Somato-Dendritic Localization and Signaling by Leptin Receptors in Hypothalamic POMC and AgRP Neurons

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Abstract

Leptin acts via neuronal leptin receptors to control energy balance. Hypothalamic pro-opiomelanocortin (POMC) and agouti-related peptide (AgRP)/Neuropeptide Y (NPY)/GABA neurons produce anorexigenic and orexigenic neuropeptides and neurotransmitters, and express the long signaling form of the leptin receptor (LepRb). Despite progress in the understanding of LepRb signaling and function, the sub-cellular localization of LepRb in target neurons has not been determined, primarily due to lack of sensitive anti-LepRb antibodies. Here we applied light microscopy (LM), confocal-laser scanning microscopy (CLSM), and electron microscopy (EM) to investigate LepRb localization and signaling in mice expressing a HA-tagged LepRb selectively in POMC or AgRP/NPY/GABA neurons. We report that LepRb receptors exhibit a somato-dendritic expression pattern. We further show that LepRb activates STAT3 phosphorylation in neuronal fibers within several hypothalamic and hindbrain nuclei of wild-type mice and rats, and specifically in dendrites of arcuate POMC and AgRP/NPY/GABA neurons of Leprb+/− mice and in Leprb+/- mice expressing HA-LepRb in a neuron specific manner. We did not find evidence of LepRb localization or STAT3-signaling in axon-fibers or nerve-terminals of POMC and AgRP/NPY/GABA neurons. Three-dimensional serial EM-reconstruction of dendritic segments from POMC and AgRP/NPY/GABA neurons indicates a high density of shaft synapses. In addition, we found that the leptin activates STAT3 signaling in proximity to synapses on POMC and AgRP/NPY/GABA dendritic shafts. Taken together, these data suggest that the signaling-form of the leptin receptor may therefore play an important role in leptin’s central effects on energy balance, possibly through modulation of synaptic activity via post-synaptic mechanisms.

Introduction

Leptin, an adipocyte-derived hormone, acts on the central nervous system to regulate energy balance, glucose metabolism and neuroendocrine actions by activating the long signaling form of the leptin receptor (LepRb) [1-4]. Among several leptin-responsive brain regions, the arcuate nucleus of the hypothalamus (Arc) serves an important role in mediating these leptin actions. Within the Arc there are several distinct subsets of key LepRb-expressing neurons, including the anorexigenic pro-opiomelanocortin (POMC)-producing neurons and the orexigenic Agouti-related peptide (AgRP)/Neuropeptide Y (NPY)/γ-aminobutyric acid (GABA)-producing neurons [5-7]. Recent studies employing optogenetics and pharmacogenetic methods to modulate the activity of those neurons in vivo further demonstrate potent effects on feeding behavior [8-10]. Leptin stimulates c-Fos expression and increases firing rates of POMC neurons [11,12]. In contrast, the action potential frequency of AgRP/NPY/GABA neurons is depressed by leptin [13,14]. Consistent with these actions, fasting, a state of low circulating leptin concentrations, leads to opposite effects on action potentials on these two neuronal populations [15,16]. Mice lacking leptin receptors only in POMC or AgRP neurons exhibit increased body weight, demonstrating that both groups of cells are required for maintenance of normal metabolic homeostasis.
AgRP/NPY/GABA cells, supporting the notion that diet-induced defects in LepRb signaling within these neurons may play a primary role in the development of obesity [4,19-21].

Leptin has structural homology to cytokines and the leptin receptor bears strong sequence similarity to the class I cytokine receptor super family [22,23]. Moreover, the signaling capabilities of LepRb show direct similarities to that of the signaling subunits of the interleukin 6 (IL-6)-cytokine receptor, leukemia inhibitory factor receptor (LIFR), and ciliary neurotrophic factor receptor (CNTFR) [24-27]. In particular, LepRb activates intracellular signaling actions through cytokine-receptor-like pathways including the canonical JAK2-STAT3 pathway [28,29]. Leptin-dependent phosphorylation of the STAT3 transcription factor is a highly specific CNS cellular marker for identification of LepRb-expressing neurons in vivo [19,30] and STAT3 activation by LepRb is critically important for normal energy balance regulation [31,32].

LepRb is required for leptin-mediated cell-autonomous effects on membrane potentials and axonal firing of hypothalamic neurons [33,34]. Furthermore, studies have shown modulation of excitatory Ca2+-currents via post-synaptic LepRb-dependent mechanisms [35,36]. Yet other evidence suggests that pre-synaptic actions of LepRb can modulate glutamate release onto dopamine neurons in the ventral tegmental area (VTA) [37] and influence GABA-release from AgRP/NPY/GABA neurons onto hypothalamic POMC neurons [12].

Despite progress in the understanding of neuronal leptin receptor signaling and of leptin’s metabolic actions via arcuate POMC and AgRP/NPY/GABA neurons, direct evidence of the sub-cellular localization of LepRb within soma and neuronal fiber compartments is not known. We show here that LepRb is expressed and signals in somato-dendritic compartments of both groups of neurons. In addition, we present evidence of LepRb signaling in close proximity to post-synaptic structures on dendritic shafts. These studies suggest that leptin receptor signaling in dendrites may be important for leptin’s effect on energy balance, possibly by modulating neuronal excitability through post-synaptic signaling mechanisms.

Materials and Methods

Ethics Statement

Animal care and procedures were approved by the Institutional Animal Care and Use Committee at Beth Israel Deaconess Medical Center.

Mice and Rats

Mice and rats were housed at 22–24°C using a 14 hr light/10 hr dark cycle with standard chow diet (Teklad F6 Rodent Diet 8664, Harlan Teklad, Madison, WI). Transgenic mice with cre-dependent activation of HA-tagged LepRb expression (HA-Leprb flox)(prior name: HA-ObRb STOP) were described previously [38]. Briefly, Pomc-cre [17] and Agrp-ires-cre mice [39] were kindly supplied by Dr. B. Lowell (BIDMC, Boston, MA) and Pomc-EGFP mice were described earlier [12]. EGFP flox (Z/EG) reporter mice (B6.129(Cg)-Tg(CAG-Bgeo/GFP)21Lbe/J) were purchased from Jackson Laboratories (Bar Harbor, ME)[40]. Leprdb-Pomc-cre;HA-Leprb flox and Leprdb-Agrp-ires-cre;HA-Leprb flox mice that selectively express HA-LepRb in POMC or AgRP neurons respectively, were generated by crossing the appropriate cre/flox lines with Leprdb+/- mice, as we have described previously [38]. The above animals were on mixed genetic backgrounds (FVB; C57BL/ KSj; C57BL/6J; 129). Adult male wild type C57BL/6J mice and male Sprague Dawley rats were purchased from Jackson laboratories.

