorso-medial hypothalamic nucleus [10,11]. However, studies of CART primarily conducted in rodents have so far generated inconclusive or discrepant results [1], highlighting the complexity in the physiology of endogenous CART biosynthesis and function. In rats and mice, in addition to alternative splicing of CART propeptides, which exhibit tissue- and brain region-specific expressions and functions [8,11,12]. Hence, the in vivo production of several CART isoforms also involves processing and expression in a tissue- and site-specific manner [8,12], generating at least two predicted active CART peptides, CART I (42-89/55-102) and CART II (49-89/62-102) [5,13]. Accordingly, studies in rodents have suggested the differential physiological functions of CART variants across the periphery as well as the central nervous system [8].

In light of the multiple factors involving complex region-specific post-translational processing, along with undetermined conformational profile and binding sites for the peptide, studies are challenged with limited targeting contacts to mimic or modulate endogenous CART expression and activity. Although most traditional studies have classified CART as anorexigenic [14–16], orexigenic evidences also exist [17–19]. In mouse models, global CART knockout led to an overall increase in body weight gain and adiposity, while lacking any overt impact on food consumption [1,20]. Supporting the catabolic property of CART, mutations and polymorphisms in the human CART gene are linked to obesity and associated metabolic disorders [21,22]. On the other hand, while intracerebroventricular (i.c.v.) administration of various CART isoforms in rodents consistently and dose-dependently suppressed food intake and weight gain [1,23], overexpression of CART I (55-102) in specific hypothalamic nuclei was shown to induce either stimulatory or inhibitory response on feeding and body weight [17,18,24]. This suggests that CART originating from different nuclei is engaged in differential functions involving discrete neurocircuits. Furthermore, central administrations of different CART variants, which demonstrate varying endogenous abundance in distinct brain regions, have exhibited site-specific activities and differential potencies in affecting food intake and body weight gain [25,26].

In an effort to identify the most critical hypothalamic sites of CART action in the control of energy homeostasis, the Arc and LHA have received considerable interest. The Arc represents a pivotal structure for adiposity signal integration and metabolic modulation [27], where CART expression is enriched and colocalized with the anorexigenic pro-opiomelanocortin (POMC) [28,29]. The first-order CART-containing neurons at the Arc launch axonal projections to neurons at the LHA and PVN [30–32]. Despite implications for the Arc-LHA pathway in mediating the anorexigen effects of CART [30,31], direct injection of CART into the Arc or LHA in rodents triggered overt orexigenic responses [17,18,24]. Differential neurochemical responses were also shown at the two regions. For instance, while CART mRNA levels at the Arc are markedly reduced by fasting and restored following refeeding [33,34], such correlation was not demonstrated at the LHA [35]. Moreover, CART in the LHA is found to be colocalized with the orexigenic-acting melanin-concentrating hormone (MCH) [36,37], further hinting that CART in different hypothalamic areas and neurons may exhibit disparate functions.

In this study, we aimed to more specifically investigate the effects of CART in the Arc and LHA on the regulation of feeding and energy homeostasis. For this, we developed models that allow the CART neuron-specific introduction and processing of CART by combining Cart-cre knock-in mice with a Cart adeno-associated viral vector designed using the flip-excision switch (AAV-FLEX) technology [38]. The AAV vector encodes the long form of the mouse Cart cDNA CART I (55-102), which is focused herein due to wider documentation and the reported higher potency in influencing appetite and body weight compared to the CART II (62-102) isoform [25,39]. This approach allows for either reintroducing CART specifically into CART neurons in otherwise CART-deficient mice, or to overproducing CART in Arc or LHA CART neurons, avoiding off-target side-effects associated with simple ectopic delivery. Furthermore, our viral approach also facilitates the in vivo processing and modifications of CART propeptides in a natural physiological environment, hence producing active peptides under endogenous conditions. Mouse models generated this way were then analyzed in a comprehensive phenotyping paradigm, determining all major aspects that are involved in energy homeostasis control.

2. MATERIALS AND METHODS

2.1. Animals

All research and animal care procedures were approved by the Garvan Institute/St. Vincent’s Hospital Animal Ethics Committee and were conducted in agreement with the Australian Code of Practice for the Care and Use of Animals for Scientific Purposes. Male mice were used for all experiments and were housed under conditions of controlled temperature (22 °C) and illumination (12:12 h light—dark cycle, lights on at 07:00 am). Mice were provided with ad libitum access to water and standard chow diet (8% calories from fat, 21% calories from protein, 71% calories from carbohydrate, 2.6 kcal/g; Gordon’s Specialty Stock Feeds, Yanderra, NSW, Australia).

2.2. Generation of inducible Cart-cre knock-in mouse model

To investigate the effects of CART action at particular hypothalamic regions, CART neuron-specific introduction of genetic elements was performed in an adult-onset inducible manner. This is enabled through the use of a conditional Cart-cre knock-in mouse line, for which the detailed generating procedures were described previously [20]. As illustrated [20], the inducible Cart-cre knock-in mice (Cartcre/cre) were generated by crossing the CartloxPlox line with mice that expressed Cre recombinase specifically in the oxytocytes to homozygosity, hence combining the tamoxifen-inducible Cre gene with the endogenous Cart promoter. In this study, both the homozygous Cartcre/cre and the heteryygous Cartcre/cre mouse lines were included. Comparison between the two groups serves to highlight the contribution of CART function in specific hypothalamic nuclei to feeding and metabolic regulation in the presence or absence of endogenous CART signaling.

2.3. Generation and expression of cre-inducible AAV-FLEX-Cart vector

To selectively express the bioactive CART in CART neurons, the flip-excision switch technique was employed to create a unique Cre-recombinase-dependent AAV vector for delivery, which carries a double floxed inverted open reading frame (Figure 1). The FLEX strategy utilizes the recombination mechanism whereby DNA elements flanked by loxP recombination sites in the opposite orientation are inverted by Cre-recombinase [38]. The AAV-FLEX-Cart vector consists of a codon-optimized mouse Cart cDNA for the expression of CART I (55-102) active peptide flanked by loxP-sites as displayed in Figure 1A.
Gene expression is driven by the chicken β-actin (CBA) promoter on a self-complementary AAV vector backbone packaged in AAV capsids of serotype 1. AAV vectors are produced by plasmid cotransfection of HEK293 cells, benzoxane-treatment of cell extracts and purification by high-performance liquid chromatography using AVB-Sepharose columns, as described [40]. Within this AAV-FLEX-Cart vector, the Cart coding sequence is flanked by a set of two distinct loxP sites positioned in the opposite orientation to each other (Figure 1Ai), where recombination only occurs between the same type of loxP sequences. The Cart coding unit was placed in a reverse thus inactive orientation downstream of the CBA promoter, preventing expression prior to Cre-driven recombination. Upon exposure to tamoxifen, Cre-recombinase is activated through translation from the cytoplasm to the nucleus, thereby catalyzing the inversion of the loxP-flanked cassette in two possible ways (Figure 1Aii). Subsequent recombination between the same type of loxP sites placed in the same orientation would eliminate the alternative loxP site in-between, thereby generating the stable transgene with the Cart sequence in the correct orientation for further transcription and translation (Figure 1Aiii). In parallel, AAV-FLEX-GFP encoding the green fluorescence protein (GFP) was used as the control. As Cre expression is driven by the endogenous Cart promoter in the Cart-cre knock-in mice, the AAV-FLEX-Cart vector and the AAV-FLEX-GFP control are targeted only to CART neurons. Hence, production of the active form of CART or GFP is exclusive to CART neurons.

