Title: A multimodal cell census and atlas of the mammalian primary motor cortex

Authors: BRAIN Initiative Cell Census Network (BICCN)

ABSTRACT

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We report the generation of a multimodal cell census and atlas of the mammalian primary motor cortex (MOp or M1) as the initial product of the BRAIN Initiative Cell Census Network (BICCN). This was achieved by coordinated large-scale analyses of single-cell transcriptomes. chromatin accessibility, DNA methylomes, spatially resolved single-cell transcriptomes, morphological and electrophysiological properties, and cellular resolution input-output mapping, integrated through cross-modal computational analysis. Together, our results advance the collective knowledge and understanding of brain cell type organization: First, our study reveals a unified molecular genetic landscape of cortical cell types that congruently integrates their transcriptome, open chromatin and DNA methylation maps. Second, cross-species analysis achieves a unified taxonomy of transcriptomic types and their hierarchical organization that are conserved from mouse to marmoset and human. Third, cross-modal analysis provides compelling evidence for the epigenomic, transcriptomic, and gene regulatory basis of neuronal phenotypes such as their physiological and anatomical properties, demonstrating the biological validity and genomic underpinning of neuron types and subtypes. Fourth, in situ single-cell transcriptomics provides a spatially-resolved cell type atlas of the motor cortex. Fifth, integrated transcriptomic, epigenomic and anatomical analyses reveal the correspondence between neural circuits and transcriptomic cell types. We further present an extensive genetic toolset for targeting and fate mapping glutamatergic projection neuron types toward linking their developmental trajectory to their circuit function. Together, our results establish a unified and mechanistic framework of neuronal cell type organization that integrates multi-layered molecular genetic and spatial information with multi-faceted phenotypic properties.

INTRODUCTION

Unique among body organs, the human brain is a vast network of information processing units, comprising billions of neurons interconnected through trillions of synapses. Across the brain, diverse neuronal and non-neuronal cells display a wide range of molecular, anatomical, and physiological properties that together shape the network dynamics and computations underlying mental activities and behavior. A remarkable feature of brain networks is their self-assembly through the developmental process, which leverages genomic information shaped by evolution to build a set of stereotyped network scaffolds largely identical among individuals of the same species; life experiences then sculpt neural circuits customized to each individual. An essential

step toward understanding the architecture, development, function and neuropsychiatric diseases of the brain is to discover and map its constituent neuronal elements together with the many other cell types that comprise the full organ system.

The notion of "neuron types", cells with similar properties as the basic units of brain circuits, has been an important concept since the discovery of stereotyped neuronal morphology over a century ago ^{1,2}. However, a rigorous and quantitative definition of neuron types has remained surprisingly elusive ^{3–7}. Neurons are remarkably complex and heterogeneous, both locally and in their long-range axonal projections that can span the entire brain and connect to many target regions. Many conventional technologies analyze one neuron at a time, and often study only one or two cellular phenotypes in an incomplete way (*e.g.* missing axonal arbors in distant targets). As a result, despite major advances in past decades, until recently phenotypic analyses of neuron types remained severely limited in resolution, robustness, comprehensiveness, and throughput. Besides technical challenges, complexities in the relationship among different cellular phenotypes (multi-modal correspondence) have fueled long-standing debates on how neuron types should be defined ⁸. These debates reflect the lack of a biological framework of cell type organization for understanding brain architecture and function.

In the past decade, single-cell genomics technologies have rapidly swept across many areas of biology including neuroscience, promising to catalyze a transformation from phenotypic description and classification to a mechanistic and explanatory molecular genetic framework for the cellular basis of brain organization ^{9–12}. These technologies provide unprecedented resolution and throughput to measure the molecular profiles of individual cells, including the complete sets of actively transcribed genes (the transcriptome) and genome-wide epigenetic landscape (the epigenome). Application of single cell RNA-sequencing (scRNA-seq) to the neocortex, hippocampus, hypothalamus and other brain regions has revealed a complex but tractable hierarchical organization of transcriptomic cell types that are consistent overall with knowledge from decades of anatomical, physiological and developmental studies but with an unmatched level of granularity ^{13–19}. Similarly, single-cell DNA methylation and chromatin accessibility studies have begun to reveal cell type-specific genome-wide epigenetic landscapes and gene regulatory networks in the brain ^{20–25}. Importantly, the scalability and high information content of these methods allow comprehensive quantitative analysis and classification of cell types, both neuronal and non-neuronal, revealing the molecular basis of cellular phenotypes and properties. Further, these methods are readily applicable to brain tissues across species including humans, providing a quantitative means for comparative analysis that has revealed compelling conservation of cellular architecture as well as specialization of cell types across mammalian species.

