TRAP-seq defines markers for novel populations of hypothalamic and brainstem LepRb neurons

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ABSTRACT

Objective: Leptin acts via its receptor (LepRb) on multiple subpopulations of LepRb neurons in the brain, each of which controls specific aspects of energy balance. Despite the importance of LepRb-containing neurons, the transcriptome and molecular identity of many LepRb subpopulations remain undefined due to the difficulty of studying the small fraction of total cells represented by LepRb neurons in heterogeneous brain regions. Here we sought to examine the transcriptome of LepRb neurons directly and identify markers for functionally relevant LepRb subsets.

Methods: We isolated mRNA from mouse hypothalamic and brainstem LepRb cells by Translating Ribosome Affinity Purification (TRAP) and analyzed it by RNA-seq (TRAP-seq).

Results: TRAP mRNA from LepRb cells was enriched for markers of peptidergic neurons, while TRAP-depleted mRNA from non-LepRb cells was enriched for markers of glial and immune cells. Genes encoding secreted proteins that were enriched in hypothalamic and brainstem TRAP mRNA revealed subpopulations of LepRb neurons that contained neuropeptide-encoding genes (including prodynorphin, Pdyn) not previously used as functional markers for LepRb neurons. Furthermore, Pdyn 3lox-mediated ablation of LepRb in Pdyn-expressing neurons (LepRb 3lox/3ox KO mice) blunted energy expenditure to promote obesity during high-fat feeding.

Conclusions: TRAP-seq of CNS LepRb neurons defines the LepRb neuron transcriptome and reveals novel markers for previously unrecognized subpopulations of LepRb neurons.

Keywords Leptin; Hypothalamus; Brainstem; Neuropeptides; Dynorphin

1. INTRODUCTION

1.1. Leptin

The hormone leptin, which is produced by adipocytes to signal the repletion of fat stores, acts via the leptin receptor (LepRb) to modulate food intake and energy expenditure [1–3]. Leptin action is also crucial for the control of glucose homeostasis and other metabolic parameters [4,5]. Moreover, leptin controls energy balance and metabolism by acting on LepRb expressing neurons in the central nervous system [6–9]. Given the centrality of leptin action to the control of body weight and metabolism, it is crucial to understand mechanisms of leptin action, including the roles for each set of LepRb neurons, since these represent points of potential therapeutic intervention.

1.2. LepRb neurons

Anatomically and molecularly distinct subpopulations of LepRb neurons each play specific roles in leptin action [10]. Although hypothalamic LepRb populations have historically been the best characterized, the brainstem also contains several substantial populations of LepRb neurons, including populations in the ventral tegmental area (VTA), dorsal and linear raphe nuclei (DR, LR), Edinger-Westphal nucleus (EW), periaqueductal gray matter (PAG), parabrachial nucleus (PBN), and nucleus of the solitary tract (NTS) [11–14]. Only a few neuropeptides expressed in brainstem LepRb neurons have been identified, however, and (with few exceptions) brainstem LepRb neurons remain largely uncharacterized [15–17].

1.3. Known subsets of hypothalamic LepRb neurons

Hypothalamic LepRb neurons, including those that contain Nos1 or Vgat, play major roles in the control of energy balance and metabolism, but represent large, heterogeneous and dispersed sets of cells that are not well-suited to circuit-level analysis [8,18,19]. Smaller, circumscribed sets of LepRb neurons that reside in the hypothalamic arcuate nucleus (ARC) and express either Pomc and Carpt or Agrp and Npy contribute to the control of energy balance and glucose homeostasis [20]. The modest effects observed upon manipulation of LepRb in...
these neurons suggest important roles for other hypothalamic LepRb neurons in leptin action, however [21–24]. Similarly, SF1/PACAP-containing ventromedial hypothalamic nucleus (VMH) neurons and the PrT/expressing subset of dorsomedial hypothalamic (DMH) LepRb neurons participate in the control of energy expenditure, but only modestly contribute to the overall regulation of body weight by leptin [25–27]. A subgroup of lateral hypothalamic (LHA) LepRb neurons that express neurotensin (Nts) modulates the mesolimbic dopamine system but also contributes only a small amount to the control of energy metabolism and behavior by leptin [28–30]. Even taken together, these known subpopulations of hypothalamic LepRb neurons constitute only a fraction of total hypothalamic LepRb neurons and fail to explain the totality (or even the majority) of leptin action on feeding, metabolic control, and body weight regulation. Thus, additional, uncharacterized, groups of hypothalamic and brainstem LepRb neurons contribute importantly to overall leptin action.

