A genetically specified connectomics approach applied to long-range feeding regulatory circuits

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Synaptic connectivity and molecular composition provide a blueprint for information processing in neural circuits. Detailed structural analysis of neural circuits requires nanometer resolution, which can be obtained with serial-section electron microscopy. However, this technique remains challenging for reconstructing molecularly defined synapses. We used a genetically encoded synaptic marker for electron microscopy (GESEM) based on intra-vesicular generation of electron-dense labeling in axonal boutons. This approach allowed the identification of synapses from Cre recombinase–expressing or GAL4-expressing neurons in the mouse and fly with excellent preservation of ultrastructure. We applied this tool to visualize long-range connectivity of AGRP and POMC neurons in the mouse, two molecularly defined hypothalamic populations that are important for feeding behavior. Combining selective ultrastructural reconstruction of neuropil with functional and viral circuit mapping, we characterized some basic features of circuit organization for axon projections of these cell types. Our findings demonstrate that GESEM labeling enables long-range connectomics with molecularly defined cell types.

Brain tissue is densely packed with complex circuits comprised of neurons and other cells, their extensive fine processes, and synapses originating from multiple cell types. These circuits are being described with increased granularity by representing circuit nodes as molecularly defined cell types, based on gene expression differences between neurons. Consequently, techniques for visualizing neural circuit structures need to be adapted for molecularly defined circuits. A number of genetic approaches have been developed to selectively label and image molecularly defined hypothalamic populations that are important for feeding behavior. Combining selective ultrastructural reconstruction of neuropil with functional and viral circuit mapping, we characterized some basic features of circuit organization for axon projections of these cell types. Our findings demonstrate that GESEM labeling enables long-range connectomics with molecularly defined cell types.

A common characteristic of these methods is that they simplify the context of neural circuit connections by selective visualization of molecularly defined neurons and their connections while ignoring complex, but unlabeled, neuropil.

In contrast, nanometer-scale anatomical analysis provides unbiased insight into the organization of neural circuit connections. Ultrastructure imaging by electron microscopy (EM) remains the gold standard for unambiguously identifying synapses and provides detailed cell biological information about the connections, such as postsynaptic density, organelle distribution and subcellular targeting. Recent studies have demonstrated the feasibility of large-scale reconstruction of mammalian and insect neural circuits using serial-section EM8–10. Large-scale ultrastructural identification of connections from molecularly defined cell types has lagged as a result of reliance on antibody detection of antigens that are uniquely present in those cells. In addition, the integrity of membranous structures is challenged during immunodetection as a result of the need for membrane permeabilization with strong detergents or, in the absence of detergent, confined to surface labeling and tissue cracks11. As a result, connectivity analysis for molecularly defined neurons is frequently limited to anecdotal examples.

In principle, these issues can be addressed by genetic labeling methods12–16. Expression of horseradish peroxidase (HRP)14,17 and the engineered singlet oxygen generator (miniSOG)15 have been used to generate electron contrast in neuronal samples. HRP is not functional when expressed in the cytosol18, but the enzymatic activity is preserved when it is targeted to the extracellular plasma membrane14,17 or endoplasmic reticulum19, and its expression does not perturb basic synaptic ultrastructure17,19. HRP and miniSOG expression are visualized through enzymatic oxidation of 3,3'-diaminobenzidine (DAB) in the presence of hydrogen peroxide (as well as light for miniSOG), which generates a polymeric precipitate. Subsequent processing with osmium tetroxide makes this DAB reaction product electron dense, which increases electron scattering in transmission EM (that is, reduces electron transmission) and appears as a corresponding dark area on the image20. However, DAB reaction product can diffuse from the site of production and obscure some ultrastructural details when targeted to the plasma membrane. To address this issue for molecularly defined cell types,
we targeted expression of HRP in the lumen of synaptic vesicles. Following exposure to DAB and hydrogen peroxide, this method labeled the interior of synaptic vesicles, which permitted the identification of axonal release sites with minimal tissue processing and excellent ultrastructural preservation.

As a proof-of-principle example of long-range ultrastructure circuit mapping and quantitative analysis of molecularly defined synapses, we investigated neural circuits that control feeding behavior. These circuits involve separate molecularly defined hypothalamic neuron types as critical circuit nodes with long-range axonal projections that are critical regulators of food seeking and consumption. Two intermingled cell types located in the hypothalamic arcuate nucleus (ARC) are defined by expression of the neuropeptides Agouti-related peptide (AGRP) and proopiomelanocortin (POMC). These cell types have opposite effects on feeding behavior and energy balance\textsuperscript{21}; increased electrical activity in AGRP neurons drives food seeking and consumption\textsuperscript{22,23}, whereas, conversely, POMC neuron activation suppresses food intake\textsuperscript{22}. AGRP and POMC neurons have an overlapping axonal projection pattern spanning multiple brain regions and their axon projections to the paraventricular hypothalamus (PVH) have received particular attention as a key circuit that regulates feeding\textsuperscript{24–26}. We investigated the ultrastructure of synapses from AGRP and POMC neurons in combination with channelrhodopsin-assisted functional circuit mapping\textsuperscript{27} and anterograde trans-synaptic viral tracing to dissect these key long-range connections. Good preservation of ultrastructure and efficient labeling enabled us to extract high-resolution reconstructed examples of release sites from these cell types across a large continuous volume. This provides an initial assessment of anatomical features indicative of circuit properties, such as the relative utilization of wired and volume neurotransmission, the use of excitatory and inhibitory neurotransmitters, and the localization of their synaptic contacts. Moreover, our findings demonstrate the practical application of GESEM-labeling techniques in two mouse brains for long-range ultrastructural circuit mapping of molecularly defined cell types.