2.4. Stereotaxic delivery of AAV-FLEX-Cart to CART neurons in Cart-cre knock-in mice

Adult-onset central CART neuron-specific expression of CART1 (55-102) in the Arc or LHA was achieved by intra-hypothalamic injection of the AAV-FLEX-Cart vector in parallel with i.c.v. tamoxifen administration in Cart-cre knock-in mice. At 11 weeks of age, Cartcre/cre and Cartcre+/ mice were anesthetized with intraperitoneal (i.p.) injection of 100/20 mg/kg ketamine/xylazine (Parke Davis-Pfizer, Sydney, Australia & Bayer AG, Leverkusen, Germany) and immobilized in a stereotaxic frame (David Kopf Instruments, Tuluenga, CA, USA) in the flat-skull position. A small incision was made to expose the skull, where a small hole was opened for injection. One microliter of the AAV-FLEX-Cart vector (titer 1.0 × 10^{12} viral particles/ml) or the AAV-FLEX-GFP control vector was injected unilaterally into the Arc or LHA at a speed of 0.1 μl/min using a 2-μl Hamilton syringe connected to a syringe infusion pump (World Precision Instruments Inc., Waltham, MA, USA). The AAV-FLEX-Cart or AAV-FLEX-GFP was stereotaxically delivered into both Cartcre/cre and Cartcre+/ mice, such that mice from each genotype were separated into two groups, one injected at the right ARC (coordinates relative to Bregma:

-2.18 mm anteroposterior, −0.25 mm mediolateral, and −5.5 mm dorsoventrally), and the other injected at the right LHA (coordinates relative to Bregma:

-1.94 mm anteroposterior, −1.0 mm mediolateral, and −5.25 mm dorsoventrally). Mice from each of the four groups were further divided evenly into two groups based on body weight and fat mass, one receiving the AAV-FLEX-Cart injection and the other littermate group injected with the AAV-FLEX-GFP control. The injection needle was gently withdrawn 15 min after injection. To activate the Cre-recombinase for mediating the transgene inversion and expression, 1 μl of the active metabolite of tamoxifen 4-OH [41] (20 mg/ml) (Sigma—Aldrich Pty Ltd, Sydney, NSW, Australia) dissolved in absolute ethanol was then injected unilaterally into the left lateral cerebral ventricle (LV) at a rate of 0.1 μl/min using a 2-μl Hamilton syringe (World Precision Instruments Inc.). The injection coordinates relative to Bregma were −0.34 mm anteroposterior, +1.0 mm mediolateral, and −2.5 mm dorsoventral corresponding to the left LV [42]. Body temperature of the animals was maintained by placement on a heating pad throughout the surgical process and the overnight recovery period. Mice were allowed two weeks of recovery with regular body weight assessment prior to any phenotypic characterization procedures. According to previous experience from extensive central viral delivery experiments [43], maximal expression of the transgene is estimated to be reached within two weeks and remain stable for at least 6 months. The conditional tamoxifen-induced Cre activation and concomitant intranuclear delivery of AAV-FLEX-Cart or AAV-FLEX-GFP in the hypothalamus resulted in Cart^{Cre+} and Cart^{Cre−} mice with CART or GFP selectively expressed in CART neurons at the Arc or LHA. For simplicity, the post-surgery subjects are referred to as Arc^{Cre+} or Arc^{Cre−}, and LHA^{Cre+} or LHA^{Cre−}. At least 10 mice were included for each of the resultant 8 groups, with n = 10−12 for each site of injection with either CART or GFP for each genotype. For each targeted area and each genotype, at least two cohorts of animals were used, each cohort consisting of AAV-FLEX-Cart-injected mice run in parallel with AAV-FLEX-GFP-injected mice from the same litter.