1.4. Challenges inherent to studying gene expression in LepRb neurons

To identify, manipulate, and understand the function of potentially important but currently unrecognized subpopulations of LepRb neurons, it is necessary to identify other genes, including neurotransmitters, expressed in these cells. Since LepRb-expressing neurons comprise only a fraction of the cells within the nuclei in which they reside, it has not been possible to disentangle the transcriptome of LepRb cells from that of other cells within these areas [11,12,31,32]. Fluorescent cell sorting can isolate labeled cells from complex populations, but this approach is suboptimal for LepRb neurons, since hypothalamic and brainstem neurons survive isolation procedures poorly at ages when fluorescent markers for LepRb neurons are robustly expressed. We thus set out to examine the transcriptome of LepRb neurons by expressing an enhanced green fluorescent protein-tagged ribosomal subunit (eGFP-L10a) selectively in LepRb neurons to enable immunopurification of ribosomes and their associated mRNA.

2. MATERIALS AND METHODS

2.1. Animals

Rosa26eGFP-/+ mice were generated as previously described in Ref. [33]. The generation of LepRb+/+ mice has also been previously described in Ref. [34]. LepRb-/- mice were crossed to Rosa26FF-/+ mice to generate LepRb+/+Rosa26FF-/+ mice, which were subsequently intercrossed to generate double homozygous LepRb+/+Rosa26FF-/+ study animals. Pdyncre mice [33] or Taccre/+ mice (Taccre/+; Jackson Laboratory Stock # 021877) were crossed to Rosa26FF-/+ mice (Gt(Rosa)26Sor<tm1(EFP)Cok/J; Jackson Laboratory, Stock # 006149) to generate PdyncreFP or TaccreFP mice for study. LepRb+/+ mice were as described previously in Ref. [35]. Pdyncre+/- mice were bred to LepRb+/+ animals, producing Pdyncre+/-; LepRb+/+ mice, which were bred to LepRb+/+ mice to generate Pdyncre+/-; LepRb+/+ mice and littermate control (Pdyncre+/-; LepRb-/-; and LepRb+/+) mice for study. All procedures were approved by the University of Michigan Institutional Animal Care and Use Committee (IACUC) in accordance with AAALAC and NIH guidelines. Animals were bred at the University of Michigan and maintained in a 12 h light/12 h dark cycle with ad libitum access to food and water.

2.2. Immunoprecipitation of ribosomes (TRAP)

Adult homozygous LepRb-/-GFP-/- mice were anesthetized and their brains removed to a mouse coronal brain matrix (1 mm sections) to isolate the hypothalamus or brainstem: material from multiple animals was pooled to produce each sample. For hypothalamic dissections, a 3 × 3 × 3 mm block was dissected from the ventral diencephalon immediately caudal to the optic chiasm. For brainstem dissections, serial 1 mm sections were removed and individual LepRb-containing nuclei (including the ED, DR, LA, PAG, PBN and NTS) were dissected by hand and pooled. Messenger RNA was isolated from eGFP-tagged ribosomes, as well as from the eGFP-depleted fraction, as previously described in Ref. [36]. RNA was assessed for quality using the TapeStation (Agilent, Santa Clara, CA). Samples with RINs (RNA Integrity Numbers) of 8 or greater were prepped using the Illumina TruSeq mRNA Sample Prep v2 kit (Catalog #s RS-122-2001, RS-122-2002) (Illumina, San Diego, CA), and 0.1–3 μg of total RNA was converted to mRNA using a polyA purification. The mRNA was fragmented via chemical fragmentation and copied into first strand cDNA using reverse transcriptase and random primers. The 3′ ends of the cDNA were denatured, and 6-nucleotide-barcoded adapters ligated. The products were purified and enriched by PCR to create the final cDNA library. Final libraries were checked for quality and quantity by TapeStation (Agilent) and qPCR using Kapa’s library quantification kit for Illumina Sequencing platforms (catalog # KK4835) (Kapa Biosystems, Wilmington, MA). They were clustered on the cBot (Illumina) and sequenced 4 samples per lane on a 50 cycle single end run on a HiSeq 2000 (Illumina) using version 2 reagents according to manufacturer’s protocols.