RESULTS

A genetically encoded synaptic marker for EM

To label axonal release sites in molecularly defined neuron types, we targeted GESEM to secretory vesicles. For this, we used a chimeric protein in which HRP was fused to the C terminus of vesicle-associated membrane protein 2 (VAMP2)/synaptobrevin-2 (Syb2)\textsuperscript{27}, which localized HRP to the synaptic vesicle lumen (VAMP2::HRP; Fig. 1a). VAMP2 was chosen because of its ubiquitous presence among small clear vesicles as well as large dense-core vesicles and was therefore expected to label fast neurotransmitter and peptidergic neuromodulatory release sites, respectively\textsuperscript{28}. To achieve cell type–specific expression, we placed the VAMP2::HRP coding sequence in an inverted orientation into a Cre recombinase (Cre)-dependent recombinant adeno-associated virus (rAAV) targeting vector (rAAV2/1-\textit{CAG-FLEX-rev-VAMP2::HRP})\textsuperscript{7}. We tested the validity of this approach in mice by labeling the projections of molecularly defined AGRP and POMC neurons. We used Cre-dependent VAMP2::HRP-expressing rAAV and POMC neurons in the ARC of Agrp-Cre or Pomc-Cre mice, respectively (Fig. 1b). We focused on the long-range (>1 mm) synaptic connectivity of their axons with neurons in the PVH (Fig. 1b), which receives strong axonal input from the ARC (ARC\textsuperscript{AGRP}→PVH, ARC\textsuperscript{POMC}→PVH). Following HRP-catalyzed oxidation of DAB and subsequent processing, electron-dense staining was encapsulated in the vesicular membrane (Fig. 1c,d). This permitted unambiguous identification of the labeled boutons without obscuring other intracellular and extracellular membranes.

To further evaluate the general applicability of this cell type–specific EM labeling approach, we generated a transgenic \textit{Drosophila melanogaster} line, using the fly homolog of VAMP2 (neuronal synaptobrevin, nSyb), for broad neuronal expression of nSyb::HRP (c155-GAL4; UAS-nSyb::HRP). Similar to our results with mouse brains, we observed synapses labeled by dark, electron-dense vesicles that were similar in size to the unfilled vesicles (Fig. 1e). This technique allowed preservation of cell membranes and key ultrastructural features, such as T-bars, which indicate the sites of synaptic contact in flies. Thus, GESEM labeling by intravascular synaptic targeting with VAMP2::HRP allows \textit{in vivo} labeling of release sites from molecularly defined cell types in mouse and fly brains for ultrastructural analysis by EM.

GESEM labeling of molecularly defined synapses

We used GESEM labeling to identify axons and reconstruct synaptic connections of AGRP and POMC neurons in the PVH. After HRP-mediated DAB oxidation and subsequent processing, we prepared
serial sections from the PVH and imaged by transmission EM. For axonal reconstructions, we used two continuous volumes of neuropil (one for each cell type) that were from one Agrp-Cre mouse and one Pomc-Cre mouse. For analysis of ARC POMC → PVH axonal boutons, 11,933 PVH images were aligned, corresponding to a continuous volume of ~120,000 µm³ (~200 × 200 × 3 µm; Fig. 2a–c). Preservation of membranous structures enabled three-dimensional reconstruction of labeled axons and their boutons (Fig. 2d–g). We observed two types of vesicles that were categorized on the basis of diameter: small vesicles (SVs, 35–50 nm) and large vesicles (LVs, 80–130 nm) of varying core density (Fig. 2f). On the basis of prior immuno-EM analysis, these likely correspond to fast amino acid neurotransmitter and neuropeptide release, respectively. Most labeled boutons from POMC neurons had ultrastructural characteristics of a synaptic contact (Fig. 2h).

Similarly, for a mouse expressing VAMP2:HRP in AGRP neurons, we imaged and aligned 10,808 PVH images corresponding to a continuous volume (~120,000 µm³; ~200 × 200 × 3 µm; Fig. 2a–c).
continuous volume of ~90,000 µm³ (Fig. 3a–c). Some boutons that were reconstructed (Fig. 3d–f) showed a mixture of SVs and LVs as well as a synaptic contact (Fig. 3g), whereas others primarily contained LVs and lacked synaptic contacts (Fig. 3h). Examination of the EM stacks revealed that 32% of identified ARC AGRP → PVH boutons had a postsynaptic partner (Fig. 3i).

For GESEM labeling with VAMP2:HRP in both mouse and fly brains, not all vesicles in a bouton were labeled by the DAB reaction. AGRP axonal boutons showed higher penetration of vesicular DAB labeling (35%) than POMC terminals (13%) (Fig. 4a). In both cell types, the penetrance of labeled vesicles was higher for SVs than LVs (Fig. 4b,c). However, these differences in labeling efficacy have a
limited influence on bouton identification because only a few labeled vesicles, typically out of tens to hundreds in a bouton, are sufficient for cell type-specific synapse identification.

**Organelle profile in ARC→PVH boutons**

The organelle profile of synaptic release sites offers insight into synaptic function for these two circuits. Because GESEM labeling is confined to vesicles, the boutons were spared from obscuring deposits often seen with other HRP-labeling methods, allowing clear visualization of structural details. We selected 51 of the complete boutons from each labeled cell type and analyzed their LV and SV composition (Fig. 5a) and postsynaptic profile (Fig. 5b) and median values per bouton (Fig. 5c) overlaid with the values obtained from individual boutons (AGRP, n = 51 boutons from one mouse; POMC, n = 51 boutons from one mouse).

Other characteristics of these labeled boutons were also notable. For several boutons, including non-synaptic LV-containing boutons, we also observed the DAB reaction product on the extracellular membrane segments, which is an outcome expected from fusion of VAMP2:HRP-containing vesicles (Supplementary Fig. 3a,b). Furthermore, several membrane invaginations resembling Ω-structures were identified in non-synaptic LV-containing boutons (Supplementary Fig. 3c–h), which can be indicative of vesicle fusion.

