HypoMap—a unified single-cell gene expression atlas of the murine hypothalamus

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The hypothalamus plays a key role in coordinating fundamental body functions. Despite recent progress in single-cell technologies, a unified catalog and molecular characterization of the heterogeneous cell types and, specifically, neuronal subtypes in this brain region are still lacking. Here, we present an integrated reference atlas, ‘HypoMap,’ of the murine hypothalamus, consisting of 384,925 cells, with the ability to incorporate new additional experiments. We validate HypoMap by comparing data collected from Smart-Seq+Fluidigm C1 and bulk RNA sequencing of selected neuronal cell types with different degrees of cellular heterogeneity. Finally, via HypoMap, we identify classes of neurons expressing glucagon-like peptide-1 receptor (Glp1r) and prepronociceptin (Pnoc), and validate them using single-molecule in situ hybridization. Collectively, HypoMap provides a unified framework for the systematic functional annotation of murine hypothalamic cell types, and it can serve as an important platform to unravel the functional organization of hypothalamic neurocircuits and to identify druggable targets for treating metabolic disorders.

Hypothalamic neurocircuits are key regulators of integrative physiology and energy homeostasis1–5. In particular, the melanocortin neurocircuit, which comprises agouti-related peptide (AgRP)- and pro-opiomelanocortin (POMC)-expressing neurons in the hypothalamic arcuate nucleus (ARC), exerts effects on neurons in the hypothalamic paraventricular nucleus (PVH) and extra-hypothalamic projection sites, such as the bed nucleus of the stria terminalis (BNST)6, to control food intake and energy expenditure. Recently, studies have shown additional specialized neuronal subtypes located in the PVH and other hypothalamic regions, including the lateral (LH) and dorsomedial hypothalamus (DMH), that contribute to regulating energy homeostasis7. Single-cell RNA sequencing (sc-seq) experiments have revealed molecular heterogeneity of cell types that were previously considered homogeneous8, including POMC neurons9.

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Many sc-seq datasets exist, covering multiple brain regions and conditions. However, direct comparison of these data is challenging owing to technical and experimental variations. The integration of datasets is a key step in projects such as the Human Cell Atlas, the BRAIN Initiative, and the Cell Census Network (BICCN) (https://biccn.org/data). Recently, BICCN released an integrated single-cell reference for the primary motor cortex across different data modalities and species, underscoring the power that the analysis of systematically collected data on brain cell types and their connections can provide. The emergence of dedicated portals and applications, such as Azimuth (https://azimuth.hubmapconsortium.org/), to facilitate access to reference datasets, further highlights the usefulness of combining available resources with newly generated data.

The field of sc-seq data integration is evolving rapidly, with more than 20 available algorithms\(^{10}\). These methods use different approaches, such as shared low-dimensional embeddings (Seurat)\(^{11}\), soft-clustering strategies (Harmony)\(^{12}\), identification of nearest neighbors across datasets (Scanorama, fastMNN)\(^{13,14}\), and deep-learning strategies like variational auto-encoders (scVI)\(^{15}\). Additionally, traditional approaches developed for bulk RNA-seq, such as Combat\(^{16}\), are widely used. The two key aims of these methods are: (1) to mix datasets and correct for the technical differences originating from experimental variation, while (2) retaining the underlying biological information in each cell type.

Additionally, single-nucleus sequencing (nucSeq) has gained a lot of attention in recent years. The major advantage of nucSeq is removal of the time-consuming enzymatic cell-dissociation steps, which can potentially influence gene expression.\(^{17}\) In addition, nucSeq can be performed on frozen tissues, thus simplifying logistics, especially when dealing with precious human materials. Recent studies have shown that nucSeq is largely comparable to sc-seq, despite profiling different RNA species\(^{18,19}\).

Here, we attempted to create the first murine hypothalamic reference, HypoMap, by systematically evaluating different integration algorithms to choose the best approach for integrating data from 17 published datasets and an in-house hypothalamic nucSeq dataset from ad-libitum-fed and overnight-fasted mice. We validated HypoMap by comparing the transcriptomic profiles to: (1) sc-seq data collected from traditional Smart-Seq+Fluidigm C1 (ref. \(^{20}\); and (2) selected cell populations through bulk-bacterial artificial chromosome-translation ribosome affinity purification (bacTRAP) RNA-seq.

To further demonstrate the use of HypoMap, we molecularly and spatially characterized neurons expressing glucagon-like peptide-1 receptor (GlpTr) and prepronociceptin (Pnoc), identified from HypoMap and bacTRAP. GLP-1 is an incretin hormone secreted from the gut that has an important role in the control of food intake and satiety.\(^{20,21}\). GLP-1R agonists (GLP-1RA) are used clinically to treat type 2 diabetes and obesity, with recent studies showing that they exert their effects in the area postrema (AP)\(^{22}\); however, their effects in the hypothalamus are less known.\(^{23,24}\). Similarly, we have recently identified Pnoc neurons in the ARC (Pnoc\(^{46}\)) as a GABAergic cell population, which is readily activated upon consumption of caloric dense, highly palatable food. Activation of Pnoc\(^{46}\) neurons promotes food intake; conversely, the ablation of these neurons prevents high-fat-diet-induced hyperphagia and weight gain.\(^{25}\) Thus, Pnoc\(^{46}\) neurons represent a promising target for the treatment of obesity. Yet, a clear molecular definition of these neurons is still lacking.

**Results**

**The generation of HypoMap**

To develop a unified hypothalamic cell atlas comprising cell types from major hypothalamic regions, we combined 17 publicly available droplet-based hypothalamic sc-seq datasets\(^{20-24}\) covering different hypothalamic regions, from the preoptic area (POA) to the ventroposterior hypothalamus (VPH) (Supplementary Table 1). In addition, we performed nucSeq of 36,626 nuclei isolated from hypothalami of mice that had been either ad libitum chow fed or fasted overnight. This brought the total number of hypothalamic cells/nuclei, after quality control, to 384,925 (Supplementary Table 1).

Next, we systematically evaluated the data-integration algorithms Harmony, Scanorama, scVI, Combat and Seurat (CCA) across different parameter ranges, and benchmarked their performance on the basis of batch mixing, cell-type purity, and cluster separation (see Supplementary Information, Extended Data Fig. 1, and Supplementary Table 2 for details). We found that scVI consistently achieved the highest cell-type purity scores while retaining high cluster separation and good dataset mixing. Other methods, such as Seurat (CCA), achieved higher mixing scores, but performed worse in retaining cell-type purity (Extended Data Fig. 1b). Therefore, we proceeded with scVI and further optimized the hyperparameters on the combined dataset (see Methods and Extended Data Fig. 2) to generate the final integrated reference dataset—HypoMap, visualized here via uniform manifold approximation and projection (UMAP) in Figure 1a.

The majority of cells in HypoMap are neurons (56.9%, blue) (Fig. 1a,c), followed by astrocytes (13%, golden brown), oligodendrocytes (12.7%, orange), and oligodendrocyte precursor cells (OPCs, 9.5%, brown). HypoMap can distinguish rarer cell types, such as microglia (3.7%, light-green), endothelial cells (2.6%, light blue), tanycytes (2.5%, pink), ependymal cells (1.1%, cyan), and mural cells (0.9%, turquoise) (Fig. 1a). Figure 1b shows the expression of key neuronal markers: VGlut2 (StC17a6), VGat (StCl32a1), Th, and Hdc. We also examined the expression of regional markers, such as Sim1 for PVH, Nr5a1 (Sf1) for ventromedial hypothalamus (VMH), Tbx3 for ARC, and Rgs16 for supraoptic nucleus (SCN), which highlight the spatial origin of the neurons as one of major driving factors for segregation.

Each of the 18 datasets is distributed across multiple cell types in HypoMap (Extended Data Fig. 2d), with some areas showing over- or under-representation of cells from specific datasets. This is expected owing to the anatomically restricted sampling strategies used in some of these studies (for example Morris et al. for SCN in deep blue; Kim et al. for VMH in dark green) (Extended Data Fig. 2d). We examined the author annotations of the ARC cells from Campbell et al. for more specifically, and the dataset covers only a subset of HypoMap, as expected (Extended Data Fig. 3). Cell types identified in this study largely consist of populations from the ARC, but we also observed cell types from other regions, for example the VMH (Extended Data Fig. 3) and the pituitary, as discussed in the original study.