2.5. Verification of the expression site of targeted AAV-FLEX-Cart by in situ hybridization and immunohistochemistry

To validate the injection sites at the hypothalamic Arc or LHA regions where CART was introduced to CART neurons, and to verify functionality of the AAV-FLEX-Cart construct, in situ hybridization and immunostaining for CART mRNA and protein expression were examined in brains collected from Cart-cre knock-in mice. At the transcriptional level, efficiency of Cart reintroduction or overexpression was validated by in situ hybridization for Cart mRNA on brain cryosections from both the Arc^{Cre+} and LHA^{Cre+} Cart^{Cre+} and Cart^{Cre−} mice (n = 5 each) following procedures as described above. For the detection of CART-containing cells receiving AAV-FLEX-Cart delivery at the translational level, CART peptide expression was assessed by immunohistochemistry as previously described [20,43]. At the completion of study, a separate group of mice was anesthetized with i.p. injection of 100/20 mg/kg ketamine/xylazine (Parke Davis-Pfizer) and sacrificed via transcardial perfusion. The brains were subsequently fixed by perfusion and subjected to post-fixing and cryoprotection. Coronal brain slices (30 μm) from both the Arc^{Cre+} and LHA^{Cre+} Cart^{Cre+} and Cart^{Cre−} mice (n ≥ 5 each) were serially cryosectioned and incubated overnight with rabbit anti-mouse CART polyclonal primary antibody Cat-DVA [44] (gift from Dr. Anna Secher, Nordoisk, Copenhagen, Denmark) diluted 1:1500. Sections were then incubated with biotinylated goat anti-rabbit IgG secondary antibody (Vector Laboratories, Burlingame, CA, USA) diluted 1:250 for 2 h, followed by incubation with ExtrAvidin-peroxidase (Sigma—Aldrich, St. Louis, MO, USA). Immunostaining for CART expression was performed using diaminobenzidine (Dako, Carpinteria, CA, USA) as the chromogen. Free-floating brain sections were mounted on Superfrost® glass slides (Menzel-Glaser, Braunschweig, Germany) and covered-slipped. For visualization of CART immunoactivity (IR), brain sections containing the targeted Arc or LHA regions from respective mice were investigated using a Zeiss Axioshot microscope equipped with a ProgRes 3008 digital camera (Carl Zeiss, Jena, Germany). Bright field photomicrographs were acquired for the targeted hypothalamic regions on both the injection and contralateral sides of the brain for comparison. An increase in CART-IR on the injection side relative to the contralateral side therefore signifies the presence of AAV-delivered...
Figure 1: (A) Schematic representation of the AAV-FLEX-Cart vector targeted selectively to CART neurons in the Arc or LHA in Cart-cre knock-in mice. The Cre-inducible Cart AAV vector was designed using the FLEX switch strategy. (i) The targeting construct comprises the CBA promoter, followed by two pairs of heterotypic recombinase recognition loxP sites (here referred to as loxP-a and loxP-b) in antiparallel configuration, flanking the codon-optimized Cart coding region in the anti-sense inactive orientation relative to the promoter. (ii) Administration of tamoxifen activates Cre expression driven by the endogenous Cart promoter to mediate inversion of the floxed element in two possible manners, whereby subsequent recombination occurs only among the same type of loxP sites. Consequently, parallel loxP sites of the same type recombine, namely loxP-a with loxP-a, loxP-b with loxP-b, deleting the alternative loxP type in-between. (iii) Stable transgene consisting of the Cart coding unit in an orientation parallel to the promoter is transcribed and translated into bioactive and functional CART peptide, which is expressed specifically in the targeted CART neurons in the hypothalamus. Figure 1B–C: Representative fluorescence micrographs of coronal brain sections from Cartcre/cre and Cartcre/+ mice unilaterally injected with the AAV-FLEX-GFP control, displaying the Arc (B) and LHA (C) regions where GFP expression within the targeted neurons indicates accurate stereotaxic delivery. Figure 1D–G: Confirmation of Cart reintroduction and overexpression in the Arc and LHA by in situ hybridization in Cart-cre knock-in mice. Representative bright-field photomicrographs of coronal brain sections showing Cart mRNA at the Arc (D) and LHA (F) of Cartcre/cre and Cartcre/+ mice receiving AAV-FLEX-Cart versus AAV-FLEX-GFP injection. Scale bar = 200 μm. Hybridization signals at the Arc (E) and LHA (G) of respective brain sections are quantified as mean labeling intensity of neurons expressed as percentage coverage of neuronal surface by silver grains (RODs) within the defined areas of interest. 3V, third ventricle; fx, fornix; Arc, arcuate nucleus; LHA, lateral hypothalamic area. Data are means ± SEM and averaged for all mice from each group examined. ****p < 0.0001 for Cartcre/cre or Cartcre/+ mice targeted with AAV-Cart versus AAV-GFP in the Arc or LHA, or for comparisons between different genotypes receiving the same treatments.
CART-expressing CART neurons that are accurately targeted and functional. In parallel, brain sections from a portion of the Cartcrecre and Cartcre/+ mice with AAV-FLEX-GFP targeted into the Arc or LHA (n = 3 each) were prepared as described above and mounted on Super Frost slides for the detection of GFP expression by direct fluorescence microscopy. To detect GFP expression produced by the control targeting construct, brain sections containing the targeted Arc or LHA regions from respective mice were examined using a Leica DMS500 B fluorescent microscope (Leica, Chicago, IL, USA). Fluorescence micrographs were acquired for the visualization of GFP, indicative of successful delivery and expression of the empty control vector in CART neurons at the respective areas.

2.6. Determination of food intake and body weight
All Cartcrecre and Cartcre/+ mice were fed a standard laboratory chow and subjected to the same experimental procedures as follows. Body weight was measured weekly throughout the duration of studies. Following a two-week recovery period after the brain surgery, all mice were at 13–14 weeks of age. Mice were assessed for spontaneous food intake in the fed state as well as for fasting-induced food intake in response to 24-hr fasting. Basal daily food intake was determined as the average of duplicate readings obtained over two consecutive 24-hr periods. Twenty four-hr fasting-induced food intake was subsequently measured at 1, 3, 7, and 24 h after refeeding, while the corresponding body weight was recorded in parallel.

2.7. Indirect calorimetry of energy expenditure and assessment of physical activity
All Cartcrecre and Cartcre/+ mice with ArcGFPLHA in collaboration with the controls ArcGFP or LHA GFPl (n = 10 - 12 per group) were preferred from group housing to individual cages for acclimatization 2 days prior to imaging procedures. For area-specific exposure of skin to facilitate temperature reading, all mice were shaved at the IBAT and lumbar back regions under light isoflurane anesthesia in parallel. A high-sensitivity infrared camera (ThermoCAM T640, FLIR, Danderyd, Sweden, sensitivity = 0.04 °C) fixed on a tripod was placed 100 cm above the freely moving singly housed mouse, and thermographic images were taken for 1 min per mouse. After each 1-min measurement, the camera was moved to record surface temperatures of the next mouse. Temperature measurement was performed daily for three consecutive days.

2.9. Measurement of whole body and brown adipose tissue temperatures by infrared imaging
Whole body temperature and temperature of the interscapular brown adipose tissue (iBAT) were measured for both the Cartcrecre and Cartcre/+ cohorts by non-invasive high-sensitivity infrared imaging as described previously [20]. Briefly, all 17-week-old mice were transferred from group housing to individual cages for acclimatization 2 days prior to imaging procedures. For area-specific exposure of skin to facilitate temperature reading, all mice were shaved at the IBAT and lumbar back regions under light isoflurane anesthesia in parallel. A high-sensitivity infrared camera (ThermoCAM T640, FLIR, Danderyd, Sweden) was configured for the glucose levels obtained during intraperitoneal glucose tolerance test (IPGTT) and insulin tolerance test (IPITT). IPGTT and IPITT were conducted at 15 and 18 weeks of age, respectively. For IPGTT, mice were subjected to 6-hr fasting; food was removed from cage hoppers at 09:00 h and injection of a dose of a 10% d-glucose solution (1.0 g/kg body weight; Astra Zeneca, North Ryde, NSW, Australia) was performed 6 h later into the peritoneal cavity. For IPITT, mice were subjected to 5-hr fasting where food was removed from cage hoppers at 10:00 h and injection of a dose of insulin (0.5 IU/kg body weight) (Novo Nordisk Pharmaceuticals, Baulkham Hills, NSW, Australia) was performed 5 h later into the peritoneal cavity. For both tests, tail vein blood was collected at 0, 15, 30, 60, and 90 min following glucose or insulin injection, and glucose concentrations were measured using a glucometer (Accu-Check II; Roche, Castle Hill, NSW, Australia). For IPITT, insulin levels were subsequently quantified using a sensitive rat insulin radioimmunoassay (RIA) kit (Millipore, Billerica, MA, USA). Glucose and insulin tolerance curves constructed for the glucose levels obtained during IPGTT and IPITT respectively are presented in terms of absolute values. Area under the glucose (AUCglucose) or insulin (AUCinsulin) concentration curves between 0 and 90 min after glucose injection during IPGTT were plotted after subtraction of the initial concentrations prior to injection.