Synaptic function is also affected by mitochondria, which influence the energy supply for synaptic transmission as well as calcium buffering during synaptic activation. Therefore, we analyzed the number and distribution of mitochondria in labeled boutons. In our imaged volumes of the PVH, AGRP boutons had fewer mitochondria than POMC boutons (Fig. 5d). The number of mitochondria was correlated with the number of SVs (r = 0.4), as would be expected from energetic requirements in the synaptic vesicle cycle (Fig. 5e), but were independent of LV pool size (r = 0.04; Fig. 5f). Consistent with this, we found that boutons with the morphological characteristics of a synaptic contact usually contained mitochondria, whereas a large fraction of non-synaptic neuromodulatory boutons did not (Fig. 5g,h). Thus, the difference in mitochondria numbers in AGRP and POMC neuron projections is likely a result of the lower number of synaptic contacts for ARC→PVH projections (Figs. 2h and 3i). Collectively, however, these results demonstrate that GESEM labeling enables systematic, quantitative and information-rich analysis of intracellular organelle profiles in boutons from long-range projections of molecularly defined cell types.

**Intra-axonal bouton variability**

Do synaptic and non-synaptic types of boutons arise from separate neuronal subpopulations? Preservation of ultrastructure in GESEM-labeled samples allowed tracing and reconstruction of axonal segments from molecularly defined cell types in the imaged volume. In our axonal reconstructions, several axon segments had multiple boutons in the imaged volume (11 from AGRP and 6 from POMC; Fig. 5i).

For AGRP axons, both synaptic and non-synaptic boutons were found in the same axons (Fig. 5i). Connected, adjacent boutons often showed marked differences in SV distribution, where SVs were segregated primarily to boutons that formed a synapse (Fig. 5i–l). LV distribution was similar across adjacent synaptic and non-synaptic boutons (Fig. 5i–l). Thus, individual ARC→PVH axons in our imaged volume contained two types of specialized boutons that were either predominantly LV-containing (peptidergic) and non-synaptic or mixed LV and SV (peptide/fast neurotransmitter) synaptic release sites. In contrast, for POMC axons, neighboring boutons had a consistent synaptic or non-synaptic ultrastructure as well as vesicle pool composition (Fig. 5i–k). Thus, these two bouton configurations may arise from separate populations; however, analysis in a larger PVH volume and additional animals is required to validate the generality of these observations. Taken together, our analysis indicate that GESEM labeling is also compatible with reconstruction of axons from long-range projections of molecularly defined neurons, which reveals substantial variability in release sites both between and within axonal segments.

**Subcellular targeting of ARC→PVH release sites**

For fast chemical neurotransmission, the subcellular location of synaptic inputs strongly affects potency to perturb neuron electrical activity. Our measurements revealed that 26% of ARC→PVH synapses were directly onto somata, but this fraction was only 2% for ARC→PVH connections (Supplementary Fig. 4a).
Additional analysis of GESEM-labeled presynaptic terminals showed that 19% of ARC\textsuperscript{POMC}\textsuperscript{+} → PVH synapses were onto spines, but no ARC\textsuperscript{AGRP}\textsuperscript{+} → PVH synapses were onto spines (Supplementary Fig. 4a). We also found that 69% of ARC\textsuperscript{POMC}\textsuperscript{+} → PVH synapses, but only 9% of ARC\textsuperscript{AGRP}\textsuperscript{+} → PVH synapses, were onto dendrites with shaft calibers of < 1 µm (Supplementary Fig. 4a). Because the diameter of PVH dendritic processes decreases with branching and distance from soma\textsuperscript{12}, this indicates that AGRP synapses onto PVH neuron dendrites may be closer to the soma than POMC synapses. Thus, in our imaged volumes, reconstructions of GESEM-labeled axonal boutons, as well as postsynaptic dendrites and somata, reveal a marked difference in subcellular targeting of these molecularly defined circuit connections, although further analysis in larger volumes is required to verify these findings.

Ultrastructural prediction of synaptic sign

The ultrastructural characteristics of synaptic contacts can also be used to predict functional synaptic properties; symmetric synapses are typically inhibitory and asymmetric synapses are excitatory and release glutamate. Because GESEM labeling permits excellent preservation of synapse ultrastructure, GESEM-assisted reconstructions of boutons and postsynaptic elements clearly revealed the synapse morphology. Using this approach, we found that AGRP neuron synapses in the PVH were mostly symmetric (symmetric, 22% of release sites; asymmetric, 3%; Fig. 6a, b). This result is consistent with previous immuno-EM analysis that showed examples of symmetric AGRP neuron synapses in the PVH\textsuperscript{36}; however, GESEM labeling and reconstruction facilitated quantitative analysis across populations of these release sites. A majority (52%) of ARC\textsuperscript{AGRP}\textsuperscript{+} → PVH boutons lacked evidence
of a synaptic contact (Fig. 6a). In addition, a subset of the boutons could not be classified, either as a result of sectioning in the plane of the synaptic contact (en face, 7%) or of damaged sections or imperfect membrane labeling (undetermined, 16%). We verified the results of EM analysis of GESEM-labeled boutons using immunohistochemical and light microscopic investigation of ARC→PVH projections. The area of the postsynaptic density (PSD) reflects the synaptic source of input to POMC neurons. We looked for unwanted inputs, and the area of the PSD was significantly less than that for neighbor- ing asymmetric synapses (37). We could not classify these synapses further as a result of sectioning in the plane of contact (en face). The area of the PSD reflects (Fig. 6a) quantitatively a bouton’s width (724 ± 152 µm², n = 11 neurons from 4 mice) to unitary current amplitudes from desynchronized evoked release (54.2 ± 5 pA, n = 12 neurons from 4 mice; Supplementary Fig. 5d). This indicates that there are ~13 ARC→PVH synapses per connected neuron, which is consistent with prior anatomical reports of ARC boutons clustered around PVH neurons. Minimal optogenetic stimulation showed lower average inhibitory postsynaptic current amplitude (164 ± 25 pA, n = 7 neurons from 5 mice; Supplementary Fig. 5e) than maximal stimulation, but inhibitory postsynaptic currents were larger than the average unitary amplitude, indicating that individual ARC axons can form multiple synapses onto PVH neurons. These functional data, taken together with ultrastructural observations of synapse subcellular localization, show that strong inhibition by ARC synaptic inputs involves multiple ARC axons per cell, which often form several synapses around PVH neuron somata and proximal dendrites.