Materials

Murine leptin was purchased from Dr. A.F. Parlow (National Hormone & Peptide Program, Torrance, CA). Supplies for immunohistochemistry (IHC) were purchased from Sigma-Aldrich (St. Louis, MO), apart from the ABC Vectastain Elite kit which was purchased from Vector Laboratories (Burlingame, CA) and the tyramide-signal-amplification kit (TSA) which was purchased from Perkin Elmer (Waltham, MA). The phospho-specific-(Y705)-STAT3 rabbit antibody was from New England Biolabs (Beverly, MA) and the anti-HA mouse antibody from Covance Inc. (Berkely, CA). The rabbit anti-POMC precursor (against amino residues 27-52) antisera was obtained from Phoenix Pharmaceuticals (Burlingame, CA), the rabbit anti-β-endorphin antibody was a kind gift from Dr. Ronnekleiv (Oregon Health and Science University, Portland, OR)[41] and the sheep anti-α-MSH antisera was from Chemicon International, Inc. (Temecula, CA). The chicken polyclonal anti-GFP and the mouse monoclonal anti-PSD95 antibodies were purchased from Abcam (Cambridge, MA). Biotinylated secondary antibodies were from Jackson Immunology Research Laboratories (West Grove, PA) and secondary fluorescent immunoglobulin conjugates were obtained from Molecular Probes (Eugene, OR). The plasmid encoding murine LepRb was described earlier [42,43]. The HA-LepRb expression vector was created by standard procedures and ultimately cloned into the mammalian expression vector pcDNA3.1 from Invitrogen (San Diego, CA). As we have described earlier, the DNA sequence encoding the 9 amino acid HA tag was inserted after codon 46 of the murine LepRb cDNA [38]. The EGFP expression vector was from Clontech Laboratories, Inc. (Mountain View, CA).

Immunohistochemistry (IHC)

Transcardiac perfusion and fixation with paraformaldehyde, removal of brains, postfixation, and cryoprotection were performed as we have generally described previously [19,21,38]. Brains were cut in 25-35 µm thick coronal sections, collected in 4-5 series, and stored at -20°C until further use. For non-fluorescent light microscopy, free-floating brain sections were incubated with biotinylated secondary antibodies, followed by avidin-biotin complex labeling, and developed with nickel-diaminobenzidine (DAB), generating a brown-black precipitate for analyses by light microscopy (LM) [19,30]. For fluorescence microscopy, the free-floating sections were incubated with fluorescent-labeled secondary antibodies [44,45]. In the case of HA immunofluorescence, TSA procedures were applied to enhance sensitivity (Perkin Elmer). DAB results were visualized using bright-field light and
photomicrographs captured with a digital AxioCam camera (Carl Zeiss, Thornwood, NY) mounted on a Zeiss microscope (Axioskop2) or a Nikon Eclipse 80i equipped with a Nikon DS-U1 digital camera (Nikon Instruments Inc., Melville, NY). Fluorescent results were captured and analyzed by confocal laser scanning microscopy (CLSM) using a Zeiss LSM510 system and the LSM Image Browser 4.2.0.121 free-ware. Adobe Photoshop software (Adobe, San Jose, CA) was used to crop images and to change file types for publication.

Electron Microscopy (EM)

For transmission EM, mouse brains were first fixed via transcardiac perfusion with saline followed by 1.0% formaldehyde and 1.25% EM grade glutaraldehyde (TedPella, Redding, CA) in phosphate buffer, pH 7.4 (PB). Brains were removed from the skull and post-fixed in 2.0% formaldehyde and 2.5% glutaraldehyde in PB at 4°C overnight. Following embedding in 3% LMP agarose, 100 µm thick coronal sections were cut on a vibratome and stored in 2.0% formaldehyde and 2.5% glutaraldehyde in PB at 4°C until further use (method modified from [46]). For pre-embedding EGFP immuno-EM, free floating sections were blocked with goat serum and incubated with anti-GFP antibodies overnight at room temperature [44]. For pre-embedding pSTAT3 immuno-EM, free floating brain sections were pretreated with NaOH/H2O2, glycine and SDS, blocked with goat serum, and incubated with anti-pSTAT3 antibodies overnight at room temperature, as described [30]. For DAB staining, the sections were incubated with biotinylated secondary antibodies, subjected to avidin-biotin complex labeling, and developed with nickel-diaminobenzidine (DAB) as described above under IHC. For pre-embedding pSTAT3 immuno-Gold-EM, free floating sections were first pretreated and treated with the anti-pSTAT3 antibodies as above, washed, and then incubated with anti-rabbit IgG conjugated to 1 nm gold particles (Amersham Biosciences Inc). Post-fixation, tissue embedding, cutting, placement on grids, and counter staining were performed by the Electron Microscope Facility at Harvard Medical School.

Immuno-stained tissue sections were silver enhanced with the InteSE kit (Amersham Biosciences), then postfixed in 0.5 % osmium tetroxide for 15 minutes, rinsed in water, dehydrated in graded ethanol solutions, transferred to propylene oxide, and embedded in epoxy resin (Epon-Araldite, EMS, Hatfield, PA) between two layers of Aclar plastic (Epon-Araldite, EMS, Hatfield, PA). After the plastic was polymerized at 60°C for 48 hrs, the area of the arcuate hypothalamic nucleus sections were first pretreated and treated with the anti-pSTAT3 antibodies as above, washed, and then incubated with anti-rabbit IgG conjugated to 1 nm gold particles (Amersham Biosciences Inc). Post-fixation, tissue embedding, cutting, placement on grids, and counter staining were performed by the Electron Microscope Facility at Harvard Medical School.

Ultrafine structures in the central nervous system were identified as described by Peters et al. [49] and Stuart et al. [50]. Briefly, neuronal cell bodies were identified by their large cytoplasmic space when compared with dendritic shafts, the presence of multiple intracellular compartments, including the nucleus and organelles, and although not essential because of the orientation of the sections, the protrusion of large proximal dendrites from their membrane. Dendritic shafts were identified by having a cross section diameter of 0.5-2 µm with a relatively irregular shape, and often by the presence of a regular array of microtubules. Axon fibers were defined as either myelinated or not, smaller cross section diameter compared to dendritic shafts and tending to maintain a consistent convex cross section. In addition, axon fibers typically exhibit a denser and more organized array of microtubules relative to dendritic shafts. Gial fibers are highly irregular in shape, often containing glycogen granules, and further differ from dendritic shafts and axons by not having a regular array of microtubule fibers. Spines were defined as their relative small size (typically ~1-2 µm length) compared to the dendritic shaft and as any protrusion from the dendritic shaft that is not a dendrite branch. In addition, spines are usually devoid of mitochondria and microtubules. Asymmetric synapses form synaptic structures that are distinguished by a thickened, postsynaptic density (PSD) and a minor presynaptic density. In contrast, in symmetric synapses the pre-and postsynaptic membranes are typically more parallel than the surrounding non-synaptic membrane, and the synapse has a less prominent postsynaptic density that is similar in thickness to the pre-synaptic density.