2.11. Tissue collection
Following completion of studies, all mice from both Cartcrecre and Cartcre/+ groups were sacrificed at 19 weeks of age. Animals were culled between 13:00 to 17:00 h through cervical dislocation followed by decapitation. Brains were promptly collected and frozen on dry ice, then stored at −80 °C until subsequent analysis as described below. The brown adipose tissue (BAT) at the interscapular area and the white adipose tissue (WAT) depots from the inguinal, epidydimal,
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2.12. In situ hybridization analysis of key hypothalamic neuropeptides
In situ hybridization was performed on brain sections of Cartcre/cre and CartCre/Cre mice from both the ArcCart-cre and LHACart-cre groups (n = 5 each) to determine mRNA expression of Cart, Pomc, neuropeptide Y (Npy), corticotropin-releasing hormone (Ctth), Mch, orexin (Oxr), tyrosine hydroxylase (Th), and thyrotropin-releasing hormone (Tth) at the hypothalamus following procedures as previously described [20]. Briefly, matching coronal brain sections (25 µm) collected from the Cart-cre knock-in mice were prepared from specific brain levels with respect to the Bregma to represent the hypothalamic areas involved in the regulation of energy homeostasis. Detection of the mRNA expression was performed for Cart in the Arc and LHA, for Npy and Pomc in the Arc, for Mch and Oxr in the LHA and, for Ctth, Tth, and Tth at the PVN. The cryosections were hybridized with [35S]-thio-dATP (PerkinElmer, Boston, MA, USA) radiolabeled DNA oligonucleotides complementary to mouse Cart: (5'-TCTTCTTCTTGGGACCGATATC- 
CACGGCAGAGTAGATGTCC AGG-3'), Pomc: (5'-TGGCTGCTTCCCGAG-CACACGCACACACACTATG GAGG-3'), Npy: (5'-GAGGGTCACTCCACA CACGGGATCCCTTG TTACCTA GCAT-3'), Ctth: (5'-CGGATAATCTC 
CATAGGTCTCTGTGTGAGTC TGGCTGACTGCT-3'), Mch: (5'-TTCTCTGC 
TTGAGACTGACATCTGAACTCCAT TCTCAGCTGG-3'), Orx: (5'- 
CTTTCCCAGAGTCAGGATACCCGCAGCGTGGT TGCCAGCTCCGTGC-3'), Th: (5'-CTCTAAGAGGGCGGATGGTGAGGA CGTCCAGTACATACTGAC-3') and Tth: (5'-AACCTACTTCTCAGAGGCTCCT GTAACGCGCTTCT-CAGTGTG-3'). Hybridization signals on sections were visualized by exposure to BioMax MR film (Kodak, Rochester, NY, USA) for 7—10 days and digitalized images from the scanned autoradiograms were acquired. For structural details, the brain sections were photoemulsion-dipped and superficially counterstained with hematoxylin, with which regions of interest were visualized and captured into digital images acquired by bright-field microscopy. Quantification of the mRNA expression levels of respective genes was performed by measuring the relative optical densities (RODs) within the brain areas of interest outlined with consistent definitions across corresponding sections on the photomicrographs using the National Institutes of Health ImageJ 1.61 software (written by Wayne Rasband; available at https://imagej.nih.gov/ij/). Background labeling was consistently uniform with signal levels below 5% of specific signal levels and was subtracted from the resultant signal density. Data are evaluated and presented as percentage of ROD averaged from at least three sections per mRNA assessed per animal.
2.13. Statistical analysis
All data are presented as means ± SEM. Differences among mouse groups of various genotypes and viral treatments were assessed by ANOVA or repeated-measures ANOVA combined with Bonferroni post-hoc analysis where appropriate. Energy expenditure, RER, and physical activity over the continuous 24-hr period were averaged for the whole 24-hr period, as well as the 12-hr light and dark phases individually. Comparison of energy expenditure (kcal/hr) between groups was performed by the analysis of covariance (ANCOVA) with lean mass as the covariate. The adjusted means of energy expenditure at a common lean mass for the comparison were generated by ANCOVA as presented. Statistical analyses were performed with GraphPad Prism 7 for Mac OS X (GraphPad Software, Inc. CA, USA). Statistical significance was defined as p-value ≤ 0.05.
3. RESULTS
3.1. Generation of hypothalamic CART neuron-specific re-introducing or overexpressing CART mouse models
To investigate the role of CART in the Arc and LHA, we employed our inducible Cart-cre knock-in lines selectively injected with AAV1-FLEX-CART vectors into these two brain areas. Specifically, we used heterozygous Cartcre/+ mice, which allow for the CART neuron-specific overexpression of CART, as well as homozygous Cartcre/cre mice with global CART ablation, thus allowing for the CART neuron-specific reintroduction of CART on an otherwise CART-deficient background [20]. To demonstrate the precise delivery of the construct to the targeted brain regions, we first employed the GFP reporter for visual guidance. Two weeks after AAV-FLEX-GFP injection, GFP expression was detected exclusively in either the Arc (Figs. 1B and S1A) or LHA (Figs. 1C and S1B), as shown in the representative photomicrographs, validating accurate targeting of the viral vectors. Next, to verify targeted expression and functionality of the unilaterally injected AAV-FLEX-Cart construct, which was activated through i.c.v. tamoxifen treatment, CART neuron-specific introduction of CART expression was determined on both the transcriptional and translational levels. Successful CART reintroduction was confirmed by in situ hybridization in both the Arc (Figure 1D—E) and LHA (Figure 1F—G) of Cartcre/cre mice, where CART mRNA expression was only observed in mice receiving AAV-FLEX-Cart while absent in control groups treated with AAV-FLEX-GFP. In parallel, Cartcre/cre mice subjected to Arc (Figure 1D—E) or LHA (Figure 1F—G) AAV overexpression displayed significantly upregulated Cart mRNA expression in the corresponding areas compared to the respective AAV-FLEX-GFP controls of the same genotype.
Immunostaining for CART protein expression further confirmed the exclusive CART-IR shown on the injection side, but not on the contralateral control side, for both the Arc (Figure 2A) and LHA (Figure 2B) regions in Cartcre/cre mice receiving AAV-Cart unilaterally. CART immunostaining at the respective local sites of delivery (Figure 2A—B) displayed a pattern that is consistent with endogenous CART-containing neurons [2,3], signifying selective CART reintroduction in CART neurons only. In contrast, lack of CART-IR on the contralateral sides (Figure 2A—B) denotes the regional and CART neuron-specific targeting, with minimal diffusion to other areas in the brain. No CART-IR was detected on matching sections from ArcGFP or LHA-GFP, Cartcre/cre controls (Figure 2A—B). Similarly, notably higher CART-IR was shown on the injection compared to the contralateral side for both the Arc (Figure 2A) and LHA (Figure 2B) in Cartcre/cre mice treated with AAV-CART. Such difference was not observed in Cartcre/cre counterparts receiving ArcGFP (Figure 2A) or LHA-GFP control constructs (Figure 2B). The visual guidance from CART expression further verified the correct stereotaxic injection for both Arc and LHA groups, while only data from mice identified for positive hits were used for down-stream analysis.
3.2. Selective reintroduction of CART in CART neurons in the arc but not the LHA reduces body weight
To investigate the basic metabolic characteristics of Cartcre/cre and CartCre/Cre mice re-introduced with or overexpressing CART in the (ArcCart) or (LHACart), male mice at the age of 11 weeks were unilaterally injected with AAV-FLEX-Cart or AAV-FLEX-GFP into the Arc or LHA, respectively, and body weight and other metabolic parameters were measured following the targeted brain surgery. In Cartcre/cre mice with endogenous CART expression in place, CART overexpression targeted at either the Arc (Figure 3A) or LHA (Figure 3B) had no obvious effect on body weight compared with respective
controls injected with the AAV-GFP control construct. The lack of difference was seen for both absolute body weight as well as weight gain expressed as a percentage of initial body weight (Figure 3A-B). In contrast, in Cartcre/cre mice without endogenous CART expression, CART reintroduction into the Arc CART neurons led to a significant decrease in both absolute body weight and percentage weight gain compared to AAV-GFP controls over the entire study period (Figure 3C). On the contrary, no significant effect on body weight was seen after CART was reintroduced into the LHA of Cartcre/cre mice (Figure 3D). These results suggest that CART action originating from the Arc but not...
Figure 3: Specific introduction of CART expression into CART neurons in the Arc led to significant decline in body weight in homozygous Cartkre/cre but not heterozygous Cartkre/+ mice. Absolute body weight and corresponding body weight change as a percentage of initial body weight measured weekly in Cartkre/+ (A - B) and Cartkre/cre (C - D) mice targeted with AAV-Cart or AAV-GFP in the Arc (A, C) or the LHA (B, D) (n = 10 - 12 per group). Data are means ± SEM and averaged for all mice from each group examined. *p ≤ 0.05, **p ≤ 0.01 for AAV-Cart versus AAV-GFP treatments, or for comparisons indicated by horizontal bar.
the LHA is responsible for the establishment of a negative energy balance that is leading to a body weight loss.