2.3. RNA-seq analysis

50 bp single-end reads underwent QC analysis prior to alignment to mouse genome build mm9 using TopHat and Bowtie alignment software [37]. Differential expression was determined using Cufflinks Cuffdiff analysis, with thresholds for differential expression set to fold change >1.5 or <0.66 and a false discovery rate of <0.05 [38]. Lists of differentially expressed genes were then queried against the Uniprot Database to identify secreted proteins [39].

2.4. Leptin treatment, colchicine treatment, and immunohistochemistry

LepRbPdynKO mice had food removed at the onset of the light cycle. Animals were treated 4 h later with meteletepin (5 mg/kg, i.p.) (a generous gift from AstraZenica, Inc.) or vehicle and were subjected to perfusion 1.5 h after leptin treatment. Treatment with ICV colchicine (10 μg) to concentrate neurones in the soma for some experiments was for 2 days prior to perfusion. For perfusion, mice were anesthetized with a lethal dose of intraperitoneal pentobarbital and transcardially perfused with phosphate buffered saline followed by 10% neutral buffered formalin. Brains were removed, post-fixed overnight, and dehydrated in 30% sucrose before coronal sectioning (30 μm) using a freezing microtome (Leica). Immunostaining was performed as previously described in Ref. [40] using primary antibodies for pST3S3 (Cell Signaling #9145, rabbit, 1:250), GFP (Aves Labs #GFP1020, chicken, 1:50), VIP (Phoenix #H06416, rabbit, 1:1000), and CRF (Phoenix #H01906, rabbit, 1:500). All antibody reactions were reacted with species-specific Alexa Fluor-488 or -568 conjugated (Invitrogen, 1:200) secondary antibodies or processed via avidin-biotin/diaminobenzidine (DAB) method (ABC kit, Vector Labs; DAB reagents, Sigma), and imaged as previously described in Ref. [41]. DAB images were pseudocolored using Photoshop software.

2.5. Phenotyping of LeprPdynKO and control mice

LepRbPdynKO and control littermates were weaned into individual housing at 21 days and fed either chow (Purina Lab Diet 5001) or high
fat diet (Research Diets D12492, 60% kcal from fat). Body weight was monitored weekly. A fasted (24 h) blood glucose sample was taken at 12–14 weeks of age. Analysis of body fat and lean mass was performed between 12 and 14 weeks of age using an NMR-based analyzer (Minispec LF90i, Bruker Optics). We also analyzed a subset of mice (13–16 weeks old) for oxygen consumption (VO₂), food intake, and locomotor activity using the Comprehensive Laboratory Animal Monitoring System (CLAMS, Columbus Instruments). Insulin was assessed using a double-antibody radioimmunoassay using an 125I-Human insulin tracer (EMD Millipore), a rat insulin standard (Novo), a guinea pig anti-rat insulin first antibody (EMD Millipore), and a sheep anti-guinea pig gamma globulin-PEG second antibody (MDRTC). Leptin was assayed by commercial ELISA (EMD Millipore). No significant differences were detected between the control (Pdyn<sup>fl/fl</sup> Lepr<sup>−/−</sup> and Lepr<sup>fl/fl</sup> groups) at the conclusion of the study and thus the data from these groups was combined for subsequent analysis.

### 2.6. Statistics
Physiological data are reported as mean ± SEM. Statistical analysis of physiological data was performed using Prism (version 6.0) software. Unpaired t-tests were used to compare results between groups of two. Body weight gain between genotypes was analyzed by two-way ANOVA. p < 0.05 was considered statistically significant.