To construct a unified set of cell annotations, we adopted a multi-level clustering of cell populations using the Leiden algorithm\(^{42}\) and Multiresolution Reconciled Tree (mrtree)\(^{43}\) (see Methods). This resulted in a circular dendrogram (or tree) representing the underlying hierarchical organization of cell populations, similar to atlases of the brain transcriptome published previously.\(^{20,21}\) Tree pruning was achieved by merging clusters that could not be separated by marker genes. The final clusters represent an overview of the transcriptomic landscape of the ‘sequenced’ hypothalamus (Fig. 2a and Supplementary Table 3). In total, we generated seven levels of clusters, each with increasing granularity, although here we show only the top 5 levels, C2, C7, C25, C66, and C185 (Fig. 2a); the two lowest levels (C286 and C387) are hidden to retain visual clarity (Supplementary Table 4; the full tree and a split version of Fig. 2a are shown in Extended Data Figs. 4 and 5, respectively).

We next carried out differential gene expression (DEG) analysis to determine marker genes for all nodes at all cluster levels (Supplementary Tables 5 and 6). Each node of the tree was labeled using the most informative marker gene (see Methods). The full cluster annotation was constructed by concatenating the labels from all ancestor node(s), thereby incorporating the hierarchical structure (Extended Data Fig. 6 shows the marker expression of AgRP and POMC clusters
Neuronal populations

Focusing on the subset of 219,030 neurons, the next cluster levels consist of 16 (C-25) and 50 (C-66) clusters that further subdivide the GLU and GABA subtrees. An example of a well-defined cluster is the Pomp.GLU-54 neuronal cluster, which includes three subclusters: Anxa2, Pomc.GLU-5, Ttr.Pomc.GLU-5, and Glipr1.Pomc.GLU-5, consistent with Campbell et al. The lowest level depicted in the tree (C185) consists of 130 neuronal cell types (Fig. 2a). Interestingly, by combining all datasets together, we were able to identify a cluster of 61 extremely rare Gnrh1-expressing neurons (Gnrh1.GLU-9) (Fig. 2a).

The inner heatmap ring in Figure 2a depicts the contribution of each dataset. As expected, datasets that cover specific regions contribute strongly to clusters originating from that region, but little to
other clusters (for example, Wen et al.); other, less selective, datasets, such as Chen et al., and the in-house nucSeq cover a larger subset of the tree. Crucially, we found that no single dataset contributes to all clusters, thus emphasizing the power of the harmonized clustering on the basis of the integrated data of HypoMap. The middle ring in the heatmap in Figure 2a shows how each cluster contributes to the total number of cells in HypoMap (also in Supplementary Table 7).

Next, we performed spatial predictions for neuronal clusters, employing per-voxel enrichment analysis by overlapping the in situ hybridization data from the Allen Brain Atlas (https://brain-map.org) and cluster gene markers, followed by manual curation using known spatial origins of source datasets (see Methods). The predicted region annotation is shown in the outer ring (Fig. 2a and Supplementary Table 8). We found that regions with well-defined gene markers, such as the ARC, VMH, and SCN, were annotated with high confidence, consistent with annotations from the original studies. We also identified clusters originating from LH, such as Hcrt.Rfx4.GLU-4, which co-expresses Hcrt and Pydnd, and Pmch neurons (Pmch.GLU-7), consistent with Rossi et al. and Mickelsen et al. The DMH is a region that lacks distinctive gene marker(s); at C185, three clusters were predicted to originate from the DMH (two glutamatergic and one GABAergic clusters, Fig. 2a).

The distribution of ARC, VMH, and SCN neurons over multiple HypoMap clusters indicates that the larger cell numbers from additional datasets enhance the clustering granularity, thus allowing for more accurate stratification of cellular subtypes. For example, we observed a refinement of the clustering of VMH Nrs4a1- and FezF1-expressing populations (C25-3; GLU-3) in HypoMap (Fig. 3a), compared with the original annotations from Chen et al. (Fig. 3b,0); this is largely due to the integration with VMH-specific datasets, such as from Kim et al., while retaining clustering accuracy, even when compared with the original annotations from Kim et al. (Fig. 3c,0). The improved clustering sensitivity also allowed the assignment of previously unlabeled cells in Campbell et al. into more distinct clusters, such as Cck-expressing Cck.Vip2.GABA-2 cells (C185-73), which were previously annotated simply as Rgs16/Vip (Extended Data Fig. 3 inset).

Non-neuronal populations
There are 165,895 non-neuronal cells in HypoMap. Non-neuronal cells exhibit a lower level of heterogeneity, despite being sampled from different hypothalamic regions, and this is reflected in fewer branches in the tree (Fig. 2a).

The majority of non-neuronal cells originate from the oligodendrocyte lineage (42.1%, Fig. 1a) and are segregated into 15 clusters (Fig. 2a). A recent study from our group showed that oligodendrocyte differentiation in the median eminence is nutritionally regulated and plays a role in controlling energy balance. HypoMap captures different stages of oligodendrocyte differentiation, from progenitor cells marked with Pdgfra and Ng2, to differentiating cells with decreased Bmp4 and Olig2 expression, and mature oligodendrocytes with increased expression of myelination genes, such as Mbp and Mog (Supplementary Table 5).

The second largest non-neuronal subtree consists of astrocytes and ependymal cells (39.1%). Astrocytes with high levels of S1ca2 and Gjb6 were divided into nine clusters (Fig. 2a). These include a cluster of reactive astrocytes (C66-54: Lgal3.Astrocytes), marked with high expression of Gfap and Lgal3. Neighboring the astrocytes are Vim-expressing ependymal cells and tanycytes (four subclusters each), both of which form tight junctions around the third ventricle and regulate its permeability. Ependymal cells are marked with high expression of Cc13S3 and tanycytes have high expression of Col23a1, Fgfl0, and Crym. The tanycyte subclusters are consistent with previous division into alpha and beta tanycytes by Campbell et al. (Fig. 1b) and the Tany-seq atlas. We also identified a small cluster of hypendymal cells from the subcommissural organ, marked by expression of Spop. HypoMap also captures a large cluster of Ly86-expressing microglia (Fig. 2a), which could be further divided into ten clusters (Fig. 2a).

Nutritional challenges, such as a high-fat diet (HFD), are known to regulate the activity of non-neuronal cells in the hypothalamus; it would be of interest to investigate how such perturbations will affect these populations in future studies.

Single-nucleus sequencing of the mouse hypothalamus
We performed nucSeq from mice that were either subjected to an overnight fast or were ad libitum chow fed. The sequencing yielded data for 36,626 nuclei, which were integrated into HypoMap. Despite the difference in techniques, the nuclei are distributed throughout HypoMap with little evidence of technical artifact (Fig. 4a). At C185, nucSeq covers 163 out of 185 clusters (Supplementary Table 7). There is an under-representation of some cell types, particularly those originating from the POA, SCN, and PVH.

We next examined the difference in gene expression between nucSeq and all sc-seq datasets included in HypoMap. Consistent with recent findings, most genes showed a positive correlation (Fig. 4b). Notably, we found that neuropeptides/hormones, G-protein-coupled receptors (for example, Glp1r), ion channels (for example, Kcnq3), and nuclear receptors (for example, Nrs4a1) all have high average Pearson coefficients ($r$) of 0.789, 0.743, 0.736, and 0.689, respectively. Growth factors (for example, Vgfl) also showed a high correlation of $r = 0.737$. Other protein classes, such as transcriptional and translation regulators, have lower $r$ values of 0.544 and 0.358, respectively.

When we examined the gene expression correlation on a per-cluster basis (see Methods), we found that the expression profiles in nucSeq showed a high correlation with the Kim et al. dataset, whereas the correlation with earlier Drop-seq datasets, such as Campbell et al., was poorer, despite an emphasis on the same hypothalamic regions (Fig. 4c and Supplementary Table 9). The overall correlation of sc-seq and nucSeq per cluster, unsurprisingly, varies between clusters (middle ring in the heatmap in Fig. 4c). Nevertheless, these experiments highlight the overall concordance between the sc-seq and nucSeq results, and thus the feasibility to use them for unified data integration.