Primary Hypothalamic and Hippocampal Neuronal Cultures

NeuroPure E18 primary rat hypothalamic cells (#N400200) and Neuropure E18 primary rat hippocampal cells (#N100200) were purchased from Genlantis (San Diego, CA) and were dissociated, plated and grown in 24 well dishes on poly-D-lysine-coated coverslips as generally directed by the manufacturer. After 4-7 days in vitro (DIV), neurons were transfected with LepRb expression vectors, or co-transfected with HA-LepRb and EGFP expression vectors using NeuroFECT (Genlantis) or Lipofectamine 2000 (Invitrogen) according to the manufacturer’s instructions. One to two days later, neurons were serum-deprived for 3-12 hours and treated with 100 nM recombinant leptin or vehicle control solution. Cells were then fixed and the coverslips were subjected to pSTAT3- or HA-immunocytochemistry (ICC) as we have generally described previously [30,38].

In situ Hybridization

Free-floating brain sections were generated as described for IHC (above) and processed for in situ as described earlier [21]. Briefly, digoxigenin (DIG) labeled RNA probes were generated by PCR amplification of mouse hypothalamic CDNA utilizing sets of POMC or NPY primers that included SP6 or T7 end-sequences. Specifically, for POMC we used a 628 base long
probe (bases 147-775 of POMC mRNA (NCBI: NM_008895)) and for NPY a 308 base long (bases 67-376 of NPY mRNA (NCBI: NM_023456)). Purified PCR products were then subjected to in vitro RNA synthesis of sense or antisense RNA’s using appropriate SP6 or T7 polymerases and DIG labeling mix. Briefly, sections were mounted on Superfrost Plus slides (Fisher, Hampton, NH) and hybridized overnight with the DIG-labeled mouse Pomp anti-sense RNA probe or with the DIG-labeled mouse Npy anti-sense RNA probe, both at 0.6 µg/ml at 60°C. All brain sections were washed twice in 0.2X SSC at 60°C, blocked in PBS with 10% bovine serum, and reacted with anti-DIG antibodies fused to alkaline phosphatase (Roche, Nutley, NJ)(1:5000, 10% serum, 2 hours at room temperature). Sections were washed and incubated with alkaline phosphatase substrate (NBT/BCIP, Roche, Nutley, NJ) producing a color precipitate. The reaction was stopped by addition of EDTA.

Results

Leptin activates STAT3 phosphorylation in neuronal fiber processes in hypothalamic nuclei of mice and rats

When we first developed the immunohistochemical (IHC) assay for phosphorylated STAT3 (pSTAT3) in brain sections from leptin-treated rodents [19,30] we noted pSTAT3 immunoreactivity (IR) in many neuronal fiber processes in addition to the expected nuclear staining. This observation and the relationship to Lepr sub-cellular localization was further investigated here. Figure 1 shows microphotographs captured by light microscopy (LM) of coronal brain sections of the medio-basal hypothalamus from a vehicle and a leptin-treated wild type mouse. While almost no pSTAT3 IR is present in the vehicle-treated animal, robust nuclear staining is observed in many nuclei known to express LepRb, including the LHA, VMH, DMH, Arc, PH and the PMN (Figure 1). In contrast to rats [51], pSTAT3 IR nuclei were not detected in the paraventricular hypothalamic nucleus (PVH) of leptin-treated mice (Figure S2). Phospho-STAT3 is an established cellular marker for first-order Lepr-expressing neurons and is not activated in second-order neurons [30]. In addition to the expected nuclear staining described above, pSTAT3 IR was also observed within neuronal fiber processes (Figure 2). These fibers were largely confined to each brain nucleus where Lepr is known to be present [52,53]. Similar fiber staining was also found within the nucleus of the solitary tract (NTS) in the caudal hindbrain (Figure S1). Because the pSTAT3 IR fibers did not extend much beyond each brain nucleus, STAT3 activation by leptin may be limited to dendritic fiber structures rather than axons which project far beyond each brain nucleus. In support of this possibility, pSTAT3 IR fibers were not found in the PVH of mice, a region known to receive dense axonal innervation from arcuate leptin-responsive neurons, including POMC and AgRP neurons (Figure S2)[6,54,55]. Similar anatomical patterns of STAT3 IR neuronal fibers were also seen in hypothalamic nuclei of leptin-treated Sprague Dawley rats, including the VMH, DMH and PMN (Figure S3). These data combined suggest that LepRb itself may be localized to neuronal fibers, possibly dendrites.

Leptin activates STAT3 phosphorylation in soma and fiber processes of hypothalamic POMC neurons of mice

To investigate whether STAT3 is activated in fiber structures of known Lepr-expressing hypothalamic neurons such as the POMC neurons, we next treated Pomc-EGFP mice with leptin and applied fluorescent-IHC and confocal laser scanning microscopy (CLSM) to hypothalamic brain sections. As shown in Figure 3, pSTAT3 IR is localized to the nucleus, cytoplasm and to proximal fibers of two POMC neurons (arrows). Similar data were obtained from the Arc of leptin-treated Npy-hrGFP mice (not shown). POMC (and AgRP/NPY/GABA) neurons might therefore serve as a suitable model system to investigate the sub-cellular fiber distribution of Lepr.

HA-tagged Lepr is localized to intracellular vesicular structures and to the plasma membrane of POMC somata, and to proximal fibers of POMC neurons

To date, sufficiently sensitive and specific anti-Lepr antibodies capable of detecting endogenous Lepr proteins in the rodent brain have yet to be reported. We therefore employed a genetic strategy to express HA-tagged Lepr in mice thus facilitating detection with well characterized sensitive monoclonal anti-HA antibodies. The genetic design included introduction of two loxP sites flanking a transcriptional blocking sequence upstream of the HA-Lepr cDNA to allow HA-Lepr expression in a cre-dependent manner (e.g. expression of HA-Lepr in POMC neurons by crossing HA-Lepr flox mice with Pomc-cre mice). The detailed strategy was described earlier [38]. Successful co-expression of HA-Lepr and POMC polypeptides in Pomc-cre;HA-Leprb flox mice is shown by immunofluorescence-IHC and CLSM in Figure S4.