### 3. RESULTS

#### 3.1. Lepr<sup>fl/fl</sup>GFP-L10a mice for profiling LepRb neurons

We generated Lepr<sup>Cre</sup>Rosa26GFP-L10a (Lepr<sup>fl/fl</sup>GFP-L10a) animals that express eGFP-L10a in LepRb neurons (Figure 1A). To confirm the expression of eGFP-L10a in Lepr<sup>fl/fl</sup> neurons in the hypothalamus and brainstem of these animals, we examined eGFP-L10a and its colocalization with phosphorylated STAT3 (pSTAT3; a marker of Leprb signaling) in brain sections from leptin-treated animals. As for Leprb reporter strains ([11,12,40], eGFP-immunoreactivity (<IR> and pSTAT3-IR colocalized in hypothalamic and brainstem regions known to contain Leprb (Figure 1B–G).

#### 3.2. TRAP-seq analysis of hypothalamic and brainstem LepRb neurons

We performed anti-eGFP TRAP on hypothalamic extracts from Lepr<sup>fl/fl</sup>GFP-L10a mice and used the resultant mRNA (as well as TRAP-depleted supernatant RNA) to generate multiplexed libraries for sequencing on the Illumina HiSeq2000 platform. We sequenced five independent samples, each containing material from the pooled hypothalami of 4–6 Leprb<sup>GFP-L10a</sup> mice. To validate the derivation of TRAP mRNA from Leprb neurons, we examined the enrichment of genes known to be expressed (or not expressed) in Leprb neurons in TRAP-derived relative to TRAP-depleted sequences (Figure 1H). TRAP-depleted material was chosen as the reference data for normalization in order to increase the power of the enrichment analysis by depleting transcripts from Leprb neurons from this reference material. This analysis confirmed the enrichment of all examined markers of Leprb neurons known to express Leprb (including Lepr, Pomc, and Agrp) [25,26,29,30,42–46] in TRAP mRNA relative to depleted supernatant. Furthermore, transcripts known to be expressed exclusively in non-Leprb neurons (e.g., Pinch, Hort, Oxt, Ginfrh) [40,47,48] were enriched in the TRAP-depleted samples relative to TRAP mRNA. Thus, TRAP-seq identified genes preferentially expressed in Leprb cells by comparison to TRAP-depleted (predominantly non-Leprb) samples.

The quantification of gene expression by next generation RNA sequencing not only permits the comparison of relative expression between two conditions but also defines the level of expression for each mRNA species since the frequency with which each sequence is detected reflects its abundance within the overall sample. In addition to revealing expression levels, these frequency data also enhance the statistical power of relative expression analysis compared to the single observation derived from each sample subjected to microarray analysis, permitting us to identify >1100 genes significantly enriched (>1.5-fold) in hypothalamic Leprb (TRAP) mRNA relative to non-Leprb (TRAP-depleted) mRNA (Supplemental Table 1).

Although brainstem Leprb neurons constitute a significant percentage of all Leprb neurons, even less is known about these cells than for hypothalamic Leprb neurons. We thus also dissected brainstem tissue from Leprb<sup>GFP-L10a</sup> mice and performed TRAP-seq on two independent samples, each derived from the brainstems of 6–8 animals. We chose to sequence only two samples due to the increased number of animals required to prepare brainstem material, since the list of enriched transcripts in the hypothalamus was essentially the same for n = 2 and n = 4 samples (data not shown). This analysis revealed ~900 genes that were significantly enriched in brainstem Leprb RNA relative to non-Leprb RNA (Supplemental Table 2). Our analysis of the non-Leprb (TRAP-depleted) and Leprb (TRAP) mRNA also defined genes that were enriched in non-Leprb cells of the hypothalamus (~1800 genes) and brainstem (~900 genes) relative to Leprb cells (Supplemental Tables 3 and 4).

To understand the common properties of brain cells that express Leprb (as well as those that do not contain Leprb), we identified genes enriched in both hypothalamic and brainstem TRAP (and TRAP-depleted) mRNA (Supplemental Tables 5 and 6). To characterize the types of cells contributing to TRAP- and TRAP-depleted mRNA, we plotted fold enrichment for hypothalamic and brainstem-derived mRNA to reveal the genes most highly enriched in common between the two sites (Figure 2). The genes most highly enriched in hypothalamic and brainstem TRAP mRNA included a number of neuropeptides (Pnpy, Ucn, Prok2, Ghrh, Cartpt, Tac1) and markers of dopaminergic (DA) neurons (Slc6a3, Th), as well as some markers for subsets of vasculature-associated cells (Fit1, Aab1b4). Some cell surface receptors and intracellular signaling proteins known to be expressed in neurons important for the control of metabolism (Mec3, Gucy2c, Fabp7, Cryab, Ermm), immune cells (Lita), and neural progenitor cells (Efh1d, Cad1, Ppp1r14a). Thus, while some studies have suggested that a variety of non-neuronal cells express Leprb and respond directly to leptin [49,50], TRAP mRNAs from Leprb<sup>GFP-L10a</sup> animals are enriched for transcripts from differentiated neurons (especially peptidergic and DA neurons) and vasculature-associated cells relative to these other cell types.