Other recent technological advances have crossed key thresholds to provide the resolution and throughput to tackle brain complexity as well, for example for whole-brain neuronal morphology

and comprehensive projection mapping ^{26,27}. Furthermore, powerful new methods, including imaging-based single-cell transcriptomics, the combination of single-cell transcriptome imaging and functional imaging, and the integration of electrophysiological recording and single-cell sequencing, allow mapping of the spatial organization, function, and electrophysiological, morphological and connectional properties of molecularly defined cell types ^{28–32}. Finally, the molecular classification of cell types allows the generation of models for genetic access to specific cell types using transgenic mice and, more recently, short enhancer sequences ^{33–39}. All of these methods have been applied to brain tissues in independent studies, but not yet in a coordinated fashion to establish how different modalities correspond with one another, and how explanatory a molecular genetic framework is for other functionally important cellular phenotypes.

Recognizing the unprecedented opportunity to tackle brain complexity brought by these technological advances, the overarching goal of the BRAIN Initiative Cell Census Network (BICCN) is to generate an open-access reference brain cell atlas that integrates molecular, spatial, morphological, connectional, and functional data for describing cell types in mouse, human, and non-human primate brains ⁴⁰. A key concept is the Brain Cell Census, similar conceptually to a population census, which accounts for the population of constituent neuronal and non-neuronal cell types, along with their spatial locations and defining phenotypic characteristics that can be aggregated as cellular populations that make up each brain region. This cell type classification scheme, organized as a taxonomy, should aim for a consensus across modalities and across mammalian species (for conserved types). Beyond the cell census, a Brain Cell Atlas would be embedded in a 3D Common Coordinate Framework (CCF) of the brain ⁴¹, in which the precise location and distribution of all cell types and their multi-modal features are registered and displayed. Such a cell-type resolution spatial framework will greatly facilitate integration, interpretation and navigation of various types of information for understanding brain network organization and function.

Here we present the cell census and atlas of cell types in one region of the mammalian brain, the primary motor cortex (MOp or M1) of mouse, marmoset and human, through an analysis with unprecedented scope, depth and range of approaches (**Fig. 1, Table 1**). MOp is important in the control of complex movement and is well conserved across species. Decades of accumulated anatomical, physiological, and functional studies have provided a rich knowledge base for the integration and interpretation of cell type information in MOp ^{42,43}. This manuscript describes a synthesis of results and findings derived from eleven core companion papers through a multilaboratory coordinated data generation within BICCN. We derive a cross-species consensus transcriptomic taxonomy of cell types and identify conserved and divergent gene expression and epigenomic regulatory signatures from a large and comprehensive set of single-cell/nucleus RNA-sequencing, DNA methylation and chromatin accessibility data. Focusing on mouse MOp, we map the spatial organization of transcriptomic cell types by multiplexed error-robust

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fluorescence in situ hybridization (MERFISH) and their laminar, morphological and electrophysiological properties by Patch-seq; we report the cell-type resolution input-output wiring diagram of this region by anterograde and retrograde tracing and investigate how axon projection patterns of glutamatergic excitatory neurons correlate with molecularly-defined cell types by Epi-Retro-Seq, Retro-MERFISH (the combination of MERFISH and retrograde labeling), and single-neuron full morphology reconstruction; we describe transgenic driver lines systematically targeting glutamatergic cell types based on marker genes and lineages. Finally, we integrate this vastly diverse array of information into a cohesive depiction of cell types in the MOp region with correlated molecular genetic, spatial, morphological, connectional, and physiological properties and relating them to traditionally described cell types. Such integration is illustrated in detail in example cell types with unique features in MOp: the layer 4 intratelencephalic-projecting (L4 IT) cells and layer 5 extratelencephalic-projecting (L5 ET) cells. This multitude of datasets are organized by the BRAIN Cell Data Center (BCDC) and made public through the BICCN web portal www.biccn.org. Key concepts and terms are described in Table 2, including anatomical terms for input and output brain regions for MOp, and hierarchical cell class, subclass and type definitions.