3.3. CART neuron-specific rescue of CART expression in the arc and LHA leads to opposite effects on adiposity

To elucidate the factors contributing to the differential responses in body weight to CART delivery, body composition analysis was performed. In line with the markedly lower body weight (Figure 3C), Cartcre/cre mice expressing Arc CART demonstrated a significant decrease in whole body fat mass compared with ArcGFP controls (Figure 4A), as confirmed by the significantly reduced mass of dissected white adipose tissue depots (Figure 4B). Interestingly, whole body lean mass was unaltered following Arc CART reintroduction (Figure 4C). In contrast, despite similar body weight, Cartcre/cre mice with CART rescue in the LHA exhibited a significant increase in whole body fat mass (Figure 4D) compared to the AAV-GFP controls, consistent with the markedly higher dissected fat weights (Figure 4E). Unlike the Arc CART reintroduction, rescue of CART in the LHA was accompanied by a significant reduction in lean mass relative to LHAGFP counterparts (Figure 4F). The lower lean mass may have offset the greater fat mass (Figure 4D-E), leading to the unvaried absolute body weight observed (Figure 3D). It is therefore possible, while not directly

Figure 4: Reintroduced CART expression in CART neurons in the Arc and LHA reduces and increases fat mass respectively in Cartcre/cre mice. Body composition analysis using dual-energy X-ray absorptiometry. Whole body fat mass in Cartcre/cre mice injected with ArcCART versus ArcGFP (A) or LHACART versus LHAGFP (D) (n = 10 - 12 per group). Tissue mass of dissected white adipose tissue (WAT) depots after sacrifice in the corresponding Arc (B) and LHA (E) groups. i, inguinal; e, epididymal; m, mesenteric; r, retroperitoneal; total, summed weight of i, e, m, and r WAT depots. Whole body lean mass in the corresponding Arc (C) and LHA (F) groups. Data are means ± SEM and averaged for all mice from each group examined. *p ≤ 0.05, **p ≤ 0.01, ***p ≤ 0.001 for AAV-Cart versus AAV-GFP treatments.
involved in weight regulation, LHA CART reintroduced on an otherwise CART-negative genetic background may modulate body composition by increasing adiposity and decreasing lean body mass. No apparent difference was seen in the weights of major organs between genotypes in either brain area (Fig. S2 A-B). Interestingly, investigation of Cartcre/cre mice subjected to CART overexpression in either the Arc or the LHA, which showed unaltered body weight (Figure 3A–B), also revealed unchanged adiposity (Fig. S2 C-F) and lean mass (Fig. S2 G-H) compared to AAV-GFP controls. As part of the body composition analysis, and based on the known link between CART and bone remodeling [16,46], effects of chronic CART introduction specifically into CART neurons in the Arc or LHA on bone parameters were also examined. Interestingly, neither introduction of CART into the Arc or LHA had any significant effect on both whole body BMC and BMD for either Cartcre/cre (Fig. S3 A-D) or Cartcre/cre (Fig. S3 E-H) mice. The unchanged bone mass phenotype suggests that CART neurons in these two hypothalamic regions are not directly involved in bone homeostatic control, while CART in other brain areas may be responsible for the regulation of bone metabolism.

3.4. CART reintroduction in the arc and LHA led to differential effects on food intake and energy expenditure

In order to determine which side of the energy balance equation is responsible for the altered body weight and body composition caused by the selective CART reintroduction into Arc or LHA CART neurons, we next measured spontaneous and fasting-induced food intake in Cartcre/cre mice. Spontaneous daily food intake under fed conditions was unchanged between ArcCART and ArcGFP groups (Figure 5A), excluding food intake alteration as a significant contributing factor to the marked decrease in body weight gain seen in ArcCART;Cartcre/cre mice (Figure 3C).