#### 3.3. TRAP-seq identifies novel subpopulations of hypothalamic and brainstem Leprb neurons

The mRNA species that are both highly expressed and highly enriched in Leprb cells presumably represent the most functionally relevant genes within a specific class of proteins. Thus, to identify the neuropeptides most likely to be functionally relevant in Leprb neurons, we examined expression level and fold enrichment for TRAP-enriched
transcripts that encode secreted proteins as defined by gene-ontology (GO) analysis on the UniPROT platform [39] (Figure 3). The secreted protein-encoding transcripts that were enriched in hypothalamic LepRb neurons included peptides that define known subpopulations of hypothalamic LepRb neurons (e.g., Pomc, Agrp, Prl, Nts, Gal) (Figure 3A). In addition to revealing the enrichment of Resp18 (a marker of peptide-secreting cells), this analysis also identified a number of highly expressed and enriched neuropeptide-encoding genes not previously examined as potential markers for subpopulations of LepRb neurons, including tachykinin-1 (Tac1), prodynorphin (Pdyn), corticotrophin releasing hormone (Crt), and growth hormone releasing hormone (Ghrh).

The genes that encode secreted proteins that were highly expressed and enriched in brainstem LepRb neurons included two neuropeptide-
encoding transcripts previously shown to be expressed in brainstem LepRb cells, Ucn and Cck [16,17,51] (Figure 3B). Our analysis also revealed the expression of a number of additional neuropeptides (including Tac1 and vasoactive intestinal peptide (Vip)) that were highly expressed and enriched in brainstem LepRb neurons.

Analysis of the transcripts encoding secreted proteins from TRAP-depleted (non-LepRb) mRNA in the hypothalamic revealed genes that encode neuropeptides found exclusively in non-LepRb neurons (e.g., Pnoc, Oxt, Avp, Hcrt, Ghrh1) and in pituitary gonadotrophs and lactotrophs (Lh, Gsa, Ph) (presumably derived from pituitary material that contaminated the hypothalamic tissue). In both hypothalamus and brainstem, TRAP-depleted transcripts encoding secreted proteins were also enriched for markers for glial (Apeo, Apod, Apoc1, Tlr, Igf2, Ptgs2, Scr1, Metn, Sparc) and immune (Ii33, Ly86) cells (Figure 3C,D). These findings are consistent with the data above (Figure 2B), which suggest the failure to recover mRNA from many non-neuronal cell types by LepRb specific TRAP.

3.4. Confirmation of novel neuropeptide-expressing subpopulations of LepRb neurons

To confirm the expression of Pdyn and Tac1 in LepRb neurons, we crossed Pdynox/ox and Tac1ox/ox mice onto the cre-inducible Rosa26RFP background, generating animals that express eYFP in Pdyn and Tac1 cells (PdynRFP and Tac1RFP mice, respectively) to examine their potential expression of LepRb (Figure 4; Supplemental Figure 1). This analysis revealed the colocalization of leptin-stimulated pSTAT3-IR with Pdyn and Tac1 in largely distinct sets of hypothalamic neurons: LepRbPdyn cells lie primarily in the ARC, VMH, and DMH; few LepRbPdyn cells were found in the LHA or ventral pre-mammillary nucleus (PMv) (Figure 4A–G). In the ARC, approximately 40% of LepRbPdyn cells also contain POMC (data not shown). In contrast, hypothalamic LepRbTac1 cells were detected primarily in the LHA and PMV; fewer LepRbTac1 cells were found in the DMH, and LepRbTac1 neurons were absent from the ARC and VMH (Supplemental Figure 1). We also found that leptin-stimulated pSTAT3-IR and eYFP in Tac1RFP mice colocalized in both brainstem areas: the NTS and the ventral lateral PAG.