Convergence AGRP and POMC synaptic inputs

Ultrastructure analysis of GESEM-labeled PVH tissue suggested that different subcellular postsynaptic compartments are targeted by AGRP and POMC projections. However, given that our neuronal reconstructions are based on limited volumes from two different animals, it is not possible with these data sets to distinguish whether these two cell types can target the same postsynaptic PVH neurons.

To examine the possibility of convergence of AGRP and POMC synaptic input on PVH neurons, we employed a strategy to first label ARC→PVH postsynaptic neurons and then identify juxtaposed AGRP boutons. A Cre-dependent herpes simplex virus (HSV) anterograde trans-synaptic tracer, HSV129(ΔTK)-loxP-STOP-loxP-tdTomato:2a:TK (H129ATK-TT), was injected into the ARC of Pomc-Cre mice, which resulted in Tdtomato expression in PVH neurons and, through trans-synaptic transfer, in other ARC neurons. Because AGRP neurons are a major presynaptic source of input to POMC neurons, we looked for unwanted retrograde transfer by inspecting AGRP boutons, which did not show Tdtomato expression (Supplementary Fig. 6a,c), consistent with...
prior evidence of selective anterograde trans-synaptic transfer for this viral vector. In the PVH, some POMC axon projections expressed Tdtomato, and PVH somata postsynaptic to POMC neurons were identified by Tdtomato expression (Supplementary Fig. 6c). Many of these labeled PVH somata overlapped with clusters of AGRP-immunoreactive boutons (Supplementary Fig. 6d–h). Although some of these boutons may be non-synaptic or onto adjacent dendrites, based on our EM analysis, a substantial fraction was predicted to be somatic (on average 26%; Fig. 6a). In support of this, inspection of our EM data did not show evidence of PVH somata contacted by an AGRP neuron synapse within one bouton-length of each other as an alternative somatic target. Thus, convergent synaptic connections of AGRP and POMC axon projections onto the same neurons in the PVH are indicated for the Tdtomato-labeled PVH somata with larger numbers of juxtaposed AGRP boutons (Supplementary Fig. 6i).

DISCUSSION

Neural circuit connectivity between molecularly defined cell types provides an important framework for examining brain function. Our results demonstrate that GESEM labeling of synaptic vesicles by VAMP2:HRP enables high-resolution, cell type–specific examination of ultrastructure in mouse brains and is compatible with fly brains. This technique for identifying synapses from molecularly defined neurons is an alternative to technically difficult immuno-EM approaches and it can be readily implemented to label the axon projections of Cre-expressing or GAL4-expressing cell types. This approach enabled us to perform high-quality serial-section reconstruction of genetically labeled axon projections, which was essential for quantitative analysis of cell type–specific connectivity and could not be readily achieved with prior functional and anatomical approaches. In addition, GESEM labeling facilitated visualization of cell type–specific circuit communication channels embedded in the context of other circuit interactions, as well as sub-synaptic, cell biological characteristics of these interactions. Long-range ultrastructure mapping is therefore an information-rich method that complements existing functional and viral-tracing circuit-mapping techniques.

Technical considerations for GESEM labeling

This method for cell type–specific ultrastructural circuit mapping is based on viral expression of vesicle-targeted HRP as a synaptic marker. This release site labeling strategy offers several useful characteristics. First, genetic expression of an EM contrast generator allowed reproducible labeling of synapses from molecularly defined cell types. Second, given that the marker is already present at the synapse when the tissue is prepared for imaging, there is no need for antibody labeling. Third, as a result of ubiquitous expression of VAMP2 in vesicular structures, HRP can label terminals that may not be detectable with immunolabeling against a cell type–specific marker. For example, we found that some ARC\textsuperscript{POMC}→PVH terminals had few or no LVs (Fig. 5a), which would make them undetectable by immunoreactivity to an antibody to POMC. In addition, carboxy-terminal tagging of VAMP2 has been used extensively for intravesicular protein targeting and does not interfere with vesicle fusion. Fourth, HRP is not functional in the cytoplasm; thus, previous attempts to express a soluble version have been ineffective. Genetically encoded labeling by extracellularly targeted HRP can obscure structures that are critical for the determination of connectivity. On the other hand, vesicular labeling could be readily detected and did not obscure intracellular and extracellular compartments, as the DAB oxidation precipitate was trapped in the vesicles. Finally, DAB precipitate entrapment in the compact vesicle volume also acts as a natural contrast enhancer, eliminating the extra step of tyramide signal amplification enhancement, which is required for a membrane-targeted variant because of its sparse distribution. Avoiding this extra enhancement step in our method further minimized perturbation to ultrastructure.

Not all vesicles were labeled by the DAB reaction (Figs. 1d,e and 4). Similar incomplete labeling was previously reported for VAMP2:HRP in cultured neurons. According to an estimate from artificial liposomes (60–80-nm diameter), a single HRP molecule is sufficient to fill the vesicle lumen with DAB reaction product. Thus, the lack of labeling may indicate the absence of functional VAMP2:HRP chimera from these vesicles. However, VAMP2 is one of the most abundant proteins in the vesicle membrane (~70 copies/vesicle). Because VAMP2:HRP was expressed for more than 3 weeks under a strong promoter, we expected that most vesicles would contain at least a single copy of the chimera in our samples. It is possible that vesicles already in use before viral transduction may not readily accept new VAMP2 molecules, but extensive vesicular protein intermixing has been reported in hippocampal cultures. Thus, this scenario seems unlikely, although we cannot rule out the possibility that intermixing in a distal projection might require a longer time. In addition, it is also possible that our fixation and labeling conditions might reduce the efficiency of HRP oxidation of DAB or inactivate a portion of the HRP molecules. Nevertheless, given that only a few electron-dense, DAB-labeled vesicles are sufficient for identification, even with partial vesicle-labeling penetrance, the GESEM method is effective for cell type–specific labeling. The variable efficacy of labeling resulting from viral transduction levels could be overcome by selecting only the highest efficiency viral injections for analysis. Alternatively, generation of a Cre-dependent VAMP2:HRP reporter mouse line would facilitate high-penetration GESEM labeling.