Table 1. Experimental and computational techniques used in this study and associated datasets

Feature	Experimental or analytic technique(s)	Abbreviations	References	Samples (e.g. # of cells or nuclei) in MOp/M1	Total samples in flagship and companion papers
Transcription	Single-cell mRNA sequencing	scRNA-Seq: SMART-Seq v4, 10x Chromium v2, v3	Background: ^{15,44} Companion: ⁴⁵	SMART-seq v4: 6,288 cells (mouse) 10x Chromium v2, v3: 193,824 cells (mouse)	1,163,727 cells
	Single nucleus mRNA sequencing	snRNA-Seq: SMART-Seq v4, 10x Chromium v2, v3	Background: 18,46,47 Companion: 45,48	SMART-seq v4: 6,171 nuclei (mouse) 10,534 nuclei (human) 10x Chromium v2, v3: 294,717 nuclei (mouse) 69,279 nuclei (marmoset) 15,842 nuclei (macaque) 76,533 nuclei (human)	1,100,168 nuclei

DNA methylation Open chromatin Combined	Single-nucleus methylcytosine sequencing 2 Single nucleus Assay for Transposase- Accessible Chromatin Single-nucleus chromatin	snmC-Seq2 snATAC-Seq	Background: 49 Companion: 45,48,50 Background: 21,51 Companion: 45,52	9,941 nuclei (mouse) 5,324 nuclei (marmoset) 5,222 nuclei (human) 79,625 nuclei (mouse)	110,294 nuclei 813,799 nuclei
transcription/ Open chromatin	accessibility and mRNA expression sequencing	SNARE-seq2	Background: 53 Companion: 48	(marmoset) 84,178 nuclei (human)	94,124 nuclei
Spatially resolved single-cell transcriptomi cs	Spatially resolved single-cell rranscriptomi Multiplexed error- robust fluorescence		Background: 28,29 Companion: ⁵⁴	~300,000 cells (mouse)	~300,000 cells
Clustering and data integration methods	Clustering - Hierarchical iterative clustering	scrattch.hicat	Background: 15,44 Companion: 45,48		
	Clustering - Metacell hierarchical clustering with dynamic tree pruning	tree-based method	Companion: 48		
	Clustering of snATAC-seq data	SnapATAC	Background: 55 Companion: 52		
	Clustering - Leiden clustering		Background: 56 Companion: 48		
	Multimodality and cross-species integration	LIGER, Seurat, SingleCellFusi on (SCF),scrattch. hicat	Background: 44,47,57-60 Companion: ^{45,48}		
Statistical validation	Cross-dataset replicability analysis	MetaNeighbor	Background: 61 Companion: 45,48		
Electrophysio logy, cellular morphology and	Combined in vitro slice physiology, biocytin cell filling, cytoplasm extraction	Patch-Seq, Smart-seq v2	Background: 30,62,63 Companion: 48,64,65	1,237 cells (mouse) 6 cells (macaque) 6 cells (human)	133 cells (mouse) 6 cells (macaque) 391 cells (human)

transcriptomi	and RNA-				
cs	sequencing				
Cellular morphology	Whole brain single cell full morphology reconstructions Barcoded anatomy	fMOST, MouseLight	Background:	151 cells (full morphology)	1,708 cells (full morphology)
and	resolved by		26,66,67		10,299 neurons
projection	sequencing	BARseq	Companion: 68,69		(BARseq)
Inter-areal circuit mapping	Anterograde tracing: PHAL, Viral tracers: AAV, Cre-dependent AAV, monosynaptic anterograde AAV- Cre	AAV, PHAL	Background: 70-74 Companion: ^{69,75}	22 experiments (mouse)	
	Retrograde tracing: CTB, viral tracers	RV, rabies, TRIO	Background: 70,76-78 Companion: ^{69,75}	40 experiments (mouse)	
	Retrograde viral labeling of neurons with defined projections followed by epigenome profiling	Epi-Retro-Seq		2,111 cells (mouse)	11,827 cells
Projection- specific profiling	Combined retrograde labeling and MERFISH	Retro-MERFISH	Companion: 54,79		
Genetic tools	Transgenic mouse lines	FlpO, Cre, CreER knockin lines; TIGRE- MORF/Ai166, MORF3 reporter line	Background: 80 Companion: ^{68,75} Stafford, Daigle, Chance et al., in preparation	6 knock-in driver lines 1 reporter line	26 knock-in lines