Additionally, immunostaining for CRH or VIP peptide and eYFP in brain sections from colchicine-treated LepRbPdynKO (LepRbRFP/°) reporter mice revealed that LHA LepRb neurons contain CRH-IR (Supplemental Figure 2), and VIP-IR colocalized with LepRb in the brainstem DR and PAG nuclei of colchicine-treated LepRbRFP mice (Supplemental Figure 3).

3.5. Function of LepRbPdyn neurons

We hypothesized that TRAP-seq would identify physiologically relevant markers of LepRb subpopulations. We chose to examine the potential function of LepRbPdyn neurons, since leptin and nutritional status modulate Pdyn expression and activity (by the criterion of c-Fos accumulation) of Pdyn neurons, and since Pdyn expression overlaps with POMC cells in the ARC [52–54]. To examine the role for LepRbPdyn neurons in leptin action, we generated Pdynox/ox;LepRbtac1floxflox (LepRbPdynK0) animals (Figure 4A) along with Pdynox/°;LepRbtac1floxflox (control) littermates. Leptin-stimulated pSTAT3-IR was largely ablated from Pdyn-expressing neurons in the VMH and DMH (along with the small populations of LepRbPdyn cells in the LHA and PMV) of LepRbPdynKO mice and was reduced approximately 50% in ARC LepRbPdyn neurons (Figure 4H). The reason underlying the incomplete penetrance of Pdynox/°-mediated excision of LepRb in ARC LepRbPdyn cells is unclear, but similar idiosyncratic deletion patterns have been observed with other combinations of cre/flox alleles, including for LepRb [51]. Thus, LepRbPdynKO mice display LepRb ablation from most hypothalamic LepRbPdyn cells; roughly 7200 total neurons display LepRb disruption in these mice, ~50% of which lie in the VMH; most of the remainder are distributed between the ARC and DMH (~20% of the total disrupted LepRb neurons each).

We detected no alterations in body weight or body composition for chow-fed LepRbPdynKO male mice compared to controls (Figure 5A,B). Weekly chow intake was also unchanged (data not shown). Body weight and chow intake were also unchanged in female LepRbPdynKO mice compared to controls (data not shown), however LepRbPdynKO females had increased adiposity on a chow diet compared to controls (8.1% vs 6.7%, p = 0.02). 24-h fasting induced weight loss was not different between LepRbPdynKO mice or controls of either sex (data not shown). Also consistent with an underlying defect in energy homeostasis in LepRbPdynKO mice, male LepRbPdynKO mice gained significantly more weight and adiposity than controls when challenged with 9 weeks of high fat diet (HFD) (Figure 5A,B). As expected, based upon their increased fat mass, circulating leptin concentrations were higher.
in LepRbPdynKO mice (Figure 5C). While insulin concentrations trended up in ad libitum-fed LepRbPdynKO mice, this increase was not significant; fasted glucose levels were not different (Figure 5D,E). While food intake assessed over three days in CLAMS was similar between high fat-fed LepRbPdynKO mice and controls (Figure 5F, 24 h data shown), calorimetry revealed decreased energy expenditure (VO2) in the HFD-fed LepRbPdynKO mice compared to controls (Figure 5G,H). While we did not have sufficient numbers of animals to analyze the data by ANCOVA, the finding that VO2 was decreased in the LepRbPdynKO mice compared to controls whether the data were normalized to body weight or not reinforces the conclusion of diminished energy expenditure in the LepRbPdynKO mice. Thus, leptin action via the Pdyn-sensitive LepRbPdyn neurons promotes energy expenditure, rather than controlling food intake, to modulate overall energy balance. Furthermore, these data confirm the identification of functionally relevant population of LepRb neurons by TRAP-seq.