Infering state-dependent neuronal activation from nucSeq data
In sc-seq, the Fos signal is often unreliable owing to the short half-life of mRNA, as well as artifacts originating from the enzymatic cell-dissociation procedure, which was reflected in the absence of difference in Fos between fasted and ad-libitum-fed states in Campbell et al. (Extended Data Fig. 7a). In the nucSeq dataset, we detected an upregulation of Fos in AgRP neurons (C66-46: Agrp.GABA-4) in the fasted state (Fig. 5a), indicative of increased neuronal activity. In addition, we also found that the effect of fasting is strong enough to

Fig. 2 | Harmonized annotation of hypothalamus cell types. a, A circular hierarchical tree of clusters of HypoMap. The first 5 levels with up to 185 clusters are shown, highlighting the diversity of hypothalamic cells when combining data across regions. Individual clusters at levels 4 and 5 are named with the most informative marker gene, given as edge labels. The inner (red) circular heatmap depicts the percentage contribution of each dataset to the clusters at the lowest tree level. The middle heatmap (blue) depicts the relative percentage contribution of each cluster at the lowest tree level to the total cell number. The scale is limited to 2%. The outer ring depicts the most likely region of origin (R) for each neuron cluster on the lowest level of the displayed tree. If support was insufficient for a cluster, no region was assigned, and the cluster was colored gray (see Methods). b,c, Dot plots displaying marker genes used for annotating the clusters at level 4 (C66) of the tree in a. For clusters with a proper name (for example, ‘Astrocytes’), the most specific gene that would have been used for annotation is included. Dot color corresponds to average log-normalized expression levels of each gene in a cluster and dot size to the percentage of cells expressing a marker in the cluster. b, Neuronal cell types. c, Non-neuronal cell types. See also Supplementary Tables 5 and 6.
influence the clustering of AgRP neurons (Fig. 5a). Thus, in the following analysis, we used the higher-level classification C66-46: Agrp.GABA-4 for AgRP neurons and C286 for all other clusters. We further examined other immediate early genes (IEGs) in AgRP neurons and found that they also exhibited comparable fasting-induced expression (Fig. 5b). We therefore aggregated the response of all IEGs (see Methods) and identified 15 additional neuronal clusters that showed increased neuronal activity in the fasting state (Fig. 5c and Supplementary Table 10). The effect was strongest in C66: Agrp.GABA-4, with an upregulation of ~90% of expressed IEGs. This is followed by two other ARC clusters: C286-149: Grp.Ppp1r17.GABA-1 and C286-139: Myo5b.Sox14.Lef1.GABA-1 (Fig. 5c). Here, we conducted the differential analysis on C286 with
Fig. 3 | Comparison of HypoMap and original clusters. **a**, HypoMap UMAP highlighting the cluster C25-3: GLU–3, which contains NrSat1– and Fezf1–expressing neuronal populations from the VMH that are compared in **b**–**e**. **b**–**e**, UMAP plot of cells from the C25-3: GLU–3 cluster from Chen et al.**34** (b) and Kim et al.**32** (c) overlaid on all cells of the cluster (gray) and colored by the original author annotations. **d**, Sankey diagrams showing the original author annotations of Chen et al.**34** and Kim et al.**32** (c) vs. the HypoMap cell labels). **e**, VMH-specific dataset from Kim et al.**32** (c) partitioned. (See Supplementary Table 20 for a full overview of all original and HypoMap cell labels).

high granularity, because we observed that changes were restricted to specific subclusters, while other daughter nodes under C185-88: Sox14. Lefl.GABA-1 showed no difference (Fig. 5c). Lefl has previously been shown to be crucial for energy homeostasis**15,16**, and the identification of Lefl–expressing subclusters pinpoints the specific cellular subtypes and their molecular characteristics for future studies.

Next, we examined DEGs in the fasted state (Extended Data Fig. 7c and Supplementary Table 11). The top cluster was C66: Agrp.GABA-4 neurons, with 797 DEGs. This was followed by C286-45: Rai14.Hmcn2. Gpr149.GLU–3, with 757 DEGs. There were 172 DEGs detected in C286-149: Gpr149.GLU–3, with 757 DEGs. Top DEGs in AgRP neurons and other activated cell types were significantly regulated in at least one cluster, and 33 and 155 of the overlapping DEGs between at least 20% of all clusters revealed pathways other than AgRP neurons, suggesting that it has a more global role. We also examined all DEGs across all clusters: 1,738 genes were significantly regulated in at least one cluster, and 33 and 155 of the DEGs were found to be significantly regulated in at least 50% or 20% of all clusters, respectively. The Gene Ontology enrichment analysis of overlapping DEGs between at least 20% of all clusters revealed pathways including protein translation and cell death (Extended Data Fig. 7d).

Evaluation with Smart-Seq2 Fluidigm C1 and bacTRAP datasets

To further evaluate the utility of HypoMap, we projected external datasets onto the reference map from (1) a hypothalamic sc-seq dataset by Romanov et al.**20** and (2) bulk bacTRAP RNA-seq of specific neuronal populations across different levels of cellular heterogeneity (Figs. 6 and 7 and Supplementary Tables 13–18).
Fig. 4 | Comparison of nucSeq and single-cell data. a. UMAP visualization of the nucSeq data colored and annotated by cluster level 3 (C25) on all HypoMap cells (gray), demonstrating that the nucSeq data are evenly integrated in HypoMap. b. Heatmap of per-gene correlation (Pearson's $r$) between sc-seq and nucSeq. Each row shows the density (color) of all genes in a specific gene class (number of genes shown on the right). Also see Supplementary Table 9. c. Heatmap of cluster-level correlation shown on the hierarchical tree of neuron clusters. For each cluster, the marker genes (M, number depicted in inner heatmap in red) were used to calculate Pearson's $r$ between all sc-seq and nucSeq data (middle heatmap in blue–green) or between individual HypoMap datasets and nucSeq (outer heatmap in blue–red). If there were fewer than ten cells per cluster and dataset, the comparison was omitted (white).
Fig. 6 | Projection of new data. a, HypoMap UMAP colored by cluster level 3 (C25) and overlaid with the projected ‘locations’ of cells from Romanov et al.20. Even clusters represented by only few cells in the query dataset can be accurately embedded into the reference. B, Probability scores (see Methods) of projection accuracy of Romanov et al.20 cells from C66-46: Agrp. GABA-4. Right, UMAP plots of the same cells showing Fox expression in fasted and ad-libitum-fed conditions. The changes in nucSeq AgRP neurons after fasting were strong enough to cause a shift in the cluster. b, IEGs with high log2(fold change) (log2 FC) in AgRP neurons. The violin plots show the per-cell expression between conditions. c, Neuron clusters activated by fasting. The bar plot depicts the percentage of significantly up-regulated IEGs in the fasted state over the total number of expressed IEGs (left number, based on presence in at least 10% cells of clusters in either condition). AgRP neurons are strongly activated, as indicated by the high number of changing IEGs. The bars are colored by mean log2 FC, and the number of cells in each cluster is shown on the right. d, Transcriptional changes in AgRP neurons induced by fasting. In the volcano plot (log2 FC versus adjusted P values from a (two-sided) Wald test), differentially expressed genes (DEGs) are highlighted. The dot plot shows Gene Ontology (GO) terms enriched in up-regulated genes. The P values are based on a hypergeometric test from an over-representation analysis and were corrected using false discovery rate (FDR). e, Comparison of transcriptional changes in AgRP neurons between nucSeq and Campbell et al.1 data. The scatter plot of log2 FCs is color by DEGs in either dataset. f, Per-cell expression levels of selected DEGs between conditions. For each gene, the expression is shown across multiple activated cell types as well as POMC neurons and a reference containing all remaining cells. P values of DEGs were obtained by Wilcoxon rank-sum tests and were adjusted for multiple comparisons using Bonferroni correction. See also Supplementary Tables 10 and 11.

Romanov et al.20 reported one of the earliest sc-seq datasets generated for the hypothalamus using traditional Smart-Seq2+Fluidigm C1 (ref. 20). We successfully projected cells from this dataset onto HypoMap (Fig. 6a and Supplementary Table 12). At C185, cells were assigned to 125 of 185 clusters, although only 41 clusters contained 10 or more mapped cells. Most cell type labels were projected with high confidence (Fig. 6b). Consistent with the original study, the majority of cells were oligodendrocytes (41% of all cells). Among neuronal clusters, Shox2-expressing neurons (C185-1: Shox2.Glu-1) (5%) were the most common assignment, containing 163 cells, which were mostly left unclassified before (Fig. 6a).

Further inspection of the original annotations demonstrated that the inferred cluster labels are an accurate reflection of the true identity of these cells. For example, 12 of 23 cells annotated as ‘GABA 13 (Galanin)’ were projected onto Gal- and Th-expressing C185-91: Gal.Hmcrn1. GABA-1 cluster of HypoMap, and others onto closely related clusters and other Gal-expressing clusters. Furthermore, HypoMap enhanced the stratification of the original annotations: out of 27 cells annotated as ‘GABA 7 (Pomc +/-)’, only 7 are Pomc-positive and 5 of these cells were mapped to the Pomc.Glu-5 cluster, whereas all Pomc-negative cells were relocated to other HypoMap clusters.

bacTRAP RNA-seq is a useful tool to obtain marker-gene-specific molecular expression profiles with a great sequencing depth and the possibility to compare between different conditions21. We generated bacTRAP of specific hypothalamic cell types previously shown to be involved in energy homeostasis, that vary in their heterogeneity and anatomical distribution. To this end, we first crossed mice expressing Cre-recombinase in either AgRP, POMC, Pnoc, or Glp1r neurons with mice allowing for Cre-dependent expression of ribosomal protein L10a fused with green fluorescent protein. We then mapped these bacTRAP datasets onto HypoMap to examine the concordance of gene expression and determine whether HypoMap could be used to deconstruct the cellular composition in bacTRAP.