We also observed boutons with electron-dense DAB reaction product on the extracellular portion of the plasma membrane of GESEM-labeled boutons (Supplementary Fig. 3). This labeling indicates extracellularly directed HRP, which is the orientation expected of the VAMP2:HRP chimeric protein after vesicle exocytosis. Thus, extracellular labeling may be associated with recent synaptic vesicle fusion. The possibility that VAMP2:HRP could report recent synaptic activity in the context of ultrastructure circuit mapping could be developed to provide an additional level of insight into circuit function.

Synaptic organization of a feeding circuit

Energy-sensing AGRP and POMC neuron populations function in an antagonistic fashion to regulate feeding behavior. The interaction of these neurons involves axon projections to the PVH and reciprocal regulation of PVH neuron melanocortin receptors by release of the agonist, α-melanocyte stimulating hormone from POMC neurons and the corresponding inverse agonist, AGRP, from AGRP neurons. Optogenetic and chemogenetic experiments in POMC and AGRP neurons suggest that the melanocortin pathway influences food consumption over a timescale of hours to days. Recent work on the ARC→PVH circuit in behaving mice has emphasized key roles for NPY and GABA to induce food seeking and consumption within minutes through a melanocortin receptor–independent pathway. This observation suggests that there is a more complex circuit than was evident in models focused solely on regulation of melanocortin receptor signaling. Consistent with the prominent role for both neuropeptide and fast amino acid neurotransmission in these circuits, we observed that GESEM-labeled boutons showed vesicle profiles that typically had a mix of SVs and LVs, ranging from predominantly SV-containing...
synaptic connections to LV-only boutons (Fig. 5a). Overall, non-synapticboutons for both cell types had abundant LVs and often had fewer SVs than boutons with a synaptic contact. The non-synaptic ARGC and POMC neuron boutons that we imaged have features similar to non-synaptic release sites for other neuromodulators and neuropeptides30. ARCA GRP→PVH projections showed a greater proportion of non-synaptic boutons, and some of these showed electron-dense DAB reaction product on the extracellular membrane (Supplementary Fig. 3a,b). This is indicative of synaptic vesicle exocytosis and is evidence that these non-synaptic boutons are LV release sites. Furthermore, we observed several putative Ω-structures in non-synaptic boutons (Supplementary Fig. 3c–h), which can be a result of recent vesicle fusion events32. Taken together, EM analysis of GESEM-labeled non-synaptic ARCA GRP→PVH boutons indicates that some are likely release sites for their LV contents (presumably peptidergic). Notably, for ARCA GRP→PVH axons in which multiple connected boutons were reconstructed in the volume imaged by EM, many (9 of 11) showed examples of SV-rich synaptic contacts adjacent to predominantly LV-containing non-synaptic boutons. Although this observation is from a limited PVH volume from a single mouse, it indicates that the same ARCA GRP→PVH axons can contain two types of boutons that are either predominantly peptidergic or combined neurotransmitter and peptide release sites.

In these molecularly defined circuits, we also observed differential distribution of mitochondria between synaptic and non-synaptic boutons, irrespective of the cell type analyzed. In light of the multifaceted functional roles of mitochondria, these observations might indicate potential differences in the energetic regulation and calcium handling in distinct, specialized release sites that mediate non-synaptic peptidergic transmission and synaptic fast neurotransmitter/peptidergic co-transmission. One possible functional implication is that, because LV fusion is regulated by a cumulative rise in intra-terminal calcium49, this configuration may facilitate neuropeptide release at lower electrical activity levels from the subset of release sites lacking mitochondria, which are typically non-synaptic. Given the necessity of peptidergic release for AGRP neuron evoked feeding24, the presence of peptidergic release sites tuned to different activity levels would be consistent with the observation of evoked food intake even at low AGRP neuron firing frequencies22.

Our initial observations of GESEM-labeled boutons indicate that the structure of these circuits is more specialized than expected from a model focused solely on antagonistic peptidergic regulation of melanocortin receptor signaling. In PVH neurons, dendrites are targeted by predominantly excitatory POMC neuron synapses (Fig. 6a) and synapses from other cell types. As in other circuits30, distal synaptic input is likely insufficient to drive neuronal excitation and would require coincident activity from other dendritic inputs. In support of this, the identified excitatory POMC synapses had small PSD areas and synaptic current amplitudes. On the other hand, the somatocentric, multi-synapse ARCA GRP→PVH connections, in combination with the asynchronous release property of these synapses24, mediate strong PVH neuron inhibition24. Thus, these two circuits appear to target different PVH neuronal compartments and are not configured as directly opposing interactions. However, because these ultra-structural data were obtained from a single animal for each experiment (AGR or POMC), a larger sample size (and, ideally, larger volume reconstructions) will be needed to draw robust conclusions about the organization of the ARC→PVH circuitry. Furthermore, trans-synaptic viral-tracing experiments (Supplementary Fig. 6) revealed that some PVH neurons receive convergent input from both cell types. On the basis of these data, we speculate that the synaptic arrangement found here is consistent with a circuit in which distal inputs, including those from POMC neurons, are summed to influence PVH neuron activity, and this can be vetoed by long-range somatic AGRP neuron input. Consistent with this, experiments that simultaneously coactivate AGRP and POMC neurons result in voracious eating similar to that seen when activating AGRP neurons alone24, and direct suppression of PVH activity elicits intense food intake24. Thus, ultrastructural analysis shows a circuit configuration that favors feeding behavior under conditions of elevated AGRP neuron activity.