In contrast, despite unaltered body weight in Cartcre/cre mice expressing LHACART (Figure 5D), a trend towards higher spontaneous food intake was seen compared with LHAGFP controls (Figure 5B), which is consistent with the significantly elevated fat mass seen in these mice (Figure 4D–E). Food intake in response to 24-hr fasting was unaffected by replenishing CART expression in Cartcre/cre mice at either the Arc (Fig. S4 A) or LHA (Fig. S4 B). However, despite similar weight loss from the 24-hr fasting, ArcCART mice experienced an accelerated weight regain during refeeding compared to ArcGFP controls (Fig. S4 C). No significant difference in body weight loss and subsequent recovery was seen in the LHA-CART mice (Fig. S4 D).

To also investigate the other component of the energy balance equation - energy expenditure, open-circuit indirect calorimetry was employed. Interestingly, a marked increase in energy expenditure was demonstrated in ArcCART compared to ArcGFP groups (Figure 5A), indicating non-shivering thermogenesis [48] when compared to AAV-GFP controls. Interestingly, the pronounced reduction in energy expenditure in LHA-CART;Cartcre/cre mice was not matched with strong effects on whole body temperature which shows a trend to a reduction relative to controls (Figure 5I). Furthermore, significantly lower BAT temperature (Figure 5J) and BAT weight (Figure 5K) were seen in Cartcre/cre mice with LHA-CART but not ArcCART mice when compared to the respective GFP controls, suggesting an involvement of LHA CART in the regulation of BAT thermogenesis. In addition, as the tail is regarded as a major thermoregulatory organ for heat dissipation [49], tail temperature was also examined and was demonstrated to be significantly increased in LHA-CART but not ArcCART;Cartcre/cre mice compared with controls (Figure 5L). The elevated tail heat loss possibly contributed to the reduced temperature seen in the whole body (Figure 5I) and the IBAT region (Figure 5J), indicating a potential involvement of CART function in the LHA to the regulation of vasodilatation and vasoconstriction [49].

Determination of RER, a measure for fuel source preference, revealed no significant difference between treatments for both Arc (Figure 5M) and LHA (Figure 5N) groups throughout the entire photoperiod as well as when averaged for 24-hr (Figure 5O). However, a modest increase in RER was demonstrated for ArcCART compared to ArcGFP;Cartcre/cre mice (Figure 5M), possibly suggesting higher relative preference for carbohydrate over fat as the metabolized fuel, which may be associated with the diminished fat availability (Figure 4A–B).

3.5. CART reintroduction in the arc and LHA induced differential effects on the hypothalamic expression of key peptides in the control of energy homeostasis

To elucidate the neurochemical alterations and the central mechanisms underlying the phenotypes manifested following CART reintroduction into Arc or LHA CART neurons, hypothalamic expression of several neurotransmitters and hormones that are pivotal in the regulation of energy homeostasis were profiled. As CART neurons in the Arc co-express the anorexigenic POMC and are adjacent to the orexigenic NPY neurons [27,32], in situ hybridization was performed to measure the transcript expression of Pomc and Npy in the Arc of ArcCART mice. Cartcre/cre mice expressing ArcCART demonstrated a notable trend to a decrease in Pomc mRNA level (Figure 6A) and significantly reduced Npy expression (Figure 6B) compared to ArcGFP controls. In contrast, no difference was observed in the Arc mRNA level for either neuropeptide in ArcCART;Cartcre/cre mice (Figure 6A–B). Since CART has been implicated as a major regulator of CRH neurons of the hypothalamic-pituitary-adrenal (HPA) axis [50,51], and CART neurons in the Arc are known to innervate CRH neurons in the PVN [30,52], we also investigated the effects of CART targeted to the Arc on Crh mRNA expression in the PVN in either Cartcre/cre or Cartcre/cre mice. Interestingly, neither of the genotypes showed any significant impact on Crh mRNA expression (Figure 6C).

In the LHA of both humans and rodents, CART has been reported to have a high level of colocalization with the orexigenic neuropeptide melanin-concentrating hormone (MCH) [36,37]. Orexin, another...
Figure 5: CART reintroduction in Cart\textsuperscript{cre/cre} mice upregulates energy expenditure when targeted in the Arc while downregulating energy expenditure and BAT temperature when introduced into the LHA. Daily spontaneous/basal food intake during fed state in Cart\textsuperscript{cre/cre} mice expressing Arc\textsuperscript{CART} versus Arc\textsuperscript{GFP} (A) or LHA\textsuperscript{CART} versus LHA\textsuperscript{GFP} (B) (n = 10 - 12 per group). Indirect calorimetric assessments for the 24-hr time course of energy expenditure (C, D), physical activity (F, G) and respiratory exchange ratio (M, N) to compare Arc\textsuperscript{CART} against Arc\textsuperscript{GFP} and LHA\textsuperscript{CART} against LHA\textsuperscript{GFP} treatments in Cart\textsuperscript{cre/cre} mice respectively. Corresponding average values for energy expenditure (E), physical activity (H), and respiratory exchange ratio (O) over the total 24-hr period were shown in adjacent bar graphs. Energy expenditure was adjusted for lean mass and compared between groups by analysis of covariance. The adjusted means of energy expenditure were presented at the common lean mass of 22.515 g (Arc) and 21.921 g (LHA) respectively. Open and filled horizontal bars indicate the light and dark photoperiods, respectively. Temperatures of the lumbar back region (I), the interscapular brown adipose tissue (IBAT) (J), and the tail (L) measured by high-sensitivity infrared imaging. (K) Dissected BAT mass. Data are means ± SEM and averaged for all mice from each group examined. *p < 0.05, ****p < 0.0001 for AAV-Cart versus AAV-GFP treatments, for comparisons between mice receiving the same treatments at the Arc versus LHA, or for comparisons indicated by horizontal bar.
Figure 6: CART neuron-specific CART reintroduction in the Arc and LHA of Cart<sup>cre/cre</sup> mice differentially impacts mRNA expression of key hypothalamic peptides in the control of energy homeostasis. In situ hybridization for Pomc (A) and Npy (B) mRNA expression in the Arc and Crh (C) mRNA expression in the PVN in Cart<sup>cre/cre</sup> and Cart<sup>cre/</sup> mice expressing Arc<sup>CART</sup> versus Arc<sup>GFP</sup> (n = 5 per group). In situ hybridization for Mch (D) and Orx (E) mRNA expression in the LHA, for Crh (F), Th (G), and Trh (H) mRNA expression in the PVN, and for Pomc (I) and Npy (J) mRNA expression in the Arc in Cart<sup>cre/cre</sup> mice expressing LHA<sup>CART</sup> versus LHA<sup>GFP</sup> (n = 5 per group). Bright-field photomicrographs of coronal brain sections showing mRNA expression for Pomc (A) and Npy (B) in the Arc, and for Crh (F) and Th (G) in the PVN. Scale bar = 200 μm or 100 μm. Hybridization signals are quantified to obtain mean labeling intensity of neurons expressed as percentage coverage of neuronal surface by silver grains (RODs) within the defined areas of interest. PVN, paraventricular nucleus; Arc, arcuate nucleus; Pomc, proopiomelanocortin; Npy, neuropeptide Y; Crh, corticotropin-releasing hormone; Mch, melanin-concentrating hormone; Orx, orexin; Th, thyroxine hydroxylase; Trh, thyrotropin-releasing hormone. Data are means ± SEM and averaged for all mice from each group examined. **p ≤ 0.01 for AAV-Cart versus AAV-GFP treatments.
feeding-stimulating peptide, can also be found in the LHA [2,44]. To examine the potential of these two neuropeptides acting together with CART to control feeding and energy homeostasis, mRNA levels of Mch and Orx were evaluated. Interestingly, CART rescue in the LHA of Cartcre/cre mice did not affect the mRNA level of either Mch (Figure 6D) or Orx (Figure 6E), despite the significant trend towards an increased spontaneous food intake (Figure 5B) seen in these mice. However, LHA-CART;Cartcre/cre mice showed significantly higher Crh levels (Figure 6F) and a significant decrease in Th expression (Figure 6G) in the PVN compared with GFP controls, consistent with the role of these PVN peptides in the regulation of energy expenditure and BAT thermogenesis [43,53]. The reduced Th transcript expression was also in line with the potential decrease in vasoconstriction reflected by the greater tail temperature thus heat loss in these mice (Figure 5L). In contrast, the mRNA level of Trh in the PVN, another key regulator of the HPA axis with implications in the control of energy balance and thermoregulation [54] was unchanged in LHA-CART;Cartcre/cre mice (Figure 6H). Similarly, CART reintroduction into the LHA in Cartcre/cre mice had no impact on the expression levels of Pomc (Figure 6I) or Npy (Figure 6J) in the Arc.