To enable investigation of possible Lepr localization in neuronal fibers, we created mice expressing HA-Lepr and EGFP in POMC soma and fiber processes (Pomc-cre;HA-Leprb flox;EGFP flox). Figure 4A shows the cell body of a POMC neuron (GFP IR) expressing HA-Leprb (HA IR). A large fraction of somatic HA-Leprb IR is found in vesicular-like structures within the cytoplasm. In addition, a minor fraction of HA-Leprb appears localized to the plasma membrane of the POMC soma. A POMC cell body is represented as a collapsed Z-stack and is reconstructed in 3-dimensions in Figure 4B. A non-transparent 3-D surface representation is shown in red. A semi-transparent representation of the POMC soma is also depicted. Green represents HA-Leprb localization in larger vesicular-like structures. A large fraction of HA-Leprb was detected in peri-nuclear regions, possibly representing Golgi and trans-Golgi networks (orange). Similar results were obtained from 3-D reconstruction of one additional POMC neuron (not shown). In Figure 5, proximal neurites extending from a POMC neuron are shown. HA-Leprb is found in a punctuate pattern of all primary and secondary fibers. These results indicate that HA-Leprb is at least localized to dendritic fibers of POMC neurons (i.e. only one axon per cell).
In Figure S5A we show that HA-LepRb is expressed in proximal neurite fibers of primary hypothalamic neurons transfected with HA-LepRb plasmids. In addition, leptin activates STAT3 phosphorylation in neuronal fibers that contain the dendritic marker, PSD95 (Figure S5B), thus supporting a dendritic localization of LepRb.

**HA-LepRb is localized to dendrites, but not axons, of POMC neurons**

We next aimed to determine if HA-LepRb is expressed in axonal fibers of POMC neurons by investigating possible colocalization with neuronal POMC axon-markers using antibodies against the POMC-polypeptide precursor or the POMC-derived β-Endorphin neuropeptide [54,56]. By fluorescence-IHC and CLSM we show in Figure 6A a POMC cell body with a long fiber (>200 µm) that expresses HA-LepRb (green) in a punctuate pattern. This fiber is relatively thick,
does not show evidence of varicosities (boutons) and does not express the (red) POMC-polypeptide axonal marker (right enlargement), and is therefore likely a dendrite. In contrast, the POMC-precursor (red) can be found in the parent soma and in thin en-passant fibers with varicosities that are therefore likely axons (left column,). Furthermore, as shown in Figure 6B, HA-LepRb is not expressed in β-Endorphin positive fibers (white arrows) which exhibit bouton-like structures that are characteristic for axons. Based on these results, we conclude that the long-form leptin receptor is principally localized to the soma and dendrites, but not to axons of POMC neurons.

**Leptin activates STAT3 phosphorylation in dendrites, but not in α-MSH-containing axonal fibers of POMC neurons**

We next examined if leptin can stimulate down-stream signal transduction in dendritic, but not axonal, cellular compartments of POMC neurons. In order to restrict signaling events to POMC neurons, we generated Lepr<sup>db/db</sup>;Pomc-cre;HA-Leprb flox mice as we have described earlier [38]. These mice are 6-9 weeks old, obese and hyper-leptinemic [38]. In Figure S6, we applied IHC and LM to brain sections to first validate that leptin-treated Lepr<sup>db/db</sup>;Pomc-cre;HA-Leprb flox mice exhibit pSTAT3 activation only in the arcuate hypothalamic regions that are consistent with the well known anatomical location of POMC neurons. As expected, STAT3 phosphorylation was not found in leptin-treated Lepr<sup>db/db</sup> control mice. Importantly, leptin-treated Lepr<sup>db/db</sup>;Pomc-cre;HA-Leprb flox mice showed evidence of pSTAT3 IR in fiber processes (and POMC nuclei) (Figure S6), as we have shown in neuronal fibers of wild type mice (Figure 2) and in processes of POMC neurons of Pomc-GFP mice (Figure 3).

In brain sections from leptin-treated Lepr<sup>db/db</sup>;Pomc-cre;HA-Leprb flox mice we next applied fluorescent-IHC and CLSM to investigate fiber localization of pSTAT3 and the POMC-polypeptide-derived axonal marker, α-melanocyte stimulating hormone (α-MSH)[54,55,57,58]. As shown in Figure 7, pSTAT3 IR was observed in fiber processes that did not exhibit α-MSH IR. In contrast, α-MSH was localized to thin fibers with varicosities, consistent with being POMC axonal processes.
Combined, these results indicate that leptin activates STAT3 phosphorylation in POMC dendrites, but not in axons, consistent with somato-dendritic localization of LepRb.

**Ultrastructural analyses of STAT3 phosphorylation by leptin in dendrites of hypothalamic POMC and AgRP/NPY neurons**

To further validate and investigate LepRb localization and signaling in dendrites at the ultrastructural level of hypothalamic neurons, we applied immuno-electron microscopy (EM) methodology. Multiple attempts unfortunately failed to detect specific signals for HA-LepRb by immuno-HA EM, possibly due to antigen interference by glutaraldehyde in the EM fixative. We therefore focused these studies on investigations of the ultrastructural sub-cellular localization of leptin-induced STAT3 phosphorylation, as pSTAT3 immunostaining was not negatively affected by glutaraldehyde. These analyses were expanded to include investigations of the localization of leptin-induced pSTAT3 in AgRP neurons. To this end, brain sections from leptin-treated *Lepr*<sup>db/db</sup>;*Pomc*-cre;HA-*Leprb* flox mice and *Lepr*<sup>db/db</sup>;*AgRP*-ires-cre;HA-*Leprb* flox mice were subjected to pSTAT3 immuno-EM. Validation of proper anatomical localization of pSTAT3 IR consistent with the known location (ventro-medial Arc) of AgRP neurons in *Lepr*<sup>db/db</sup>;*AgRP*-ires-cre;HA-*Leprb* flox mice is shown by IHC and LM in Figure S7. Of note, *Lepr*<sup>db/db</sup>;*AgRP*-ires-cre;HA-*Leprb* flox mice (6-9 weeks old) are nearly as obese and hyperleptinemic as *Lepr*<sup>db/db</sup> controls (not shown). As in POMC fibers of *Lepr*<sup>db/db</sup>;*Pomc*-cre;HA-*Leprb* flox mice, pSTAT3 IR was also observed.
Figure 4. Localization of HA-tagged LepRb at the plasma membrane and cytoplasm of hypothalamic POMC neurons. A. CLSM of brain sections from a Pomc-cre;HA-Leprb flox;EGFP flox mouse. Shown is the soma of a hypothalamic POMC neuron expressing HA-LepRb (EGFP IR (red); HA IR (green); Nuclear DAPI (blue)). Within the cytoplasm, HA-LepRb is expressed in a vesicular/punctate pattern. Right: Enlargement of box area in A, depicting plasma membrane localization of HA-LepRb (arrows). Shown are single plane sections (0.42 µm). B. Left: CLSM was used to generate a collapsed Z-stack of the soma of a hypothalamic POMC neuron (EGFP IR (red)) expressing HA-LepRb (HA IR (green)). The Z-stack is ~16 µm deep comprising of 39 confocal sections. Right: The soma of the neuron is reconstructed in 3-dimensions using all 39 sections and the Reconstruct computer software. A non-transparent surface representation of the cell is shown in red (insert). A grey semi-transparent surface representation is also depicted. Blue is the nucleus (non-transparent). A number of vesicular-like structures that express HA-LepRb are shown in green. In addition, peri-nuclear localization of HA-LepRb in the presumed Golgi/trans-Golgi networks is represented in orange.