Table 2. Glossary

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Table 2. Glossal y						
Glossary			Definition	InterLex Identifiers		
Neuroanatomical regions described		described				
	(mouse), M1	Primary motor cortex, the main target of cellular diversity analyses	Primary motor cortex as defined in Fig 1 of the paper.	ILX:0770115		

	non-human primate)			
	L1, L2/3, L4, L5, L6, L6b	Layers within MOp	Cortical layers in primary motor cortex	ILX:0770170 ILX:0770171 ILX:0770172 ILX:0770173 ILX:0770179 ILX:0770180
_	s receiving axo etrograde label	nal projections from MOp ling studies		ILX:0770177
Cortical	(Secondary n	y somatosensory cortex), MOs notor cortex), TEa (Temporal rea), ACA (Anterior cingulate	Subset of cortical regions that receive axonal projections from primary motor cortex that were targeted in BICCN retrograde labeling studies.	ILX:0770178 ILX:0770117 ILX:0770116 ILX:0770118 ILX:0770120
Subcortical	STR (Striatum), TH (Thalamus), SC (Superior colliculus), VTA (Ventral tegmental area), HY (Hypothalamus), MB (Midbrain), P (Pons), MY (Medulla), claustrum		Subset of subcortical regions that receive axonal projections from primary motor cortex that were targeted in BICCN retrograde labeling studies.	ILX:0770167 ILX:0770122 ILX:0770123 ILX:0770124 ILX:0770137 ILX:0770165 ILX:0770126 ILX:0770127 ILX:0770127 ILX:0770125 ILX:0770128
Germinal sou	rces of cortica			
	MGE, CGE	Medial and caudal ganglionic eminences	The MGE is a progenitor domain within the ventral telencephalon that, together with the lateral ganglionic eminence (LGE), are the source of the majority of interneurons in the neocortex, hippocampus and olfactory bulb. In addition, oligodendrocytes arise from these regions and migrate into the developing cortex. The CGE is a progenitor domain within the ventral telencephalon that is a source of cortical interneurons in the striatum, neocortex and limbic system. The CGE is defined as a posterior region in which the medial and lateral eminences are fused to one structure. This structure is also a source of oligodendrocytes. Adapted from https://discovery.lifemapsc.com .	ILX:0770129 ILX:0770130

Definition of hierarchy	of terms used to	o describe cellular		
Cell class: T	op branches of	hierarchical tree	The top branches of theCN transcriptomic cell type hierarchy comprising neuronal and non-neuronal cells. Neuronal cells comprise inhibitory GABAergic cortical neurons and excitatory glutamatergic cortical neurons. Non-neuronal cells comprise glia and non-neural cells.	ILX:0770094
	Inh	GABAergic or inhibitory neurons, derived from MGE and CGE	Neuronal cell with a soma located in the cortex that uses GABA as a neurotransmitter and which exerts an inhibitory post-synaptic effect and derived from MGE and CGE.	ILX:0770098
Neuronal	Exc	Glutamatergic or excitatory neurons	Neuronal cell with a soma located in the cortex that uses glutamate as a neurotransmitter and exerts an excitatory post-synaptic effect.	ILX:0770097
	Glia	Non-neuronal cells of neuroectoderm origin	Non-neuronal brain cells of neuroectoderm origin	ILX:0770169
Non- neuronal	Non-neural	Cells of mesoderm, neural crest or yolk sac origin	Non-neuronal brain cells of mesoderm, neural crest or yolk sac origin	ILX:0770187
		ss, major groupings with across data modalities	Subset of class, major groupings with highly convergent evidence across data modalities	ILX:0770095
	Vip, Sst, Sst Chodl, Pvalb, and Meis2) Glutamatergic subclasses (L2/3 IT, L4/5 IT, L5 IT, L6 IT, L6 IT Car3, L5 ET, L5/6		Subclasses of GABAergic neurons distinguished by one or more marker genes	ILX:0770149 ILX:0770150 ILX:0770151 ILX:0770152 ILX:0770153 ILX:0770154 ILX:0770155 ILX:0770156 ILX:0770157 ILX:0770157 ILX:0770158 ILX:0770159 ILX:0770160
			Subclasses of glutamatergic neurons distinguished by anatomical location and projection pattern.	ILX:0770161 ILX:0770162 ILX:0770163
		IT: Intratelencephalic projecting	Excitatory glutamatergic neuron that projects to other telencephalic structures.	ILX:0770100
		ET: Extratelencephalic projecting	Excitatory glutamatergic neuron that projects to structures not derived from telencephalon	ILX:0770101