3.6. CART neuron-specific reintroduction of CART in the LHA reduced glucose tolerance and insulin responsiveness

Based on the well-established link between fat metabolism and glucose homeostasis [55] and given the distinctly altered fat content in Cartcre/cre mice expressing Arc-CART and LHACART, various parameters of glucose metabolism were investigated. Following i.p. glucose challenge, blood glucose levels throughout the 90 min duration were indistinguishable between Arc-CART and ArcGFP Cartcre/cre mice (Figure 7A), as reflected in the unchanged AU Glucose. In contrast, LHA-CART;Cartcre/cre mice demonstrated significantly higher blood glucose levels compared with GFP controls during the course of IPGTT as shown by the increased AUC (Figure 7B), which suggests reduced glucose tolerance. Interestingly, despite different responses to glucose challenge between the Arc and LHA groups (Figure 7B), serum insulin levels during IPGTT were not different compared to corresponding controls when reintroduced into either the Arc (Figure 7C) or LHA (Figure 7D).

Similar to the GTT, insulin responsiveness using the IP-ITT test was not altered between the Arc-CART and ArcGFP mice displaying unvaried blood glucose levels in response to i.p. insulin (Figure 7E). However, LHA CART mice with similar initial fasted levels demonstrated significantly higher blood glucose excursion throughout the 90 min period compared to LHA-GFP counterparts (Figure 7F), as reflected by the corresponding, prominent increase in AU Glucose compared to controls (Figure 7F). The reduction in insulin responsiveness may be due to decreased lean mass as well as increased adiposity in these mice, which may thus contribute to the reduced glucose tolerance observed in LHA-CART mice (Figure 7B).

4. DISCUSSION

This study presents the first characterization of energy homeostatic functions of CART specifically expressed in the Arc or LHA in a CART neuron-specific manner, through targeted unilateral AAV-FLEX CART delivery in either the homozygous or heterozygous version of the Cartcre knock-in mice. Intriguingly, in Cartcre/cre mice with existing endogenous CART expression, CART overproduction in Arc or LHA CART neurons had little effect on any of the body composition and metabolic parameters measured. On the other hand, significant and distinctive changes were demonstrated in Cartcre/cre mice subjected to CART neuron-specific reintroduction of CART into the Arc and LHA on an otherwise CART-deficient background. In Cartcre/cre mice, CART selectively targeted to CART neurons in the Arc markedly reduced body weight and fat mass without altering lean mass, whereas CART reintroduction into the LHA led to significant fat mass gain and lean mass loss, resulting in unchanged body weight. Although food consumption was unaltered, Arc-CART;Cartcre/cre mice showed an increase in both energy expenditure and physical activity, accompanied by a significant decrease in Npy mRNA expression in the Arc, which may have contributed to the observed reduction in adiposity. In comparison, the elevated fat mass in LHA-CART;Cartcre/cre mice was associated with greater spontaneous food intake and lower energy expenditure, with significantly reduced Th and notably increased Crh mRNA levels in the PVN. In line with this, the LHA CART-mediated suppression in energy expenditure was paralleled with a significant drop in BAT temperature and BAT weight, while both parameters remained unchanged in Arc-CART;Cartcre/cre counterparts. Interestingly, glucose homeostasis was also differentially regulated in the two groups. While LHA-CART;Cartcre/cre mice displayed reduced insulin responsiveness and reduced glucose tolerance, attributed at least in part to the higher fat and lower lean body masses, Arc-CART;Cartcre/cre mice, on the other hand, were not influenced in this aspect. These findings demonstrate that CART neurons in the Arc and LHA play differential roles in the control of feeding behavior, body composition, and energy homeostasis.