We first focused on bacTRAP obtained from AgRP and POMC neurons, which are relatively homogeneous cell types, and we further explored more heterogeneous data obtained from Pnoc and Glp1r neurons. We selected genes that were significantly enriched in the immunoprecipitation (IP) samples compared to the control samples of the bacTRAP datasets to define cell type signatures (see Supplementary Tables 13–18 for all DEGs). Subsequently, we compared bacTRAP marker genes and the signature fold changes using rank biased overlap (RBO), which allows the comparison of ranked sets with more weighting for the top part of the list that contains the most relevant markers23. In Figure 6c, we highlight the AgRP bacTRAP signature enrichment as small bars on top of the UMAP plot, showing the expression of Agrp in the sc-seq of HypoMap. This demonstrates that the AgRP signature is mapped successfully onto the cluster containing AgRP neurons in HypoMap (C66-46: Agrp.GABA-4), and indicates that the sc-seq integration grouped all AgRP neurons into one branch of the tree, and therefore could be used for further interpretation (Fig. 6c and Supplementary Table 19). There were other enriched ARC clusters with lower enrichment scores, indicating that they share some of the marker genes with AgRP neurons.

Next, we analyzed the POMC bacTRAP data (Fig. 6d): the C286-75: Anxa2.Pomc.Glu-5 cluster at level C286 (score = 0.256) was more enriched than the other POMC-expressing clusters, C286-77: Trt.Pomc.Glu-5 (score = 0.197) and C286-76: Glp1r.Pomc.Glu-5 (score = 0.185), an effect that is at least partly driven by the higher abundance of canonical marker genes for POMC neurons, such as Cartpt (Supplementary Table 5). In addition, we observed low-grade mapping of the POMC bacTRAP data onto Aggrp-expressing clusters (Fig. 6d), and the enrichment score for all other clusters was lower.

We recently found that leptin receptor (Lepr) and Glp1r-expressing POMC neurons showed little overlap and have distinct molecular signatures and functions6. This study, we profiled these two POMC subpopulations using intersectional Cre/Dre-dependent targeting. This approach builds on the use of POMC CRE as well as LeprCre and Glp1rCre transgenic mice. In contrast to the POMC Cre transgenic model, POMC CRE expression exhibits a progressive increase in recombination later in adulthood, thereby circumventing the developmental marking of AgRP neurons6. Analysis of the intersectional POMC bacTRAP gene signatures revealed that the POMc-Lepr signature overlaps mostly with C286-75: Anxa2.Pomc.Glu-5 (score = 0.413), but scores lower in other POMC-expressing clusters (Fig. 6e), as previously validated functionally. This is consistent with the expression of Pomc and Lepr depicted in the UMAP of Figure 6e (only cells expressing both genes are highlighted). The enrichment of the POMc-Glp1r signature was highest in C286-77: Trt.Pomc.Glu-5 (score = 0.325), which is concordant with the expression of Glp1r and Pomc in HypoMap (Fig. 6f). The high
enrichment score in C286-75: Anxa2.Pomc.GLU-5 (score = 0.299) was likely driven by the high expression of marker genes, such as Pomc and Cartpe, similar to POMC-only bacTRAP.

Validation of predicted Glp1r-expressing neurons

Next, we turned to the more heterogeneous Glp1r- and Pnoc-expressing neurons. Again, we utilized the bacTRAP signature for Glp1r-expressing neurons.
hypothalamic cells to investigate their heterogeneity in HypoMap. At C286, the highest enrichment of Glp1r bacTRAP was found in the Agrp- and Oxt-expressing neurons C286-81: Ebf3.Caprln2.GLU-6 (score = 0.19) and C286-82: Oxt.Caprln2.GLU-6 (score = 0.166), both from the PVH, consistent with previous findings\(^\text{24}\) (Fig. 7a). Interestingly, Glp1r-expressing POMC neurons (C286-77: Ttr.Pomc.GLU-5)\(^\text{24}\) were enriched, with a lower score of 0.19. We identified four additional populations on the basis of high Glp1r expression and enrichment score: C286-175: Ghrh.GABA-3, C286-174: Trh.Nxk2.4.GABA-3, C286-181: Il1rapl2.0tp.Sst.GABA-4, and C286-130: Tbx19.Il1rapl2.GABA-1 (Fig. 7a and Supplementary Table 19).

We performed multiplexed single-molecule in situ hybridization to identify the spatial distribution of Glp1r-expressing populations (Fig. 7). Using probes specific to Glp1r, Pomc, and Anxa2 for the Ttr, Pomc and Anxa2.Pomc subclusters, we found that although 49% of all POMC-ARC neurons expressed Glp1r, only 14.6% co-expressed Pomc and Anxa2 (Fig. 7b,c). For the SST\(^\text{24}\)-positive neuronal, we used probes against Sst and Unc13c to target the C286-181: Il1rapl2.0tp.Sst.GABA-4 cluster and found that 64.5% of Sst.Unc13c-positive cells expressed Glp1r (Fig. 7b,c). When we used Sst as the only marker gene, the percentage of Glp1r-positive cells decreased to 31.4%, indicating that C286-181: Il1rapl2.0tp.Sst.GABA-4 is indeed the most relevant SST\(^\text{24}\)-subtype (Fig. 7b,c).

Ghrh is a distinct marker of C286-175: Ghrh.GABA-3 neurons, which control growth via the growth hormone (GH)–insulin-like growth factor 1 (IGF1) axis\(^\text{25}\). We found that 47.9% of Ghrh-expressing cells expressed Glp1r (Fig. 7b,c). However, GLP-1 action has not been investigated in these neurons, and these findings offer the possibility that they may serve as an integrator in the adaptation of metabolism and growth.

Anxa2 is expressed in C286-130: Tbx19.Illrapl2.GABA-1. Using probes specific for Anxa2 and Tbx19, we found that 38.1% of Anxa2- and Tbx19-positive cells expressed Glp1r (Fig. 7b,c), which was markedly higher than the number of triple-positive cells identified in sc-seq (Extended Data Fig. 8). Trh and Nxk2.4 were used to distinguish C286-174: Trh.Nxk2.4.GABA-3 from other Trh-expressing cells. Although, we detected only a few Trh-positive cells in ARC, and not all of them were Nxk2.4-positive, we found a striking overlap with Glp1r expression: 90.9% of Trh- and Nxk2.4-expressing cells were Glp1r-positive (Fig. 7c). Last, we examined the expression of Glp1r in Oxt-expressing cells in the PVH, which had the strongest bacTRAP enrichment; 47.9% of the Oxt-expressing cells were Glp1r-positive (Fig. 7c), consistent with the previous findings\(^\text{26}\).

**Validation of predicted Pnoc-expressing neurons**

Pnoc is widely expressed in the hypothalamus, thus the molecular signature obtained through bacTRAP reflects the heterogeneity of this diverse population (Fig. 7d). Projecting Pnoc bacTRAP onto HypoMap resulted in lower enrichment score overall than that of the AgRP and POMC bacTRAP, but resulted in more enriched clusters (Fig. 7d). Many of these clusters express Pnoc at high levels in HypoMap, indicating that Pnoc bacTRAP is identifying these cell types correctly. At tree level expressing cell types in HypoMap. e, f, RNAscope of Pnoc and marker genes of selected ARC neuronal cell types based on Pnoc-bacTRAP and gene expression in (d). e, f, Representative images (e) and quantification (f) of Pnoc in Sst or Crabp1 co-expressing subclusters. Points refer to individual sections, in total 14 ARC sections along the rostral-caudal axis from 4 mice were included for each experiment. Mean ± s.e.m: Sst/Pnoc: 59.9 ± 3.24; Sst/Pnoc/Nts: 6.78 ± 3.56; Sst/Pnoc/Unc13c: 52.28 ± 7.17; Sst/Pnoc/Nts/Unc13c: 2.29 ± 1.1; Crabp1/Pnoc: 76.35 ± 2.2; Crabp1/Pnoc/Tmen215: 41.02 ± 4.81; Crabp1/Pnoc/Htr3b: 32.18 ± 3.04; Crabp1/Pnoc/Tmen215/Htr3b: 10.02 ± 1.17. In all dot plots, the red point depicts the mean and red error bars the s.e.m. of all sections. We used a two-sided Wilcoxon rank-sum test (multiple testing correction with Benjamin–Hochberg) to test for differences between the means of relevant groups and added the resulting P values to the quantification in (e) and (f).
reference. Harmony, Scanorama, and Combat also performed well, when tuned properly, especially when using a large number of latent variables\textsuperscript{12,14,16}. Overall, while computational metrics can assist in determination of an optimal integration method, these should be combined with careful evaluation and experimental validation.