4. DISCUSSION

We examined gene expression in brain LepRb neurons, revealing over 1100 mRNAs that are enriched in hypothalamic LepRb cells and approximately 900 genes whose expression is enriched in brainstem LepRb cells. These LepRb enriched transcripts encode a variety of neuropeptides and other classes of proteins found primarily in peptidergic neurons (along with genes characteristic of DA neurons and an uncharacterized set of cells that express vascular markers). Most of these genes were not previously known to be expressed/enriched in LepRb neurons. Translational profiling of these cells thus revealed markers for previously unrecognized subpopulations of LepRb neurons, as well as genes of many classes that are likely to be important for the function of LepRb cells (and thus, for the control of body weight and metabolism). We have demonstrated that one of these populations (LepRbPdyn cells, contained primarily in the VMH, DMH and ARC) plays a crucial role in leptin-regulated energy balance through the control of metabolic rate/energy expenditure.

4.1. Validity of LepRb-specific TRAP-seq

The enrichment of genes encoding known LepRb-expressed mRNAs in our TRAP samples (e.g., Pomc, Agnp, Pthr, Nls, Gal, etc.) [25,26,29,30,42-46], together with our verification that many TRAP-enriched transcripts not previously known to be expressed in LepRb neurons colocalized with LepRb neurons, demonstrates the accuracy/specificity of the TRAP-seq method for cell type-specific transcriptome analysis. Although Pdyn expression and the activity of Pdyn neurons have previously been shown to be regulated by leptin [52,54], this regulation has never been shown to be cell-autonomous. Furthermore, while some ARC LepRbPdyn neurons contain POMC, LepRbTac1 and LepRbCRH neurons were absent from the ARC and thus distinct from both POMC and AgRP cells. Hence, TRAP-seq not only revealed previously unknown neuropeptide transmitters employed by discrete populations of LepRb neurons, but also defined markers for previously unrecognized subsets of LepRb cells.

4.2. Limitations of TRAP-seq

Markers for some previously defined sets of hypothalamic LepRb neurons [18,19], such as Nos1 (~1.5-fold enriched) and Sla32a1 (vGat, ~1.2-fold enriched), were poorly enriched. This presumably reflects the wide expression of Nos1 and Sla32a1 throughout the
hypothalamus in both LepRb and non-LepRb neurons. Thus, TRAP-seq more robustly reveals the enrichment of genes with restricted expression (e.g., neuropeptides) than it does widely expressed genes, since the fold enrichment of narrowly expressed genes in TRAP mRNA is enhanced by comparison to the relative dilution of these genes within the tissue as a whole. Hence, the mRNAs that can be unambiguously assigned to the LepRb neuron transcriptome are biased toward genes more highly expressed in LepRb cells than other cells in the tissue, and this method is less sensitive for widely expressed transcripts that are also found in some LepRb neurons. The genes that are highly enriched in TRAP mRNA, however, often represent the best markers for circumscribed, functionally related, populations of cells that are tractable for circuit analysis.

Figure 4: Pdyn expression defines a distinct subpopulation of LepRb neurons. (A) Pdyn<sup>YFP</sup> mediates the excision of the transcription-blocking cassette from Rosa26<sup>YFP</sup> in LepRb cells, promoting the expression of eYFP in these cells. Pdyn<sup>YFP</sup> can also mediate the excision of exon 17 from Lep<sup>flox</sup> mice resulting in the ablations of LepRb from Pdyn cells (Lep<sup>RbPdynKO</sup> mice). (B–F) Representative images showing colocalization of pSTAT3- (purple) and GFP-IR (green; detects eYFP) in the ARC (B), VMH (C), DMH (D), LHA (E), and PMv (F), of Pdyn<sup>YFP</sup> mice following leptin treatment (5 mg/kg, i.p., 90 min). (G) Cells containing both Pdyn (eYFP-IR) and LepRb (pSTAT3-IR) were quantified in the hypothalamic regions shown (plotted as mean ± SEM). (H) Colocalization of pSTAT3 and Pdyn was reduced in Lep<sup>RbPdynKO</sup> mice on the reporter background relative to control (C) Pdyn<sup>YFP</sup> mice (mean ± SEM is shown; *p < 0.05 by t-test). Arrows indicate double labeled cells. Scale bar = 100 μm. Cell counts were performed on serial sections (1:4) from n = 3 control and n = 2 Lep<sup>RbPdynKO</sup> mice treated with leptin (5 mg/kg, i.p., 90 min). For (G), cell counts were multiplied by 4 to approximate total hypothalamic cell numbers. NEO, neomycin selection cassette; black arrow heads, LoxP sites; pA, polyadenylation site.
that some cell types (including other neurons) might express low levels of LepRb, however, if cre expression from Leprcre and/or reporter expression from Rosa26eGFP-L10a is too low to detect in such cells. Thus, our analysis is necessarily biased towards those cell populations of which have active expression at both the Lepr and Rosa26 locus. In either case, our TRAP-seq analysis appears to be specific, if not perfectly sensitive, for neurons (and some vasculature-associated cells) that contain LepRb.