		NP: Near-projecting	Excitatory glutamatergic neuron that projects axons locally rather than long distance	ILX:0770103
		CT: Corticothalamic projecting	Excitatory glutamatergic neurons that project to the thalamus	ILX:0770102
	Glial subclass	ses (Astro, Oligo, OPC)	Subclasses of glial cells including astrocytes (Astro), oligodendrocytes (Oligo) and OPC cells (OPC)	ILX:0770141 ILX:0770140 ILX:0770139
	Non-neural st SMC, Peri, M	ubclasses (Endo, VLMC, licro, PVM)	Subclasses of non-neural cells including endothelial cells (Endo), vascular leptomeningeal cells (VLMC), smooth muscle cells (SMC), pericytes (Peri), microglia (Micro) and perivascular myeloid cells (PVM)	ILX:0770142 ILX:0770143 ILX:0770144 ILX:0770145 ILX:0770146 ILX:0770147
Cell type: Subset of subclass, finest resolution clustering achieved for a modality or a consensus clustering across modalities and/or species			Subset of subclass, finest resolution clustering achieved for a modality or a consensus clustering across modalities and/or species	ILX:0770096
Cluster: Data-driven cell set, synonymous with type		, synonymous with type	Data-driven cell set, synonymous with type	ILX:0770164

Major findings from this coordinated consortium project include:

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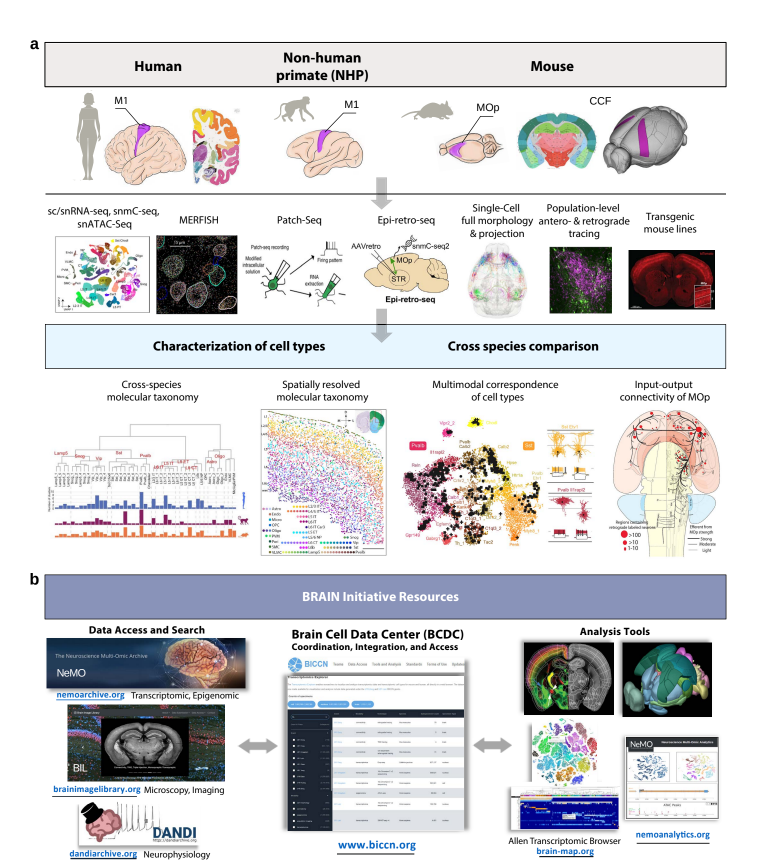
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- Combined single-cell transcriptomic and epigenomic analysis reveals a unified molecular genetic landscape of adult cortical cell types that integrates gene expression, chromatin state and DNA methylation maps.
- Combination of single-cell -omics, MERFISH-based spatially resolved single-cell transcriptomics and Patch-seq generates a census and atlas of cell types, including their population demographics of type, proportion, and spatial distribution across cortical layers and sublayers.
- Comparative analysis of mouse, marmoset and human transcriptomic types achieves a unified cross-species taxonomy of cortical cell types with their hierarchical organization that reflects developmental origins; transcriptional similarity of cell type granularity across species varies as a function of evolutionary distance.
- We observed both highly conserved gene expression and epigenomic signatures of cell identity across species, as well as a large set of species-specific cell type gene expression profiles suggesting a high degree of evolutionary specialization.
- The overall correspondence among transcriptomic, epigenetic, spatial transcriptomic, morphological, and intrinsic physiological datasets reinforces the transcriptomic classification of neuronal subclasses and distinctive types, demonstrating their biological validity and genomic underpinnings, and also reveals continuously varying properties along these axes among some neuronal subclasses and types.
- Multi-faceted anatomic studies yield a cellular resolution wiring diagram of mouse MOp anchored on major transcriptome-defined projection types, including input-output