We showed that nucSeq covers many of the clusters in HypoMap (163 of 185). Comparing the transcriptomic profiles of sc-seq and nucSeq highlighted a dataset-related bias: it correlates better with some datasets (for example, Kim et al.\textsuperscript{32}), which could be related to the single-cell technology used (that is, Drop-seq versus 10x). However, because some 10x-based datasets (for example, Moffit et al.\textsuperscript{30} and Mickelsen et al. (Flynn10x)\textsuperscript{27}) show a lower correlation, this may not be the only factor. Differences also exist between cell types; for example, the Trh-expressing cluster C185-12: Ebf1.Trh.GLU-2 showed a markedly lower $r$ value of 0.017 (across all datasets) than that of its sister-cluster C185-11: Cbln2.Trh.GLU-2 ($r = 0.789$) (Fig. 4c). This divergence could be related to differences in RNA species measured between sc-seq and nucSeq.

Changes in IEGs can be used to infer neuronal activity\textsuperscript{17,32}. Here, AgRP neurons are known to be activated during fasting\textsuperscript{57} and are used as positive controls. No other cell type showed an effect similar in strength compared to AgRP neurons, but some, such as C286-149:

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**Figure Legends**

1. Logo-normalized expression of Pomc, Anxa2, Glp1r, and Merge.
2. Logo-normalized expression of Sst, Unc13c, Glp1r, and Merge.
3. Logo-normalized expression of Nkx2-4, Trh, Glp1r, and Merge.
4. Logo-normalized expression of Oxt, Glp1r, and Merge.
5. Logo-normalized expression of Ghre, Glp1r, and Merge.
6. Logo-normalized expression of Tbx19, Anxa2, Glp1r, and Merge.
Grp.Ppp1r17.GABA-1 and C286-139: Myo5b.Sox14.Left.GABA-1, could be interesting targets for future studies. When comparing cells from fasted and ad-libitum-fed mice, we detected DEGs in many cell types. Interestingly, many of the DEGs are common across multiple clusters. This indicates that these changes may not be cluster specific, but might follow a common program (for example, protein translation) in response to a change of the metabolic state.

To demonstrate the utility of HypoMap, we showed that the underlying scVI model is able to project new hypothalamic sc-seq data, such as the Smart-Seq+ Fluidigm C1 data by Romanov et al. onto the existing reference, thereby allowing quick and reliable annotation of cells in new datasets.

We also attempted to project bacTRAP bulk RNA-seq data of known neuronal populations onto HypoMap, to demonstrate that HypoMap was able to segregate these neurons over a wide range of cellular complexities and heterogeneity. Importantly, the POMC transgenic-based translational profiling not only faithfully captured bona fide POMC-expressing cells, but also detected the AgRP neuron cluster, which expressed the POMC transgene developmentally and had thus been lineage-traced in this approach, as previously described. However, HypoMap was also able to differentiate distinct POMC neuron subpopulations, as revealed by selective ribosomal profiling of Lepr+ and Glp1r-expressing POMC neurons via intersectional recombinase based targeting. Our recent study using this system allowed us to unravel the microcircuits and functional consequences of these heterogeneous cell groups. Mapping of the more complex Pnrc and Glp1r bacTRAP datasets further illustrates how HypoMap can be successfully used to disentangle complex cell type mixtures. Compared with AgRP and POMC neurons, Pnrc and Glp1r neurons are less well characterized populations, and HypoMap offers an opportunity to better define these diverse, physiologically relevant neuron subtypes.

Here, we validated six Glp1r-expressing cell populations in the ARC and other hypothalamic regions. GLP-IRa can activate POMC neurons and contributes to an acute anorexigenic effect. Consistent with this, acute chemogenetic activation of Glp1r-expressing POMC neurons rapidly and potently suppressed food intake, but less so in Lepr-expressing POMC neurons. Activation of Glp1r-expressing cells in the PVH has also been described to mediate an acute anorexigenic effect, and HypoMap can identify these clusters: C286-82: Oxt.Caprin2.GLU-6 and C286-81: Ebf3.Capr1.GLU-6, which were both validated by in situ hybridization. Integrating this information with the identification of additional Glp1r-expressing clusters in HypoMap may guide functional studies to further delineate the cells responsible for mediating the weight-reducing actions of GLP-1RAs. Candidate Glp1r-expressing clusters include Trh and Nkx2.4 co-expressing populations and Grh-positive and Tbx19-positive populations. The enrichment observed in C286-75: Anxa2.Pomc has also been described to mediate an acute anorexigenic effect, and HypoMap can identify this cluster, which expressed the POMC Cre transgene developmentally and had thus been lineage-traced in this approach, as previously described. However, HypoMap was also able to differentiate distinct POMC neuron subpopulations, as revealed by selective ribosomal profiling of Lepr- and Glp1r-expressing POMC neurons via intersectional recombinase based targeting. Our recent study using this system allowed us to unravel the microcircuits and functional consequences of these heterogeneous cell groups. Mapping of the more complex Pnrc and Glp1r bacTRAP datasets further illustrates how HypoMap can be successfully used to disentangle complex cell type mixtures. Compared with AgRP and POMC neurons, Pnrc and Glp1r neurons are less well characterized populations, and HypoMap offers an opportunity to better define these diverse, physiologically relevant neuron subtypes.

For each Drop-seq library, the sequence alignment of the biological reads was performed using STAR 2.7.5 to the mouse genome GRCh38 (mm10); and (2) perform the UMI and gene count sequence reads from 17 publicly available datasets were downloaded using 'fastq-dump' from the SRA toolkit (version 3.0.0), with the exception of (1) GSE132355, GSE133576, GSE167927, and SRP135960, for which the original 10X BAM files were downloaded and used to regenerate fastq files via the 'bamfastq' command from the Cellranger software (version 6.0.1, 10X Genomics); and (2) Kim et al., for which the original reads could not be found and the count table was used instead.

## Methods

### Dataset download

An overview of all datasets included in the reference map is available in Supplementary Table 1. Sequence reads from 17 publicly available datasets were downloaded using ‘fastq-dump’ from the SRA toolkit (version 3.0.0), with the exception of (1) GSE132355, GSE133576, GSE167927, and SRP135960, for which the original 10X BAM files were downloaded and used to regenerate fastq files via the ‘bamfastq’ command from the Cellranger software (version 6.0.1, 10X Genomics); and (2) Kim et al., for which the original reads could not be found and the count table was used instead.

### Sequence alignment, UMI and gene count

For 10X datasets (including nucSeq), Cellranger Version 6.0.1 (5.0.1 for nucSeq) was used to (1) map sequence reads to the mouse genome GRCh38 (mm10); and (2) perform the UMI and gene-level counts against Ensembl gene model V100, with Gm28040 removed to recover Kiss1 expression. The per-cell gene count tables (in HDF5) generated by the software were then used for downstream analyses.

For each Drop-seq library, the sequence alignment of the biological reads was performed using STAR 2.7.5 to the mouse genome GRCh38 to generate a BAM file containing mapped reads. Read 1, which contains the cell barcode (CBC)/UMI read was first split into two separate fastq files, one containing just the CBC and the other containing the UMI, using ‘fastx_trimmer’ from the FASTX Toolkit (version 0.0.13), and were then tagged (XC and XM, respectively) onto the mapped biological reads using Egbio’s ‘AnnotateBamWithUmis’ command (Fulcrum Genomics, version 1.5.1). All the untagged reads were removed from annotated BAM file using Samtools (version 1.14), and PCR duplicates based on UMI (XM tag) were removed using Picard’s ‘MarkDuplicates’ command (version 2.22.2).

The deduplicated BAM file was then further annotated with genes using the gene model above (GRCh38, Ensembl V100 minus Gm28040) using ‘TagReadsWithGeneFunction’ from Drop-seq tools (version 2.3), and digital gene expression was performed using ‘DigitalExpression’ command from Drop-seq tools, using a minimum number of transcript cut-off of 800 UMIs to generate a gene-level expression matrix for all cells detected, in a tab-delimited text format.
Dataset quality control
We used the expression matrices of each dataset and ran a basic preprocessing using Seurat (version 4.0.2)\. Each dataset was further filtered to contain cells with a minimum of 1,000 UMI per cell and a maximum fraction of mitochondrial genes of 0.1. We curated a common set of metadata features, including unique cell, sample and dataset IDs, sample description (sex, mouse strain, experimental condition), dataset descriptions (technology, regional identities), and cell details (author annotations, UMI counts, cellcycle scores), among others. We harmonized the annotations of major cell types (‘classes’) to be comparable across datasets, including: neurons, tanycytes, astrocytes, oligodendrocytes, ependymocytes, immune cells, vascular/endothelial cells, and fibroblasts. For the data from Kim et al.\(^3\), no raw read files were available, hence we used the UMI count matrices provided by the authors and subset the cells to those annotated by the authors as hypothalamic cells (removing doublets, low-quality cells, and other unannotated cells). After identifying within-dataset batches, we merged all datasets to one combined dataset as input for our integration pipeline.