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in AgRP neuronal fibers of \textit{Lepr}^{db/db};AgRP-ires-cre;HA-Leprb \textit{flox} mice at the LM level (Figure S7).

Figure 8 shows pSTAT3 immuno-EM microphotographs at different levels of magnification from the medial Arc of a leptin-treated \textit{Lepr}^{db/db};Pomc-cre;HA-Leprb \textit{flox} mouse. In the left image, pSTAT3 IR neuronal nuclei (black arrow) and non-pSTAT3 IR nuclei (white arrow) can be seen. At higher magnifications, pSTAT3 IR dendritic shaft structures are identified, including dendrites in the photographic plane (top right). An example of a cross-section of a large dendrite with an array of microtubules (MT) and one mitochondrion (M) is presented at bottom right. In several brain sections from several grids, we identified random pSTAT3 IR structures and counted a majority of neuronal nuclei (N=49) and dendrites (N=70), and no axonal fibers or presynaptic structures (N=0). At this time-point after leptin treatment, immunostaining was
Figure 6. HA-LepRb is localized to dendrites, but not axons, of POMC neurons. A. Middle Image (*): CLSM was applied to depict a POMC neuron with a >200 µm long dendrite (4 arrow heads) in a brain section from a Pomc-cre;HA-Leprb flox mouse. POMC IR is shown in red and HA (LepRb) IR in green. To present the entire length of the dendrite, the image was merged from three different focal planes. Left Column: The POMC soma expresses both HA-LepRb and POMC polypeptides. None of the POMC axonal fibers (red arrows) in the field exhibit HA IR. Right Image. Magnification of box area depicting a fiber segment containing punctate HA-LepRb expression. POMC IR is not detectable in the fiber (dendrite). Shown are single confocal planes. B. Depicted are two POMC somata co-expressing β-endorphin IR (red) and HA-LepRb (HA IR (green)) in a brain section from a Pomc-cre;HA-Leprb flox mouse. Two nearby POMC IR-axonal fibers (arrows) do not express HA-LepRb. Nuclear DAPI is colored blue. These are single confocal planes.

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Figure 7. Leptin does not activate STAT3 phosphorylation in α-MSH-containing axonal fibers of POMC neurons. CLSM showing pSTAT3 IR (red) and α-MSH IR (green) and DAPI (blue) in the arcuate nucleus of a leptin-treated (5 mg/kg; 30 min) Leprb<sup>db/db</sup>;Pomc-cre;HA-Leprb flox mouse (8 weeks old). Top: Shown are single confocal planes. Examples of two POMC somata with nuclear pSTAT3 and cytoplasmic α-MSH are indicated with arrows in merged image. 3v; 3<sup>rd</sup> ventricle. Bottom: Magnification of box area. Many α-MSH IR axonal fibers with boutons are found (single arrows). These do not co-express pSTAT3. In contrast, pSTAT3 IR fibers (double arrows) do not exhibit α-MSH IR and do not show presence of axonal boutons.

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rarely seen in the cytoplasm (soma) of neurons with pSTAT3 IR nuclei. Similarly, shown in Figure 9 are different direct magnification levels of brain sections from a leptin-treated Lepr^{db/db};AgRP-ires-cre;HA-Leprb flox mouse. Many pSTAT3 IR neuronal nuclei and dendritic structures are identified. Analysis of randomly selected pSTAT3 IR structures identified neuronal nuclei (N=16), dendrites (N=358) and axons (N=1). Because these above studies were done in genetically modified and obese Lepr^{db/db} mice, we also investigated pSTAT3 IR cellular structures in the medial arcuate nucleus of leptin-treated wild type C57BL/6J mice (Figure S8). Of randomly selected pSTAT3 IR cellular structures, we counted neuronal nuclei (N=16) and dendrites (N=79), and found no evidence of pSTAT3 IR in axonal structures (N=0). We did not find pSTAT3 IR in macroglia, including in astrocytes, microglia or ependymal cells within the arcuate of these wt mice suggesting that these cell types may not express LepRb. Altogether, these EM studies show somato-dendritic localization of leptin-stimulated pSTAT3 both in random arcuate neurons of wild type mice, and in POMC and AgRP neurons of transgenic Lepr^{db/db};Pomc-cre,HA-Leprb flox and Lepr^{db/db};AgRP-ires-cre,HA-Leprb flox mice, consistent with the somato-dendritic expression pattern of LepRb shown above.

**Leptin activates STAT3 in proximity to post-synaptic structures on dendritic shafts from POMC and AgRP/NPY neurons**

To further explore the sub-cellular localization of leptin receptor signaling within dendrites, we examined possible leptin-induced STAT3 phosphorylation within dendritic spines and in proximity to post-synaptic structures on dendritic shafts of POMC and AgRP neurons. In randomly selected pSTAT3 IR (POMC) dendritic structures in brains of Lepr^{db/db};Pomc-cre,HA-Leprb flox mice we did not find direct evidence of spines by EM. This is consistent with earlier reports by Hentges et al. [59] and Liu et al. [16] showing a low number of dendritic spines on POMC neurons. By CLSM, which allows rapid analysis of much...
larger fields and longer fiber distances, we did however observe pSTAT3 IR within rare spine-like structures on POMC dendrites (Figure S9). By EM, phospho-STAT3 IR was detected in dendritic shafts in close proximity to both asymmetrical and symmetrical synapses (Figure 10). The pSTAT3 IR did not appear to concentrate directly within the post-synaptic density (PSD) itself, but was mostly associated with microtubules. Of note, pSTAT3 IR was not detected within mitochondria. Because it can be difficult to clearly differentiate the black DAB deposits from the electron-dense post-synaptic density, we employed pSTAT3 immuno-Gold EM. As shown in Figure 10, gold particles were indeed targeted to microtubules and not directly to the PSD itself. We also demonstrated presence of asymmetrical shaft synapses on POMC dendrites in brains of lean POMC;EGFP lox mice by immuno-GFP EM (Figure 10). In AgRP dendrites, we similarly detected leptin-induced pSTAT3 IR in proximity to both symmetrical and asymmetrical synapses on dendritic shafts (Figure 11). However in contrast to the reported high-density of dendritic spines on AgRP neurons relative to POMC neurons [16], we did not find evidence of spines in pSTAT3 IR dendritic structures in leptin-treated Lepr<sup>db/db</sup>;Pomc-cre;HA-Leprb flox mice.