CONCLUSIONS

GESEM labeling enables long-range, high-resolution mapping of cell type–specific connections in complex molecularly defined neural circuits. Because this approach allows excellent preservation of ultrastructure, it facilitates reconstruction of molecularly defined axon segments as well as systematic and quantitative analysis of their axonal boutons and synapses. Future refinement of this method could involve combination with EM markers targeted to different subcellular compartments13–16 to enable simultaneous labeling of pre- and postsynaptic neuronal cell types. In concert with functional circuit mapping techniques, GESEM labeling could be a powerful tool for achieving cell type–specific connectomes.

METHODS

Methods and any associated references are available in the online version of the paper.

Accession codes. GenBank (KM923926).

Note: Any Supplementary Information and Source Data files are available in the online version of the paper.

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AUTHOR CONTRIBUTIONS

D.A. and S.M. Sternson designed the experiments and wrote the paper. D.A. and S.M. Sertel analyzed EM images and reconstructed synapses. J.H.S. suggested VAMP2HRP labeling and generated the UAS-N56k–HRP line. J.N.B. performed immunohistochemistry and image analysis. H.H.S. performed molecular cloning. R.D.F. developed tissue preparation and staining protocols. W.-P.L. and R.D.F. performed sectioning and transmission EM imaging. R.D.F. and L.K.S. performed electron micrograph registration and alignment.

COMPETING FINANCIAL INTERESTS

The authors declare no competing financial interests.

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ONLINE METHODS

All experimental protocols were conducted according to US National Institutes of Health guidelines for animal research and were approved by the Institutional Animal Care and Use Committee at Janelia Research Campus.

Mice. Agrp-Cre
t, Pomc-Cre
t male mice were used for GESEM labeling and ultrastructural analysis. Other mice used were Agrp-ires-Cre
t/A09 (ROSA-loxP-StoploxP-ttdTomato)
, and Pomc-topazFP
t. Background strains of mice were C57Bl6 or C57Bl6/FVB hybrid. Mice were given ad libitum access to food and were housed in a 12-12 h light-dark cycle, with lights on at 6:00 a.m. Prior to viral injection surgery, mice were housed in cages in groups ≤5. Following surgery, mice were singly housed.

Stereotactic rAAV and HSV injections. Vamp2:HRP was obtained by removing the stop codon in the Vamp-2 coding sequence (Bos taurus, amino acids 1–115), ligating a sequence encoding a linker peptide (TAGGSGTGGSGGTG) and a 2× Myc-tag polypeptide sequence (EQKLISEEDLEEQKLISEEDL), which was followed by the HRP cDNA at the 3′ end (GenBank accession code KM923926). The cDNA encoding the chimeric protein was then inserted into rAAV2-CAG-FLEX backbone in the inverted orientation to create the Cre-dependent viral expression vector (serotype 1), rAAV2-CAG-FLEX-rev-Vamp2:HRP (construct available at http://www.addgene.org/Scott_Sternson/). Viral injections were performed as described previously.

Anterior/posterior: −1.4 mm; dorsal/ventral: −5.9 mm, −5.8 mm, −5.7 mm). Viral vector (serotype 1), rAAV2-CAG-FLEX-rev-Vamp2:HRP, was produced by the University of Pennsylvania Gene Therapy Program Vector Core (titer: 3 × 1011 GC ml
). Mice were injected with the viral vector between postnatal days 21–25.

For circuit mapping experiments, rAAV2/1-CAG-FLEX-rev-ChR2:tdTomato was used as described previously. HSV infections, HSIV21-loxP-STOP-loxP-ttdTomato:Za:TK (H129ATK- TT), was obtained from the Center for Neuroanatomy with Neurotropic Viruses, strain H356). Viral infections (100 nl at each site) were made with BSL2 precautions into ARC coordinates (medial/lateral: 0.25 mm; dorsal/ventral: −5.9 mm, −5.8 mm, −5.7 mm). Viral titer was 1.3 × 109 pfu ml
. After a short incubation time (~36 h, which is associated with anterograde transfer across one synapse) in BSL2 housing, mice were perfused and processed for immunohistochemistry (see below). Tdtomato expression was Cre dependent, as injection of the H129ATK-TT viral vector into C57Bl6 mice lacking Cre-recombinase did not show Tdtomato expression.

EM. AGRP and POMC neurons in the ARC of 3-week-old Agrp-Cre and Pomc-Cre mice were transduced with a Cre-dependent virus expressing VAMP2:HRP (rAAV2/1-CAG-FLEX-rev-Vamp2:HRP). 4 weeks post-infection animals were transcardially perfused during the light period with 4% paraformaldehyde and 0.8% glutaraldehyde (wt/vol) in 0.1 M phosphate buffer. Brain tissue was vibratome sectioned (100 µm) in 0.1 M sodium cacodylate buffer and processed for EM at 20–24 °C. We injected several mice and used only the best transduced brains for EM processing. For this, we screened brain sections for viral transduction using Tyramide-FITC or tyramide-Cy3 treatment (TSA, PerkinElmer). HRP derived fluorescence signal is monitored in the PVH area of slices, and the adjacent untreated vibratome sections were selected for processing for large-scale EM imaging. The brain slices were rinsed 2 × 10 min with 50 mM glycine in 0.1 M sodium cacodylate buffer followed by 0.1 M sodium cacodylate buffer containing 0.1% saponin (wt/vol) 2 × 15 min to increase membrane permeability without compromising membrane ultrastructure. The brain slices were then incubated in 1 ml of 3,3’-diaminobenzidine tetrahydrochloride (DAB) (Sigma Chemical, cat. no. D9505-50TAP) at 0.3 mg ml
 in 0.1 M sodium cacodylate 0.1% saponin buffer for 30 min in the dark. The DAB reaction was initiated by the addition of 10 µl 0.003% H2O2 (wt/vol) and allowed to proceed for 1.5 h in the dark. The samples were rinsed with 0.1 M sodium cacodylate buffer, and then post-fixed with 1% reduced OsO4 (wt/vol) in 0.1 M sodium cacodylate for 30 min. The samples were then rinsed in distilled water, dehydrated in ethanol followed by propylene oxide and embedded in Eponate 12 resin (Ted Pella).