From existing literature, CART has been classified primarily as anorectic, as summarized from i.e.v. CART studies, as well as global CART knockout rodents [1,20], which showed an increase in body weight gain and adipose mass with unchanged feeding behavior [15,20]. Consistent with this classic view, CART reintroduction into the Arc of otherwise CART-deficient mice leads to reduced body weight and lower fat mass, which is accompanied by increased energy expenditure and physical activity, with unaltered food intake. This confirms the Arc as a crucial hypothalamic site for the anorexigenic potency of CART, since the limited gene dosage from the unilateral CART reintroduction was sufficient to trigger profound catabolic effects. Mechanistically, a substantial decline in the Arc Npy level was observed in these mice, which may be responsible, at least in part, for the increased energy expenditure [27], indicating the critical involvement of Arc NPY in CART action in regulating energy homeostasis. As opposed to the catabolic effects displayed in Arc-CART;Cartcre/cre mice with unaffected feeding behavior, direct evidence for an orexigenic role of CART is also supported by the LHA-CART-directed elevation in food intake and adiposity under CART-deficient condition. This finding is in line with a prior study where heightened Cart expression in the LHA was associated with elevated feeding and body weight [56]. More importantly, the increased adiposity in these mice may be achieved by the alterations in Th and Crh expression in the PVN, indicating the possible control function of LHA CART neurons on TH and CRH neurons in the PVN. Additionally, the anabolic profile of LHA CART;Cartcre/cre mice comprises a reduction in lean mass, lower energy expenditure, and reduced BAT temperature, suggesting that CART in the LHA regulates energy homeostasis through multiple mechanisms. Furthermore, as the LHA is traditionally described as the hunger center [57], and lesions in the LHA were reported to cause hypophagia and weight loss [58], our results further confirm the significance of CART in the orexigenic activity of the LHA. Moreover, since CART knockout mice showed obesogenic traits [14,15,20], Arc CART activity may exhibit greater baseline potency than CART in the LHA, and is therefore regarded a major contributor to the overall CART system in energy homeostasis. Contrary to our results, previous studies in rodents subjected to intra-Arc or -LHA CART administration showed body weight gain and
increased feeding [17–19]. The different responses to CART action are most likely due to the different techniques employed. While the AAV-FLEX strategy employed in this work enables the endogenous processing and synthesis of endogenous CART selectively in CART neurons, previous intra-hypothalamic CART delivery by intranuclear injection or chronic infusion leads to widespread and uncontrolled action of CART peptide in these nuclei and areas. Furthermore, when chemically synthesized CART fragments are used, it is only presumed that these peptides would represent the active form of CART produced by CART neurons. The representativeness of such methods for replicating physiological CART function may be questionable, as evident in the discrepancy between pharmacological studies [1]. In addition, whilst few studies have identified any impact of CART injection on energy expenditure and physical activity, the two parameters were profoundly and differentially regulated in ArcCART and LHACART Cartcre/cre mice. Further verifying the efficacy of this viral approach for CART expression, this study also provided the first evidence for LHA CART in the regulation of glucose metabolism. However, the reduced

Figure 7: CART neuron-specific rescue of CART expression in the LHA of Cartcre/cre mice reduces glucose tolerance and insulin responsiveness. Blood glucose levels during a 90-min intraperitoneal glucose tolerance test (1.0 g/kg body weight) performed in 6-hr fasted Cartcre/cre mice expressing ArcCART versus ArcGFP (A) or LHACART versus LHAGFP (B) (n = 10 - 12 per group). Adjacent bar graphs showing the area under the glucose concentration curve (AUCglucose) between 0 and 90 min after glucose injection. Corresponding serum insulin levels during the 90-min IPGTT procedure for the Arc (C) and LHA (D) groups. Adjacent bar graphs showing the area under the insulin concentration curve (AUCinsulin) between 0 and 90 min after glucose injection. Absolute blood glucose levels during a 90-min intraperitoneal insulin tolerance test (0.5 IU/kg body weight) performed in 5-hr fasted Cartcre/cre mice expressing ArcCART versus ArcGFP (E) or LHACART versus LHAGFP (F). Adjacent bar graphs showing AUCglucose between 0 and 90 min after insulin injection. Data are means ± SEM and averaged for all mice from each group examined. *p ≤ 0.05, **p ≤ 0.01 for AAV-Cart versus AAV-GFP treatments, or for comparisons indicated by horizontal bar.
glucose tolerance and insulin responsiveness resulted from CART reintroduction into the LHA could also be secondary to the higher fat and lower lean masses [59] in these mice. Importantly, while hypothalamic CART has been reported to be involved in bone physiology [16,46], our data here show that CART reintroduction into CART neurons in either of the two nuclei had only a minor effect on bone mass, suggesting that other hypothalamic CART neuronal population are more critical for this CART-related control mechanism. In addition, our study also revealed that overproducing CART in specific CART neurons in the Arc and LHA of Cartc/cre mice does not confer any additive function compared to controls with existing endogenous CART, or enhance any effect seen in the respective Cartc/cre mice with reintroduced CART expression. The lack of dose response in Cartc/cre mice could either be due to a saturation effect above a certain expression threshold for CART at the particular region or the insufficiency of unilateral CART delivery to generate any detectable physiological shift. Future work involving bilateral targeting of CART may provide additional insights into the validity of this conjecture. Potentially, any effects from the introduced CART in the Cartc/cre mice may also be subject to counterbalancing actions from endogenous CART expressed locally or at other central areas. This further highlights the usefulness of our Cartc/cre mice as a valuable instrument for determining region-specific CART activity. In conclusion, via CART neuron-specific reintroduction of CART in CART-deficient mice, the catabolic and anabolic homeostatic influences of CART were demonstrated in the Arc and LHA, respectively. This work supports the notion of an emerging, multifaceted, and nuclei-specific function of CART in appetite control and energy homeostasis regulation. Collectively, our results consolidate the postulation that CART in discrete hypothalamic nuclei performs distinct functions [30–32]. While appetite-stimulating properties were indicated for LHA CART neurons in these two regions may serve as therapeutic entry points for appetite control and body weight maintenance in the treatment of obesity. Future work applying the same design with bilateral CART delivery may facilitate the fine-tuning of CART dosage effects in the presence and absence of endogenous CART. In parallel, comprehensive neurochemical profiling upon CART reintroduction or overexpression in other areas may provide further invaluable insights into the elements and neuronal signaling pathways that interact with the CART system, thereby unraveling the underlying mechanisms linked to CART action in the regulation of feeding and energy homeostasis.

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ACKNOWLEDGMENTS

The authors are grateful to Mr. Ronaldo Enriquez for providing technical assistance for the animal work. We thank the staff of the Garvan Institute Biological Testing Facility for facilitation of these experiments. This research was supported by the National Health and Medical Research Council of Australia (NHMRC) with grant number 1081290, a Research Fellowship to H.H., a Postgraduate Scholarship to J.L., and the Austrian Science Fund (FWF grant J-3814) to A.F.

CONFLICT OF INTEREST

No potential conflicts of interest associated with this article were reported and there has been no significant financial support for this work that could have influenced its outcome.

APPENDIX A. SUPPLEMENTARY DATA

Supplementary data related to this article can be found at https://doi.org/10.1016/j.molmet.2017.10.015.

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**MOLECULAR METABOLISM** 7 (2018) 102 – 118

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