3D-reconstruction of hypothalamic POMC and AgRP/NPY dendritic segments

To further examine LepRb-dependent STAT3 phosphorylation in POMC and AgRP dendrites, we performed serial-EM 3D-reconstruction of pSTAT3 IR dendritic segments from leptin-treated Lepr<sup>db/db</sup>Pomc-cre;HA-Leprb flox and Lepr<sup>db/db</sup>AgRP-ires-cre;HA-Leprb flox mice. Figure 12A shows a cross-section of a large pSTAT3 IR dendritic shaft in a brain section from a leptin-treated Lepr<sup>db/db</sup>Pomc-cre;HA-Leprb flox mouse. One mitochondrion is present. The dendrite was followed through 160 adjacent 50 nm thick EM sections and reconstructed in 3-dimensions using the Reconstruct software. Figure 12B shows a semi-transparent representation of the ~8.0 μm long segment with the single intracellular mitochondrion (green). This segment did not contain any spine structures consistent with the above
analyses of random pSTAT3 IR dendritic structures and the earlier report by Liu et al [16]. A different dendritic segment (11 µm long) was also reconstructed and similarly did not show evidence of spines (not shown). Figure 12B (middle) shows a non-transparent surface presentation. Finally, we identified shaft synapses on the same segment. A total of 15 synaptic structures were found along the segment and are presented in red on a semi-transparent presentation in Figure 12 (bottom).

We similarly reconstructed a pSTAT3 IR dendritic shaft-segment from a leptin-treated Lepr\textsuperscript{db/db};POMC-cre;HA-Leprb flox mouse. Figure 13A shows microphotographs of two cross sections of the pSTAT3 IR dendrite at different levels (a and b). The 3D-reconstruction of the ~12.5 µm long segment including...
its multiple mitochondria (green) and one spine-like structure is presented in Figure 13B (left). A different AgRP dendrite segment (3.2 μm long) was also examined and did not exhibit any spines (not shown). A non-transparent surface representation is shown in Figure 13B (middle). This AgRP dendritic segment had 20 identifiable synapses (Figure 13B, right).

Combined, these ultrastructural data show leptin-dependent STAT3 phosphorylation within POMC and AgRP/NPY dendrites in close proximity to both symmetrical and asymmetrical shaft synapses.

Discussion

The principal finding of these studies is that the long signaling form of the leptin receptor, LepRb, exhibits a somatodendritic expression pattern in hypothalamic arcuate POMC and AgRP/NPY neurons. In addition, we show evidence of leptin-dependent LepRb signaling in close proximity to synaptic structures on dendritic shafts. Combined, these results suggest that leptin has dendritic actions that may involve modulation of synaptic function and be important for mediating leptin’s in vivo metabolic actions.

The dendritic localization of LepRb is supported by the following findings: 1) Leptin-dependent STAT3 phosphorylation is found in relatively short neuronal fibers that do not extend significantly beyond each brain nucleus; 2) HA-LepRb is directly localized to proximal and distal dendritic fibers of POMC neurons; and 3) ultrastructural studies show STAT3 activation by leptin in dendrites in POMC and AgRP neurons. The lack of axonal LepRb expression and signaling is attributed to: 1) HA-LepRb immuno-reactivity is not found in POMC IR or β-Endorphin IR fibers; 2) leptin-activated pSTAT3 is not detected in α-MSH processes; 3) leptin-stimulated pSTAT3 is lacking in known axonal terminal beds of POMC and AgRP neurons, such as the PVH; 4) EM analyses did not show evidence of pSTAT3 in axon fibers of POMC and AgRP/NPY neurons in transgenic mice selectively expressing LepRb only in those neurons; and 5) EM analyses show negative results for pSTAT3 in axon fibers within the arcuate nucleus of wild-type animals. Despite this evidence, we cannot rule out the possibility that low levels of LepRb are present in axons and/or

Figure 11. Leptin activates STAT3 signaling in proximity to post-synaptic structures on dendritic shafts from AgRP neurons. Left Side: Immuno-EM for pSTAT3 (DAB) from a leptin-treated (4 mg/kg ip; 10 min) Lepr<sup>db/db</sup>;AgRP-cre;HA-LepRb flox mouse. The dendrite (D) is marked by black stippled line. Bottom: Magnification of symmetrical synaptic structure from stippled box above. Double arrows indicate the PSD. Right Side: Immuno-EM for pSTAT3 (DAB) in cross section of dendritic shaft from a leptin-treated (4 mg/kg ip; 10 min) Lepr<sup>db/db</sup>;AgRP-cre;HA-LepRb flox mouse. The dendrite (D) is marked by a black stippled line. Bottom: Magnification of the asymmetrical synapse from above. Single black arrow indicates the PSD. D: dendrite (pSTAT3 IR positive); D*: dendrite (pSTAT3 IR negative); M: mitochondrion; NT: presynaptic nerve terminal.

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Figure 12. 3D-reconstruction of a leptin-responsive hypothalamic POMC dendritic segment. A. Immuno-EM for pSTAT3 (DAB) in a leptin-treated Lepr<sup>db/db</sup>;Pomc-cre;HA-Leprb flox mouse. Cross-section of a pSTAT3 IR dendritic shaft with a single mitochondrion are shown. Two non-pSTAT3 IR dendrites also marked (D*). B. Serial-EM 3D-reconstruction of the POMC dendritic shaft of the pSTAT3 IR dendrite in Figure A. Top: Shown is a partially transparent presentation (mitochondrion in green and the dendritic surface in gray). Middle: Non-transparent surface representation of same POMC dendritic shaft segment (after rotation). Bottom: Shown is a partially transparent representation with shaft synapses depicted in red (N=15). Total length of segment is ~8.0 µm (reconstructed from 160 sections - each ~50 nm thick). D: dendrite; M: mitochondrion.

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Figure 13. 3D-reconstruction of a leptin-responsive hypothalamic AgRP dendritic segment. A. Immuno-EM for pSTAT3 (DAB) in a leptin-treated *Lepr<sup>db/db</sup>*;*AgRP-ires-cre;HA-LepRb flox* mouse. Left (a): Cross sections of the dendritic shaft and the spine-like structure are outlined by stippled lines and by yellow transparent color. Right (b): Shown is a cross section of same dendritic shaft, but at a different level. Two mitochondria are labeled (M). B: Left: 3D-reconstruction of the dendritic segment. The location of the two EM microphotographs from A are marked (a and b). Shown is a partially transparent representation with intracellular mitochondria in green and the dendritic surface in gray. Middle: Non-transparent surface representation. Bottom: Shown is a partially transparent representation with shaft synapses depicted in red (N=20). Total length is ~12.5 µm (250 sections – each ~50 nm thick).

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that STAT3 is activated in axons at either longer or shorter time points following in vivo leptin administration.

Somato-dendritic expression and signaling by LepRb is consistent with a recent study reporting effects of leptin on translation of brain-derived neurotrophic factor (BDNF) mRNA in dendritic fibers of VMH neurons [60]. In addition, a similar localization pattern has been reported for CNTF-activated pSTAT3 in cranial and spinal motor neurons [61]. Also, ultrastructural immuno-EM analyses in rodent brains of non-cytokine receptors such as somatostatin receptor subtypes (sst1) [62], the mu-opioid receptor (MOR) [63] and the corticotrophin-releasing factor receptor (CRFβ) have shown a predominant somatodendritic sub-cellular compartmentalization [64].