Brains from 7- d-old adult wt118 [iso] 5905 × [iso] Canton 5 G1 flies were isolated by dissection in cold saline and fixed for 1 h in 1% glutaraldehyde (wt/vol) in 0.1 M sodium cacodylate buffer on ice. Subsequent processing steps beginning with rinses in 0.1 M sodium cacodylate buffer containing 50 mM glycine were identical to those for HRP staining of mouse brain tissue.

Ultrathin serial 60-nm sections were cut with a Diatome diamond knife using a Leica UC6 ultramicrotome and picked up on Polyfilm films on slot grids. Sections on grids were stained with 1% uranyl acetate (wt/vol) followed by lead citrate. Sections were examined and photographed using an FEI Spirit BioTWIN TEM equipped with a Gatan U905-4k camera (3.8 nm per pixel) using the Legoion automated imaging software54, resulting in a mosaic of overlapping images for each section. Images were assembled using Fiji with fine alignment using Raveler software as described elsewhere. Briefly, alignment points, both within overlapping images and between adjacent sections, were found by cross-correlation of image patches. A least-squares fit then generated an affine transformation per image, mapping it into a global solution. Remaining discontinuities, caused by lens distortion and other nonlinearities, were minimized by defining 21 control points for each image, which were adjusted by least-squares fit to bring corresponding features into alignment, resulting in a single large flat image per section.

In each of these assembled images, DAB reaction product–labeled features were identified manually and selected for analysis. Synapses were modeled with Fiji/TrakEM2 (refs. 60,61). The first 51 complete boutons identified from each sample were used for SV, LV and mitochondria quantification (Fig. 5a–h). To check for possible artificial differences that could be caused by slight differences in region selection from different mice or sample preparation and imaging, investigators blind to sample identity compared the same anatomical parameters taken from the unlabeled parts of same images. Unlike for the labeled boutons and their targets, we did not observe significant differences in several parameters (number of SVs, Agrp-Cre control boutons: 463 ± 53, Pomc-Cre control boutons: 631 ± 75; number of LVs: Agrp-Cre control boutons: 22 ± 3, Pomc-Cre control boutons: 21 ± 3; number of synaptic mitochondria, Agrp-Cre control boutons: 1.7 ± 0.25, Pomc-Cre control boutons: 1.9 ± 0.2; dendritic diameter, Agrp-Cre control boutons: 1.6 ± 0.07 µm, Pomc-Cre control boutons: 1.7 ± 0.08 µm (n = 34 unlabeled boutons nearby labeled AGRP boutons from one Agrp-Cre mouse and n = 41 unlabeled boutons nearby labeled POMC boutons from one Pomc-Cre mouse, P > 0.05 in all comparisons between control bouton measurements from Agrp-Cre and Pomc-Cre mouse data sets)).

Axonal bouton and synapse classification. An axonal segment was classified as a bouton if it contained clustered vesicles and its diameter was >3 times neighboring axonal segments (typically 200–400 nm in width). For a bouton to be classified as having synaptic contact, we used two criteria: parallel pre- and postsynaptic membranes (extending >200 nm), and clustered SVs in contact with this membrane. Some boutons were completely reconstructed in the imaged volume, but did not have an identifiable postsynaptic partner. Another group of boutons were partially reconstructed and had a clear identifiable postsynaptic partner. For classification of boutons as symmetric and asymmetric, we examined the entire bouton for presence of a postsynaptic density (an electron dense protrusion into the postsynaptic cytosol) and focused on the sections that contained docked vesicles. In cases with two synaptic contacts from the same axon, each showed the same morphology (ARCAPR + PVP: 1 axon, both synapses symmetric; ARCPOMC→PVP: 4 axons, all synapses asymmetric; 1 axon, both synapses symmetric). A subset of synapses were classified as en face if they had clustered SVs near contact sites, but the synaptic cleft was obscured because the tissue sectioning angle was in the plane of the contact site. For some boutons, missing sections or a damaged membrane did not allow clear determination of synaptic or non-synaptic type, and they were classified as undetermined.

Electrophysiology. Experimental techniques were similar to those reported previously, and only the differences are described here. After viral infection (10–14 d incubation) mice were deeply anesthetized with isoflurane and decapitated. Coronal brain slices containing the PVH (300 µm) were prepared in chilled cutting solution containing 234 mM sucrose, 28 mM NaHCO3, 7 mM dextrose, 2.5 mM KCl, 1 mM MgCl2, 1.25 mM CaCl2, 3 mM sodium pyruvate and 1.25 mM NaH2PO4, aerated with 95% O2/5% CO2. Slices were transferred to artificial cerebrospinal fluid (aCSF) containing 119 mM NaCl, 25 mM NaHCO3, 11 mM D-glucose, 2.5 mM KCl, 1.25 mM MgCl2, 2 mM CaCl2, and 1.25 mM NaH2PO4, aerated with 95% O2/5% CO2. Slices were incubated at 34 °C for 30 min and then maintained and recorded from at 20–24 °C. Axons were identified and targeted by Tdtomato fluorescence emission. Neurons were patched using electrodes with tip resistances 4–5 MΩ. The intracellular
solution for voltage clamp recordings contained 125 mM CsCl, 5 mM NaCl, 10 mM HEPES, 0.6 mM EGTA, 4 mM Mg-ATP, 0.3 mM Na3GTP, 10 mM lidocaine N-ethyl bromide (QX-314), pH 7.35 and 290 mOsM. The holding potential for voltage-clamp recordings was −60 mV. In most recordings, internal GTP was replaced by GDP-BS (0.5 mM, Sigma).

Photostimulation. A laser (473 nm, CrystaLaser) was used to deliver photostimuli ranging from 0.01–1 mW. Neutral density filters were used to control the laser power at the specimen. Light pulse duration (1 ms) was controlled by a Pockels cell (ConOptics) and a mechanical shutter (Vincent Associates). A focal spot was targeted onto the specimen with two x-y scanning mirrors (Cambridge Technology) through a 4x or a 6x objective (Olympus). Laser power was monitored with a photodiode for each light pulse.