Batch identification
We considered datasets as independent datasets if they were split into multiple datasets by the original authors (for example Wen et al.\(^1\)). For all individual datasets, an automated detection of batches was used: starting with a default processing using Seurat, consisting of log-normalization, feature detection (2,000 features), and principal component analysis (PCA), a low-dimensional embedding with 50 PCs was obtained. Afterwards, randomForest R package (version 4.6) (https://cran.r-project.org/package=randomForest) was used to predict the sample ID using the PCs as features and cells as observations. Using hierarchical clustering with spearman correlations \(\rho\) of the pairwise out-of-bag (oob) probabilities as distances, a clustering of samples was obtained. We iteratively merged the closest samples in the associated dendrogram and trained another random forest only on the currently merged samples. The entropy of the oob class probabilities, normalized by the logarithm of the total number of samples and averaged over all cells, was used to estimate how well samples are separated. An entropy close to 1 means high similarity of samples; decreasing entropy indicates that the random forest can tell the merged samples apart and that there could be a batch effect. Merging of samples was stopped when the averaged entropy over all (merged) samples fell below 0.9. We calculated the entropy \((H)\) as:

\[
H(X) = -\sum_{i=1}^{c} p(X_i) \times \log(p(X_i))
\]

where \(p(X_i)\) is the vector of class probabilities.

Doublet removal
We used DoubletFinder (https://github.com/chris-mcginnis-ucsf/DoubletFinder)\(^4\) to identify doublets within each batch (as defined above), independent of all other data sets. We used 70 PCs and a fixed value for the variable numbers of artificial doublets (pN) of 0.25. For the neighborhood size (pK), we iterated over multiple values to find an optimal one for each batch but limited it to a maximum of 0.1. For 10x sc-seq data, we set the expected rate of doublets to 0.05; for Drop-seq data and pre-filtered data (for example, Kim et al.\(^3\)) we set the expected rate of doublets to 0.01. Using the DoubletFinder predictions and data and pre-filtered data (for example, Kim et al.\(^3\)) we set the expected rate of doublets to 0.05; for Drop-seq data, we set the expected rate of doublets to 0.01. Using the DoubletFinder predictions and pre-specified genes such as ‘mt-’ or ‘Rp’ and any genes that were already selected Leiden clustering results with increasing granularity, aiming to roughly triple the number of included clusters with each level. In total, 7 clustering levels yielding between 2 and 680 clusters were generated. These clusters were subsequently combined into a clustering tree using the Multiresolution Reconciled Tree (mmtree)\(^5\) algorithm. We used the mmtree function from the original package (https://github.com/pengminshi/MRtree) with standard parameters. We did not cut the resulting tree at an optimal level, but instead used it as a representation of the complex subtypes within neuronal cell types. After determining marker genes distinguishing each cluster node from its siblings in the tree (as described below), we merged sibling clusters with less than 10 relevant markers (specificity > 1, see ‘Cluster annotation’) into one cluster node to avoid over-clustering.

For visualization of the tree we used the ggtree R package (version 2.4.2) (https://github.com/YuLab-SMU/ggtree). VMH neuron cluster comparisons were visualized using the sankeyNetwork function from the networkD3 R package (version 0.4) (https://cran.r-project.org/package=networkD3).

Marker gene detection
We used the van Elteren test implementation for Seurat objects (https://github.com/KChen-lab/stratified-tests-for-seurat) to detect marker genes for each node of the cluster tree, comparing the current cluster either against all other nodes on the same level or against the sibling nodes to obtain a set of markers per cluster. We subset the tree into its neurons and non-neurons and calculated the global marker genes only versus other cells in the respective subtrees to restrict the marker sets to more relevant genes.

Cluster annotation
We calculated a specificity score, \(S\), per gene and cluster, \(c\), as

\[
S_{c, ref} = \frac{\text{Average log}_2\text{Foldchange}_{c, ref} \times \frac{p_{c, ref}}{p_{c, all}}}.
\]

which includes the foldchange of the average expression of cluster \(c\) and a set of reference cells (for example, all others), and the percentage (Pct) of expressing cells in cluster \(c\) and the reference. We used the specificity \(S\) to rank genes, and additionally the adjusted \(P\) values from the van Elteren test to reduce this ranking to potential marker genes. To determine the most relevant marker genes and to select a cluster name, each node (cluster) of the tree was visited from top to bottom. We ranked the most descriptive markers by calculating a score (7) for each cluster \(c\) as

\[
T_c = S_{c, all} \times S_{c, siblings} \times \frac{1}{\max (S_{c, siblings})}.
\]

which includes the specificity compared with all other clusters, as well as only the sibling clusters and the specificity for all children clusters \(i\). Additionally, we excluded pre-specified genes such as ‘mt-’ or ‘Rp’ and any genes that were already used as a name in the ancestor nodes. To incorporate a node’s parent...
cluster, the final cluster annotation was constructed by concatenating the node’s best marker gene with its parent’s name. If no siblings existed, no additional gene name was added. For the first three levels of the tree, we manually set the names of clusters on the basis of the previously curated major cell types ('class') and, in some cases, specific neuronal markers.

Region annotation
In order to predict potential spatial locations of clusters, we combined the likely regions of origin based on the original dataset (if specific to certain subregions), with a prediction based on ISH expression values from the Allen Brain Atlas62 (https://mouse.brain-map.org/). We used the cocoframer R package (version 0.1.1) (https://github.com/AllenInstitute/cocoframer) to query the API and obtain ISH expression values for each voxel from the coronal sections.

For each voxel we used the ‘energy’ value to rank all probes. Then, for each neuronal cluster from level 6 of the HypoMap tree (C286), we selected all probes with matching marker genes (specificity > 4) and calculated the median rank in that voxel as an enrichment score. We aggregated the median ranks (normalized to total number of expressed probes) of the four voxels with the highest median ranks. We subtracted the aggregated normalized median ranks from 1 and used the result as the final score for each region, so that a score close to 1 indicates an enrichment of the cluster marker genes in the top ranked probes of the voxels of each region. In Supplementary Table 8, we have reported the top ten results for each cluster.

To build one final regional prediction, we assigned the highest scoring region as the possible spatial origin (if the score was >0.8). However, to further refine the prediction, we included the known dataset origin for many studies. For each subregion-specific dataset, we manually curated a set of likely Allen Brain Atlas regions and checked their contribution to the cells of each cluster. On the basis of this, we down-scaled the median rank predictions if insufficient cells originated from the predicted region (for example, a high mammillary region score was down-scaled if the majority of cells originated from preoptic/anterior datasets). We further checked many clusters manually and changed the final annotation shown in the paper figures to the second- or third-highest prediction for well-documented cell types. To reduce the number of categories for the color-coded annotation in Figure 2a, we summarized some regions by spatial proximity (for example, grouping all mammillary and pre-mammillary clusters or all preoptic region clusters together).

Mapping of new data
We stored the reference scVI model using scvi.model.SCVI.save. The scvi.model.SCVI.load_query_data method allows the embedding of a query dataset into the original UMAP 9. The nearest neighbors via scArches algorithm15,63. We used Seurat’s ProjectUMAP function (version 1.26.0) to calculate RBO scores for each bacTRAP signature on each cluster of the mrtree clustering64 (https://www.bioconductor.org/packages/release/bioc/html/gespeR.html).

NucSeq comparative analysis
To compare the molecular profiles of single-cell and single-nucleus data in HypoMap, we calculated the correlation of gene expression between our in-house nucSeq and the public sc-seq datasets. For the comparison, we used the projected level 5 clustering (C185, corresponding to the lowest level of the hierarchical tree in Figs. 2a and 4c). We selected genes that were expressed in more than 20% of cells of a cluster or that had a mean expression > 0.2 (log-normalized) in at least one cluster. Using the mean expression per cluster (single-cell versus nucSeq), we calculated Pearson’s r for each gene and grouped genes by different classes on the basis of the Ingenuity pathway analysis (Qiagen) for further analysis. For neuropeptides we included genes annotated with the gene ontology (GO) term GO:0035179 or annotated ko04080 in the KEGG pathway. In order to compare the correlation of sc-seq and nucSeq on the cluster level, we used marker genes of each cluster with specificity > 0.25 and Pearson’s r > 0.3. Pearson’s r for each cluster was calculated between nucSeq and HypoMap or the individual sc-seq datasets in HypoMap across all marker genes.