Relative to the soma, dendrites contain the vast majority of synapses on a given neuron. Our ultrastructural experiments showing leptin-dependent STAT3 phosphorylation in proximity to symmetrical and asymmetrical synapses, suggest that LepRb signaling might influence post-synaptic responses to GABA and glutamate. Indeed, electrophysiological slice studies of hippocampal neurons show that leptin can enhance the amplitude of excitatory post synaptic currents (EPSCs) via N-methyl-D-aspartate receptors (NMDAR) and α-amino-3-hydroxy-5-methyl-4-isoxazolepropionic acid receptors (AMPAR) [35,65]. Leptin did not modulate the paired pulse ratio (PPr), further indicating involvement of a postsynaptic mechanism rather than affecting glutamate release (pre-synaptic action). In addition, leptin facilitates NMDA-induced Ca2+ influx in dissociated hippocampal and cerebellar neuronal cultures and in Xenopus oocytes transfected with LepRb- and NMDAR-encoding plasmids [35,36]. Combined, these findings are consistent with dendritic expression of LepRb.

An early study of leptin’s electrophysiological effects suggested that LepRb is localized to pre-synaptic axon-terminal structures of hypothalamic NPY (AgRP) neurons serving to influence GABA release onto POMC neurons [12]. However, this conclusion was based on measurements of mini inhibitory-postsynaptic currents (miPSCs) in only 3 POMC neurons. In addition, the effect of leptin on POMC IPSC frequency was slow (~12-15 minutes) leaving a pre-synaptic somato-dendritic based mechanism feasible. For example, LepRb localized to the soma of NPY neurons might influence transfer of GABA-related substances from the soma to nerve terminals via the microtubule transport system [66-68]. Indeed, fast anterograde axonal transport can occur at rates between 140-280 µm/minute, and conveyed in this system is a variety of proteins, synaptic vesicles and enzymes associated with neurotransmitter generation [66-68]. Alternatively, with the delayed leptin time-response in mind [12], a post-synaptic mechanism involving restructing of the dendritic receptive zone (synaptogenesis) via LepRb expressed in POMC dendrites might also be possible [69-71]. Future electrophysiological studies involving mice with specific deletion of LepRb from POMC or NPY/AgRP/GABA neurons might differentiate between a pre-synaptic versus a post-synaptic mechanism for leptin to modulate PSCs in this neurocircuitry.

A more recent study reported inhibition of glutamate release by leptin onto neurons in the ventral tegmental area (VTA) and similarly to Cowley et al. [12], also showed a schematic figure depicting localization of LepRb to the nerve terminal of the presynaptic neuron [37]. This conclusion was based on measurements in brain slices showing a reduction in the probability of glutamate release onto VTA neurons. In addition, the depressive effects of leptin on AMPAR and NMDAR mini excitatory-postsynaptic currents (mEPSCs) affected frequency, but not amplitude. Such results are typically consistent with a pre-synaptic mechanism of LepRb action as proposed in the paper, and it is therefore possible that LepRb is localized differently in POMC and AgRP/NPY neurons versus other types of LepRb neurons (e.g. the glutamatergic neurons that synapse onto VTA neurons). On the other hand, the studies by Thompson et al. [37] cannot entirely exclude the possibility that the pre-synaptic actions of leptin are mediated by LepRb localized to the soma (and/or dendrites) of the glutamatergic neurons. As also discussed above, the relatively slow time-course for the effect on mEPSCs (15-18 minutes after leptin) and PPr (30 minutes after leptin) could allow enough time for leptin-action on the soma to influence transport of transmitter-related substances to nerve terminals. Indeed, we show evidence of LepRb membrane localization in the soma of arcuate POMC neurons (although these are likely not the glutamatergic neurons innervating the VTA). Future studies will be needed to decisively determine whether the above reported pre-synaptic actions on VTA neurons occur via LepRb signaling in the soma/dendrites or within axon terminals of the pre-synaptic neurons.

We also show evidence that STAT3, a transcription factor, is activated in distal dendrites. However its function in this neuronal compartment is unknown. One possibility is that STAT3 is eventually transported to the soma/nucleus to influence gene expression. Consistent with this possibility we found that pSTAT3 is associated with microtubules, the major transportation machinery in dendrites [72]. Earlier studies in non-neuronal cell models have reported interaction between STAT3 and microtubule-associated proteins, and effects of STAT3 on microtubule stabilization [73,74]. The LepRb-STAT3 pathway might thus affect dendritic growth/branching and/or neuronal migration and/or synaptic plasticity [58,70,75-77]. Alternatively, STAT3 may serve a novel function in dendrites that has yet to be determined. Of additional interest for further studies is the possibility of a dendritic localization-signal-sequence within LepRb, as has been reported in other proteins with somato-dendritic compartment localization [78-82].

Consistent with the recent studies by Liu et al. [16], we found only few spines on POMC dendrites. However, in contrast to those studies by Liu et al., we found that dendrites of AgRP/NPY neurons also have few spine structures. One possible reason for this discrepancy is that our EM studies are limited to investigations of a low number of dendritic segments that might not be representative. It is also possible that AgRP dendrites have lost spine expression due to lack of LepRb expression in neurons (except AgRP neurons) in the LepRbΔ6-60/Δ6-60; Agrp-ires-cre; HA-Leprb flox mice, or indirectly because of the obesity or other abnormalities of these animals. Alternatively,
STAT3 may be activated by leptin in regions of AgRP/NPY/GABA dendrites that have a low density of spines. Our 3D-reconstructions further indicate that the number of shaft synapses may greatly outnumber spine synapses on both POMC and AgRP/NPY dendrites. We also show evidence that many of the shaft synapses can be categorized as either symmetrical or asymmetrical, which typically represents GABAergic and glutamatergic synapses, respectively [83,84]. The GABAergic and glutamatergic synaptic input measured on POMC and AgRP/NPY/GABA neurons may therefore primarily involve dendritic shaft synapses rather than dendritic spine synapses [12,16,85].

Further implications of our finding of dendritic LepRb localization include the possibility that leptin-resistant obesity may be caused, at least in part, by altered LepRb expression or signaling in this neuronal compartment. In addition, it will be important to determine if LepRb itself is localized and signals directly within the post-synaptic density to affect synaptic activity.