Evoked ARC(ΔF) → PVH quantal amplitude measurements were performed under the same conditions as above except that, in the aCSF, Ca2+ (2 mM) was replaced by Sr2+ (2 mM). Quantal events were chosen from a window immediately following stimulation until the event frequency dropped to three times above the baseline spontaneous event frequency.

For minimal stimulation, whole-cell voltage clamp was established and synaptic connectivity was tested using maximal laser intensity. Afterwards, the light intensity was reduced progressively using a neutral density filter, while monitoring synaptic release until photostimulation failed to induce synaptic release in >50% of the trials for at least 20 consecutive trials. Additional photostimulation trials were typically obtained.

Pharmacology. Drugs were bath applied with a gravity perfusion system. Blockers and final concentrations were as follows: glutamate receptors (AP-5, 50 µM; CNQX, 10 µM) and GABA_A receptors were blocked (picrotoxin, 50 µM; Sigma). In initial POMC and AGRP connectivity recordings, first measurements were performed in the absence of blockers, then sequentially AP-5 and CNQX or picrotoxin were added to confirm the nature of connection. In all other measurements with AGRP connections, saclofen (50 µM, Tocris), AP-5 and CNQX were used. Saclofen was included to prevent metabotropic GABA-mediated modulation of the postsynaptic neuron.

Antibodies. Rabbit antibody to NPY22 (1:2,000, Immunostar 22940), goat antibody to AGRP (1:5,000, Neuritox G15023)23, guinea pig antibody to RFPe2 (1:25,000, Covance), rabbit antibody to POMC (27–52)25 (1:2,000 Phoenix Pharmaceuticals, H02930), rabbit antibody to GFP (1:5,000, Invitrogen A-11122), sheep antibody to GFP (1:3,000, Abd Serotec 47451051), rabbit antibody to vGlut1 (1:2,000, SYS, 135502)26, guinea pig antibody to vGlut2 (1:2,000, SYS, 135404), rabbit antibody to vGluT2 (1:1,000, SYS, 135502)26; mouse antibody to vGat (1:100, SYS, 131011), rabbit antibody to vGat2 (1:4,000, Covance) and goat antibody to vAcht (1:2,000, Chemicon ARN100). Fluorophore-conjugated, minimalist cross reactivity secondary antibodies were from Jackson Immuno (1:500).

Immunohistochemistry and imaging. Mice were perfused with 4% paraformaldehyde in 0.1 M phosphate buffer fixative (pH 7.4). Tissue was post-fixed in this solution for 3–4 h and washed overnight in phosphate-buffered saline (PBS) (pH 7.4). Brain slices (20–200 µm thick, for imaging in the first 10 µm from the surface) were incubated overnight at 4°C with primary antibodies diluted in PBS, supplemented with 1% BSA (wt/vol) and 0.1% Triton X-100 (vol/vol). Slices were then washed three times and incubated with species appropriate and minimally cross-reactive fluorophore-conjugated secondary antibodies for 2 h at 20–24°C. Slices were rinsed in PBS (2 times) and mounted for imaging using hardset Vectashield (H-1400).

PVH boutons were imaged with a Zeiss LSM 510 using a 63x objective, NA: 1.4 to obtain confocal images with a z-section of less than 0.9 µm per image for excitation with 488- nm, 561- and 633-nm laser lines. Confocal stacks of 10–15 sections were obtained to facilitate complete reconstruction of individual AGRP or POMC boutons.

Colocalization analysis. For examination of molecular characteristics of AGRP boutons, Agrp-IRES-Cre mice were crossed with Ai9 (ROSA-IoxPSTOPloxP-topazFP) mice. POMC boutons were identified in Pomc-topazFP transgenic mice25. We performed colocalization analysis by first using fluorescence from Tdtomato or topazFP to identify AGRP and POMC boutons, respectively, without regard to the presence of other synaptic markers. We also determined that the complete bouton was captured in our image stack. This subset of boutons was then assessed for the presence of peptidergic and neurotransmitter transporters (AGRP, NPY, vGat, vGlut1, vGlut2 and vAcht).

Fluorescence image analysis of trans-synaptically labeled PVH neurons. Images were obtained from ten sections, two mice (five from each mouse) and were collected with a Zeiss LSM 510 microscope using a 63x objective, NA: 1.4. The fluorescence signal acquired for each channel was maintained in the linear range, the pinhole was constrained to less than or equal to 1 airy unit (AU), and sampling in the z axis was performed at 0.5x the optical resolution of the objective. To identify AGRP boutons on trans-synaptically labeled (Tdtomato expressing) PVH neurons, images were obtained in the PVH containing complete neuronal somata of AGRP-labeled neurons in the X, Y and Z axes. Juxtaposed AGRP boutons with Tdtomato-labeled PVH somata and proximal dendrites were identified by analyzing the images in the XY, XZ and YZ planes. Only AGRP boutons that overlapped or contacted Tdtomato labeling in all three axes were scored as contacts.

Statistics. Values are represented as mean ± s.e.m. P values for pair-wise comparisons were calculated using Microsoft Excel and SigmaPlot by Mann-Whitney U test (because conditions of equal variance were violated). For Figure 5I, we used two-tailed paired Student’s t tests, for which the normality condition was satisfied based on the Shapiro-Wilk test. No statistical methods were used to predetermine sample sizes, but our sample sizes are similar to those employed in the field37. Sample sizes of 51 boutons reflected the first manually identified 51 boutons, which were chosen as the subset to be used for detailed quantitative analysis. Sampling was semi-random as it reflected the order of identification of the labeled boutons and no specific feature of the labeled boutons. In the case of identification of boutons connected by an axonal segment, these were found by analyzing the entire sample of labeled boutons identified in the imaged volume.

A Supplementary Methods Checklist is available.

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