4.3. Hypothalamic and brainstem LepRb neurons

Interestingly, while some genes were enriched in both hypothalamic and brainstem LepRb neurons, many highly enriched neuropeptide encoding mRNAs were found in LepRb neurons from only one region. Indeed, while a few neuropeptide encoding genes enriched in brainstem LepRb neurons (including Tac1 and Cartp) were also found in populations of hypothalamic LepRb cells, brainstem and hypothalamic LepRb neurons contain relatively distinct sets of highly expressed neuropeptide transmitters, as most neuropeptide encoding genes that are enriched in hypothalamic LepRb neurons are absent from brainstem LepRb cells. Similarly, Cck [16,51] and Vip are enriched and expressed in brainstem, but not hypothalamic, LepRb neurons. These findings support the concept of discrete functions for individual groups of LepRb neurons and their uniquely expressed gene products [10].

4.4. Function of LepRbPdyn neurons

Of the novel subsets of LepRb neurons that we identify here, we investigated the role for leptin action on LepRbPdyn neurons. LepRb ablation from LepRbPdyn neurons in LepRbPdynKO mice decreased...
energy expenditure to increase adiposity during exposure to HFD. Thus, the Pdyn sensitive LepRb neurons promote energy expenditure in response to leptin. Approximately half of the LepRb-Pdyn cells from which LepRb was ablated in LepRb-PdynKO mice lie in the VMH, and almost all VMH LepRb neurons contain Pdyn. It is technically challenging to assess the degree of colocalization between the LepRb-Pdyn and LepRbSPfi populations; both appear to comprise ~80% of all VMH LepRb neurons, suggesting a minimum of 60% of VMH LepRb neurons are affected in both models [25,26]. The consistent phenotype observed between these two models of VMH LepRb ablation, however, both serve to highlight the known role for the VMH in the control of energy expenditure, including in response to leptin [25,26]. Non-VMH LepRbSPfi cells (primarily in the ARC and DMH) also may contribute to the modulation of energy expenditure, but they do not appear to play major additional roles in the control of energy balance.

4.5. Neuropeptides in LepRb neurons

In addition to providing markers for LepRb subpopulations, the products of transcripts enriched in LepRb neurons are likely to play important functional roles in LepRb neurons. While the contribution of POMC and AgRP peptides to leptin action has been well described, the metabolic functions of the majority of the neuropeptides identified by TRAP-seq (including dynorphin) are as yet undetermined. Furthermore, not only do LepRb-expressed neuropeptides presumably contribute to the effects of leptin on neurotransmission but also the GPCRs, transcriptional regulators, signaling proteins, and other classes of proteins enriched in LepRb neurons likely contribute to the function of these cells and thus to the control of energy balance and metabolism. Some of these LepRb enriched gene products may constitute potential targets for therapeutic intervention in obesity and diabetes.

4.6. Conclusions

Overall, by elucidating the transcriptomes of brainstem and hypothalamic LepRb neurons, our TRAP-seq analysis reveals markers for numerous subpopulations of LepRb neurons, along with genes likely to contribute importantly to central leptin action. In the future it will be important to define the roles in leptin action and metabolic control for new subpopulations of LepRb neurons and for a variety of gene products that are highly enriched in LepRb neurons. The LepRb expressed genes defined by our TRAP-seq analysis thus provide an important resource for these and other future investigations.

AUTHOR CONTRIBUTIONS

MA, CP, and DO researched data, wrote and edited manuscript; MH, MK, and BL reviewed/editd manuscript. MM wrote and edited manuscript.

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CONFLICT OF INTEREST

The authors declare that they have no conflicts of interest relevant to this manuscript.

APPENDIX A. SUPPLEMENTARY DATA

Supplementary data related to this article can be found at http://dx.doi.org/10.1016/j.molmet.2015.01.012.

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