We used a curated set of IEGs from Wu et al.17 and reduced the set to 73 genes (expressed in >300 cells and a maximum of 10,000 cells in nucSeq). We manually added 1700016P03Rik, which correlated with Fox. To determine whether 1700016P03Rik is a CREB1-target gene, we looked for cAMP response element (CRE) in the promoter66. Briefly, we used BSgenome (version 1.58) (https://bioconductor.org/packages/release/bioc/html/BSgenome.html) to retrieve promoter sequences –800 to +200 base pairs (bp) around the transcription start site (TSS) (mm10). We then used refTSS v3.3 to obtain a set of TSSs per gene66. Detection of JASPAR profiles MA0018.1 and MA0018.2 from JASPAR2020 (ref. 67) was conducted with TFBSTools (version 1.28) (https://bioconductor.org/packages/release/bioc/html/TFBSTools.html) using the function searchSeq on both strands and with a relative score of >80%. We counted the number of detected profiles per promoter and normalized it by the number of non-overlapping promoters per gene. We found multiple occurrences of CRE in 1700016P03Rik’s promoter, similar to other known CREB1-target genes (Extended Data Fig. 7b). To quantify the activation of clusters (Level 4) we counted the number of significantly up-regulated IEGs detected using the Wilcoxon test implemented in Seurat’s FindMarkers per cluster and split by diet condition, using Bonferroni correction for multiple testing.

Differential expression between cells from fasted and ad-libitum-fed mice was performed using the Wilcoxon test and negative binomial generalized linear model implemented in Seurat’s FindMarkers function (see Supplementary Tables 10 and 11). The enrichment of significantly up-regulated genes in the fasted state in AgRP neurons for GO ‘Biological Process’ terms was conducted with the clusterProfiler R package (version 3.18.1)49 (https://bioconductor.org/packages/release/bioc/html/clusterProfiler.html). We used the enrichGO function with default parameters and all genes expressed in at least 10% off all AgRP neurons in nucSeq as background gene set. The simplify function was used to remove redundant terms...
Experimental methods
All experimental methods regarding animal care and mouse lines, details on the bacTRAP, nucSeqm and in situ hybridization experiments can be found in the Supplementary Information.

Statistics
Statistical analyses are described in the respective sections. No prior power calculations were performed. For large-scale hypothesis testing (for example, marker genes or differential gene expression) dedicated R packages for these analyses were used and included correction for multiple testing (DESeq2: Benjamini–Hochberg, Seurat: Bonferroni). For individual null hypothesis tests, we used the appropriate functions in R: t.test, wilcox.test, or cor.test with alternative set to ‘two.sided.’ For the quantification in Figure 7, we additionally used functions from the R packages ggpubr (version 0.4.0) (https://github.com/kassambara/ggpubr) and rstatix (version 0.7.0) (https://github.com/kassambara/rstatix).

Reporting summary
Further information on research design is available in the Nature Research Reporting Summary linked to this article.

Data availability
Both HypoMap and the hypothalamic nucSeq are available in an interactive CellxGene viewer (available via https://www.mrl.ims.cam.ac.uk). Additionally, the Seurat object containing the HypoMap, which is required to reproduce the shown figures and to project new data, is deposited at University of Cambridge’s Apollo Repository (https://doi.org/10.17863/CAM.87955) in standard RDS format.

The nucSeq and the bacTRAP profiling data for Agr2, Glp1r, and Pomc neurons are available from the Gene Expression Omnibus (GEO), accession numbers: GSE207736 and GSE208355, respectively. The Pnoc bacTRAP data are available at GSE137626. The Pomc-Lepar and Pomc-Glp1r bacTRAP data are available at GSE153753. The published sc-seq studies used to construct HypoMap are listed in Supplementary Table 1. Source data are provided with this paper.

Code availability
The code used to create all Figures, all additional input data, the output plots, tables and source data files can be found at: https://github.com/istueurnagel/HypoMap_paper. An R package that allows mapping of new single-cell data onto the existing HypoMap scVI model is available at: https://github.com/istueurnagel/mapscvi. The pipeline for evaluation of scVI hyperparameters to optimize the HypoMap model can be found at https://github.com/istueurnagel/scIntegration. The pipeline to build HypoMap, including all downstream clustering and annotation steps, can be found at: https://github.com/istueurnagel/scHarmonization. The R packages scUtils (https://github.com/istueurnagel/scUtils) and scCoco (https://github.com/istueurnagel/scCoco) provide additional functions used in the above pipelines. We will provide any other code upon request.

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Author contributions
L. S., B. Y. H. L., P. K., G. S. H. Y., and J. C. B conceived the present study. L. S. and B. Y. H. L. conducted the computational analysis and code development. B. Y. H. L., P. K., P. D., and A. B. supervised the computational analysis. C. A. B., A. J. d. S., A. d. R. M., and W. C. performed the bacTRAP experiments. B. Y. H. L., G. K. C. D., J. A. T., and S. N. K. conducted the single-nucleus sequencing and performed the bioinformatic analysis. I. C., D. R., and A. P. C. lead the mouse work in Cambridge. C. A. B. performed the ISH experiment for validation of the Glp1r-expressing clusters via RNAscope analysis. T. S. H. performed the ISH experiment for validation of the Pnoc-expressing clusters via RNAscope analysis. G. K. C. D. and B. Y. H. L. deployed the CellxGene server for data sharing. L. S., B. Y. H. L., P. K., H. F., G. S. H. Y., and J. C. B. wrote the manuscript with support of all co-authors. All authors agreed on the final version of the manuscript.

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Competing interests
P. D. worked for Novo Nordisk A/S. G. S. H. Y. receives grant funding from Novo Nordisk A/S, and consults for them on their obesity ‘break-out’ campaign. J. C. B., I. S., C. A. B. and H. F. received project funding from Novo Nordisk A/S. The other authors declare no competing interests.

Additional information
Extended data is available for this paper at https://doi.org/10.1038/s42255-022-00657-y.
Extended Data Fig. 1 | Selection of single-cell integration method. a, Overview of the pipeline used to determine an optimal integration of data including normalization and feature selection. b, Evaluation metrics calculated on integration results of preliminary dataset (85,000 cells). Purity refers to a combined score of cell type purity and cluster separation. Mixing refers to a combined score of dataset mixing. An optimal integration achieves high mixing scores while retaining high purity scores. c, UMAPs of each integration method’s best result colored by dataset of origin to show dataset mixing and cluster separation. As indicated by the metrics, most methods are able to mix the data and retain most of the original cell types. The Raw PCA is not mixing the data fully, while the low-dimensional Harmony result is not able to represent the full complexity of the data.
Extended Data Fig. 2 | See next page for caption.
Extended Data Fig. 2 | Optimization of scVI parameters. a, Evaluation metrics calculated on scVI integration results of full HypoMap (384,925 cells). Purity refers to cell type purity only. Cell type separation (asw_norm = average silhouette width) is shown by the point size (see methods for details on metrics). PCA (orange) clearly mixes that data less well than scVI (pink). b, Evaluation metrics similar to (a), calculated on scVI integration results with comparable hyperparameters using either all cells (light blue) or only neurons (grey) as input. Using all cells as input did not affect the integration performance in mixing and purity, but the the cluster separation (asw_norm) was lower. c, Example box plots for detailed evaluation of scVI hyperparameters, visualizing the influence of the number of training rounds (epochs) and hidden layers on the three different metrics. Each point corresponds to a scVI training run on the full HypoMap data. The center of the boxplot is the median of all runs, the lower and upper hinges correspond to the first and third quartiles and the whiskers extend from each hinge to the largest value smaller than 1.5 times the distance between the first and third quartiles. Overall n = 224 scVI runs that were compared, the number differs between boxplots depending on the parameters. d, UMAP visualization of HypoMap colored by datasets to visualize mixing. e, UMAP visualization of HypoMap colored by mapped cell types for the evaluation of purity (see Methods and Supplementary Table 2 for details).
Extended Data Fig. 3 | UMAP visualization of cells and original annotations from Campbell et al. UMAP visualization of cells and original annotations from Campbell et al., highlighting which parts of HypoMap are covered by this dataset. Inset shows the enlarged view of cell clusters from the SCN colored by HypoMap clusters.
Extended Data Fig. 4 | Full hierarchical cluster tree. Full hierarchical cluster tree of HypoMap showing all 7 cluster levels. This includes level 6 (C286) and 7 (C465). Individual clusters at levels 4 – 7 are named with the most informative marker gene, given as edge labels. Full cluster names were constructed by concatenating the given gene names with those of all ancestors.
Extended Data Fig. 5 | Hierarchical cluster trees of HypoMap split into neuronal (A) and non-neuronal (B) populations. Similar to Fig. 2A, but split into neuronal (a) and non-neuronal (b) populations. The first 5 levels with up to 185 clusters are shown. Individual clusters at levels 4 and 5 are named with the most informative marker gene, given as edge labels. The inner (red) circular heatmap depicts the contribution of each dataset to the clusters at the lowest tree level in percent. The middle heatmap (blue) depicts the relative contribution of each cluster at the lowest tree level to the total cell number in percent. The scale is limited to 2%. The outer ring depicts the most likely region of origin (R) for each neuron cluster on the lowest level of the displayed tree. If support was insufficient for a cluster, no region was assigned and the cluster was colored in grey (see methods). For the non-neuronal cell types in (b) no regional prediction was conducted.
Extended Data Fig. 6 | Marker gene expression across datasets. a-b. Violin plots showing the expression of the top 5 marker genes (selected by specificity and adjusted p-value) of POMC subcluster C185-48: Anxa2.Pomc.Glu-5 (a) and AgRP subcluster C185-115: Npy.Agrp.GABA-4 across datasets, demonstrating that the expression level of key marker genes are mostly stable.
Extended Data Fig. 7 | Neuronal changes after fasting. a, there is no change in Fos expression in Campbell et al. AgRP neurons after fasting. Left: UMAP plot depicting C66-46: Agrp.GABA-4 in HypoMap. The inset shows Agrp expression in Campbell et al. cells from C66-46: Agrp.GABA-4.Agrp.GABA-4. Right: UMAP plots of the same cells showing Fos expression in fasted and ad libitum fed states. b, CREB1 transcription factor binding site (TFBS) detection in IEG promoters compared to 1000 randomly sampled background gene promoters. 1700016P03Rik contains multiple putative CREB1 binding sites in its promoter sequence, indicating it could play a role as an immediate early gene. c, Number of differentially expressed genes (DEG) after fasting in each cluster shown on the UMAP of nucSeq data. d, Enrichment of DEGs that were found in at least 20% of all clusters (155 genes) in ‘biological process’ gene ontology (GO) terms. The values are based on a hypergeometric test from an over representation analysis and corrected using false discovery rate (fdr). Globally regulated genes are enriched in translation related terms and ‘cell death’.
Extended Data Fig. 8 | Gene expression of marker genes used for Glp1r RNAscope validation. For each marker combination that was quantified together with Glp1r (see also Fig. 7a–c), double or triple positive cells are shown on UMAP subsets around the relevant cell types. The color scale is based on the square root of the product of expression levels. The colored rectangles in the grey reference in the central UMAP show which parts of the complete HypoMap are shown in the subsets. For most cell types the gene combinations are highly specific, especially compared to their neighboring cells. However, for Tbx19/Anxa2 only very few Glp1r cells exist in the single cell data.
Extended Data Fig. 9 | Gene expression of marker genes used for Pnoc RNAscope validation. For each marker combination that was quantified together with Pnoc (see also Fig. 7e-f), double or triple positive cells are shown on UMAP subsets around the relevant cell types (See Extended Data Fig. 8 for details).
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Software and code