Supporting Information

Figure S1. Leptin activates STAT3 phosphorylation in neuronal fiber processes within the NT5 of C57Bl/6J mice. Shown are light microscopy (LM) images of phospo-STAT3IR (DAB) in coronal brain sections of the hindbrain from 8 weeks old wild type C57Bl/6J male mice. Animals were given leptin (5 mg/kg, ip) and sacrificed after 30 minutes. Top: pSTAT3 IR is found within the NTS at the level of the area postrema (AP). The bottom image shows high magnification of stippled box from top. Arrows identify some of many pSTAT3 IR neuronal processes. XII: Hypoglossal nerve; NTS: nucleus of the solitary tract; DMX: dorsal motor nucleus of the vagus nerve; cc: central canal; AP: area postrema.

(TIF)

Figure S2. Leptin does not induce STAT3 phosphorylation within the PVH, a major axonal target of leptin-responsive POMC neurons. Left: pSTAT3 IR is present in the RCH/Arc region of the anterior hypothalamus, but importantly, not in the PVH of leptin-treated (5 mg/kg i.p., 30 minutes) C57Bl/6J mice. Bottom: High-magnification microphotograph demonstrating lack of pSTAT3 IR fibers (and nuclei) in the PVH. Right: Many POMC IR nuclei are found in the RCH/Arc. Bottom: Dense networks of neuronal (axonal) POMC fibers are observed in PVH. RCH: retrochiasmatic area; Arc: arcuate; 3v: 3rd ventricle; PVH: paraventricular hypothalamic nucleus.

(TIF)

Figure S3. Leptin activates STAT3 phosphorylation in neuronal fiber processes in hypothalamic nuclei of Sprague Dawley rats. Left: Shown are light microscopy (LM) images of phospho-STAT3 IR (DAB) in coronal brain sections of the mediobasal hypothalamus from Sprague Dawley rats. Rats were given leptin (5 mg/kg, ip) for 45 minutes. Right: High-magnification microphotographs of boxes in left column. Robust pSTAT3 IR is found in fibers within the VMH and PMN.

(TIF)

Figure S4. Expression of HA-tagged LepRb in hypothalamic POMC neurons. A. CLSM of POMC neurons (red (POMC-polypeptide IR)) and HA-LepRb (green (HA IR)) in a hypothalamic brain section from a Pomc-cre;HA-Leprb flox mouse (top row) and a negative control section from a HA-Leprb flox mouse (bottom row). Some non-specific HA IR (green) is observed along the lining of the 3rd ventricle and at the base of the Arc in the control section. Shown are single confocal planes. B. Top row: Example of a POMC soma (red) co-expressing HA-Leprb (green) in a Pomc-cre;HA-Leprb flox mouse. DAPI fluorescence (blue) identifies the nucleus. Bottom row: Example of a POMC neuron that does not express HA-Leprb in a HA-Leprb flox control mouse. Shown are single confocal planes.

(TIF)

Figure S5. Localization of leptin receptors and activation of STAT3 in neuronal fibers of transfected primary neurons. A. Primary hypothalamic neurons were co-transfected with plasmids encoding HA-tagged LepRb and GFP, and subjected to immunocytochemistry (ICC) for HA (green) and GFP (red). Leptin receptors are expressed in the soma and in fibers. Shown are collapsed confocal Z-stack sections B. As in A., neurons were transfected with plasmids encoding HA-LepRb. At DIV 12, cells were treated 100 nM leptin for 20 min and fixed. Slides were then subjected to ICC for pSTAT3 (green) and PSD95, a dendritic protein marker (red). DAPI (blue) was included to label nuclei. Top: Two neurons exhibit pSTAT3 IR in the soma and fibers. Bottom: Enlargement of box in top image showing punctate pSTAT3 staining in PSD95 positive fibers. Shown are single confocal planes. DIV: days in vitro.

(TIF)

Figure S6. Lepr<sup>db/db</sup>;Pomc-cre;HA-Leprb flox mice express pSTAT3 in neuronal processes of POMC neurons. Top Left: LM shows lack of pSTAT3 IR (DAB) in the mediobasal hypothalamus of a leptin-treated (5 mg/kg, ip, 20 min) obese Lepr<sup>db/db</sup> control mouse. Bottom Left: pSTAT3 IR in the arcuate (ARC), but not the VMH or LHA of obese Lepr<sup>db/db</sup>;Pomc-cre;HA-Leprb flox mice consistent with the targeting of HA-Leprb to POMC neurons. Insert: For comparison, the anatomical localization of POMC neurons is shown by in situ hybridization for Pomc mRNA. Right: Enlargement of box. Many positive pSTAT3 fibers are shown (arrows). 3v: 3rd ventricle; Arcuate: hypothalamic arcuate nucleus; VMH: ventromedial hypothalamic nucleus; LHA: lateral hypothalamic area.

(TIF)

Figure S7. Lepr<sup>db/db</sup>;Pomc-cre;HA-Leprb flox mice express pSTAT3 in neuronal processes of AgRP neurons. Left: LM shows pSTAT3 IR (DAB) in the mediobasal hypothalamus of a leptin-treated (5 mg/kg, ip, 20 min) obese Lepr<sup>db/db</sup>;Pomc-cre;HA-Leprb flox mouse. pSTAT3 IR is found in the medial arcuate (ARC), but not in the VMH or LHA,
consistent with targeting of HA-LepRb expression to AgRP neurons. Insert: For comparison, the anatomical localization of AgRP/NPY neurons is shown by in situ hybridization for Npy mRNA. Right: Enlargement of section. Several pSTAT3 IR fibers are visible (arrows). 3v: 3rd ventricle; VMH: ventromedial hypothalamic nucleus; LHA: lateral hypothalamic area. (TIF)

Figure S8. Activation of STAT3 phosphorylation by leptin in dendrites of arcuate hypothalamic neurons of wild type C57BL/6J mice. Immuno-EM for pSTAT3 (DAB) in the arcuate hypothalamic nucleus of a leptin-treated (4 mg/kg, ip, 15 min) wild type C57BL/6J mouse. Top: A pSTAT3 IR neuronal nucleus is labeled with "Nu" (black text) and several pSTAT3 IR dendrites are indicated with black arrows. A pSTAT3 IR negative neuronal nucleus is labeled "Nu" in white text. Insert: A cross sections of a pSTAT3 IR dendritic shaft. Bottom: pSTAT3 IR dendrite in the photographic plane. Nu; nucleus; D: dendrite; bv: blood microvessel. (TIF)

Figure S9. Activation of STAT3 phosphorylation by leptin in dendritic spines of POMC neurons. A Pomc-EGFP mouse was given leptin (5 mg/kg, ip, 30 min). IHC and CLSM was applied to visualize pSTAT3 IR (red) and EGFP-efluorescence (green) in a brain section of the arcuate nucleus of the hypothalamus. All images are single confocal planes. The arrow depicts a dendritic spine-like structure. (TIF)

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Author Contributions
Conceived and designed the experiments: CB LH SB SH. Performed the experiments: CB SH SB LH AG JH NH WL EJB ME STH. Analyzed the data: CB SH SB LH AG NH TZ. Wrote the manuscript: CB STH.