Policy information about availability of computer code

Data collection

Image acquisition software: Leica ASX v. 3.5.5, 19975

Data analysis

We have extensively described our computational pipeline in the methods. Please also refer to the github repositories described in the code availability statement:

The code used to create all figures, all additional input data and the output plots and tables shown here can be found at https://github.com/steuernagel/hypoMap_paper. An R package that allows quick mapping of new single cell data onto the existing HypoMap scvi model is available at https://github.com/steuernagel/mapscevi. We will share these repositories and any other code upon request, and change the repositories to public access upon publication.

This docker image combines most packages required for working with our scvi and R neural pipeline: https://hub.docker.com/r/steuernagel/r_scvi

Additionally we provide a list of the most relevant software and packages used in this study:

R (version 4.1.0)
Seurat R package (version 4.1.0)
python (version 3.8)
scvi (version 1.5.3)
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Both HypoMap and the hypothalamic nucSeq is made available in an interactive cellxgene viewer (available via https://www.mrl.ims.medschl.cam.ac.uk). Additionally, the Seurat object containing the HypoMap, which is required to reproduce the shown figures and to project new data, is deposited at University of Cambridge’s Apollo Repository (doi:10.17863/CAM.87955) in standard RDS format.

The nucSeq and the bacTRAP profiling data of Agrp, Glp1r, and Pomp neurons are available from the Gene Expression Omnibus (GEO), accession numbers: GSE207736, and GSE208355 respectively. The Proc bacTRAP data are available from GSE137626. The Pomc-Lepr and Pomc-Glp1r bacTRAP data are available from GSE153753. The published sc-seq studies used to construct HypoMap are listed in Suppl. Table 1).

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Sample size

No prior power calculations for sample size were performed. Sample sizes (n=3-5) for bacTRAP, nucSeq and RNAscope experiments were based on previous experiments (Biglari, 2021; Jais, 2020). For RNAscope validation we aimed for 4 mice with 2-4 sections (rostral and caudal) analyzed per animal and celltype.

Data exclusions

We removed single cells from the nucSeq that were of low quality (e.g. Doublets or low UMI numbers). For the creation of HypoMap removed cells that were of low quality and additionally removed samples that had fewer than 100 cells left after cell filtering. We did not remove any other data.

Replication

A substantial part of the analysis conducted in this study revolves around replicability and comparability of the different data types. We generally find that the nucSeq data is well comparable with the single-cell data but dependent on covariates such as single-cell technology. The bacTRAP RNaseq data partly recapitulates the findings from the single-cell data but does not exactly replicate the expected cell types.
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| ×   | ChIP-seq              |
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**Antibodies**

- Antibodies used: Heintz Lab TRAP anti-GFP 19F7 antibody (Cat# Htz-GFP-19F7, RRID:AB_2716736)
- Antibodies used: Heintz Lab TRAP anti-GFP 19C8 antibody (Cat# Htz-GFP-19C8, RRID:AB_2716737)

**Validation**

From: Heintz Lab; Rockefeller University. Validated in Heiman et al., 2008, DOI:https://doi.org/10.1016/j.cell.2008.10.028
Ordered from: Memorial Sloan-Kettering Cancer Center

**Animals and other organisms**

Policy information about studies involving animals, ARRIVE guidelines recommended for reporting animal research

**Laboratory animals**

- Cologne (bacTRAP & RNAscope):
  - Mice were housed in individually ventilated cages at 22–24°C and humidity at 45-55% using a 12-h light/dark cycle. Animals had access to water and food ad libitum and were fed a normal chow diet (ssniff, V1554). Food was only withdrawn during defined fasting periods. C57BL/6N mice were obtained from Charles River, France. For RNAscope experiments 16 h fast, 10 weeks old male C57BL/6N mice were sacrificed. For bacTRAP experiments 10 weeks old male Glp1rCre ROSA26lSIEGFPL10a mice, 12 weeks POMCCre ROSA26lSIEGFPL10a and 12 weeks AGRPcre ROSA26lSIEGFPL10a were sacrificed in a random-fed state.
  - Details on genetically modified mouse lines:
    - Driver lines: Glp1r-ires-Cre (Williams, 2016), AgRP-ires-Cre (Balthasar, 2004) and POMC-Cre (Anastassiadis, 2009) mice have been previously described.
    - ROSA26lSIEGFPL10a (ROSA26-CAGS-lox-STOP-lox-EGFPL10a-WPRE): This line was generated by breeding ROSA26lSfrERGFPL10a (ROSA26-CAGS-lox-STOP-lox-EGFPL10a-WPRE) 7 with a ubiquitously expressed CAGGS-Dredi er line (Anastassiadis, 2009).
    - Experimental lines: Glp1rCre ROSA26lSIEGFPL10a mice were generated via mating homozygous Glp1r-ires-Cre mice to homozygous ROSA26fl/fl mice of the EGFPL10a construct. A similar breeding strategy was used for the POMCCre ROSA26lSIEGFPL10a mice.
      - Resulting double transgenic Cre+/- ROSA26fl/wt mice were used as experimental animals.

- Cambridge (single nucleus sequencing):
  - Male C57BL/6j mice at 6-8 weeks were housed in ventilated cages in a controlled temperature (20-24°C) and humidity (45-65%) facilities with a 12-h light/dark cycle (lights on 06:00–18:00) and had ad libitum access to food (RM3(E) Expanded chow, Special Diets Services, UK) and water in the animal facility at the Anne McLaren Building, University of Cambridge. For the overnight fasted group (6 animals), the chow was removed from at 5pm until 9am the next day, the animals had access to water throughout the fasted period.

**Wild animals**

- The study did not involve wild animals.

**Field-collected samples**

- The study did not involve samples collected from field.
Ethics oversight

All animal procedures were conducted in compliance with protocols approved by local government authorities (Bezirksregierung Köln). Permission for breeding and experiments on mice was issued by the Department for Environment and Consumer Protection-Veterinary Section in Cologne.

Mouse studies performed in Cambridge were in accordance with UK Home Office Legislation regulated under the Animals [Scientific Procedures] Act 1986 Amendment, Regulations 2012, following ethical review by the University of Cambridge Animal Welfare and Ethical Review Body (AWERB).

Note that full information on the approval of the study protocol must also be provided in the manuscript.