- connectivity at subpopulation level and output pathways at genetically-defined single-cell level.
- The long-range axon projection patterns of individual glutamatergic excitatory neurons exhibit a complex and diverse range of relationships (between one-to-one and many-to-many) with transcriptomic and epigenetic types, suggesting another level of regulation in defining single-cell connectional specificity.
- Cell type transcriptional and epigenetic signatures can guide the generation of an extensive genetic toolkit for targeting glutamatergic pyramidal neuron types and fate mapping their progenitor types.
- Multi-site coordination within BICCN and data archives allows a high degree of standardization, computational integration, and creation of open data resources for community dissemination of data, tools and knowledge.

Figure 1. Summary of experimental and computational approaches taken as well as community resources generated by the BICCN. a, Comprehensive characterization of cell types in the primary motor cortex (MOp) of three mammalian species using multiple approaches spanning molecular, genetic, physiological and anatomical domains. Integration of these datasets leads to a cohesive multimodal description of cell types in the mouse MOp and a cross-species molecular taxonomy of MOp cell types. b, The multimodal datasets are organized by the Brain Cell Data Center (BCDC), archived in the Neuroscience Multi-omic (NeMO) Archive (for molecular datasets), Brain Image Library (BIL, for imaging datasets) and Distributed Archive for Neurophysiology Data Integration (DANDI, for electrophysiology data), and made publicly available through the BICCN web portal www.biccn.org.



RESULTS

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Molecular definition of cell types in MOp

The mouse MOp molecular taxonomy is derived from 9 datasets, including seven sc/snRNA-seq datasets and one each of snmC-Seq2 and snATAC-Seq datasets (companion paper ⁴⁵). The combined seven sc/snRNA-seq datasets (>700,000 cells total) had the advantages of large number of cells profiled using the droplet-based 10x Chromium v2 or v3 method and deep fulllength sequencing using the plate-based SMART-Seq v4 method, resulting in a consensus transcriptomic taxonomy for the mouse MOp with the greatest resolution compared to other data types, containing 116 clusters or transcriptomic types (t-types), 90 of which were neuronal types ⁴⁵. We used this mouse MOp transcriptomic taxonomy as the anchor for comparison and crosscorrelation of cell-type classification results across all data types. We further utilized two computational approaches, SingleCellFusion (SCF) and LIGER, to combine the seven transcriptomic with two epigenomic datasets and derive an integrated molecular taxonomy consisting of 56 neuronal cell types (corresponding to the 90 transcriptomic neuronal types) for the mouse MOp, with highly consistent molecular profiles based on transcriptomics, DNAmethylation, and open chromatin ⁴⁵ (Fig. 2a). Critically, this integrated taxonomy enabled us to link RNA transcripts with epigenomic marks identifying potential cell-type-specific cisregulatory elements (CREs) and transcriptional regulatory networks. Similarly, we established M1 cell type taxonomies for human (127 t-types) and marmoset (94 t-types) by unsupervised clustering of snRNA-seq data, followed by integration with epigenomic datasets (companion paper ⁴⁸).

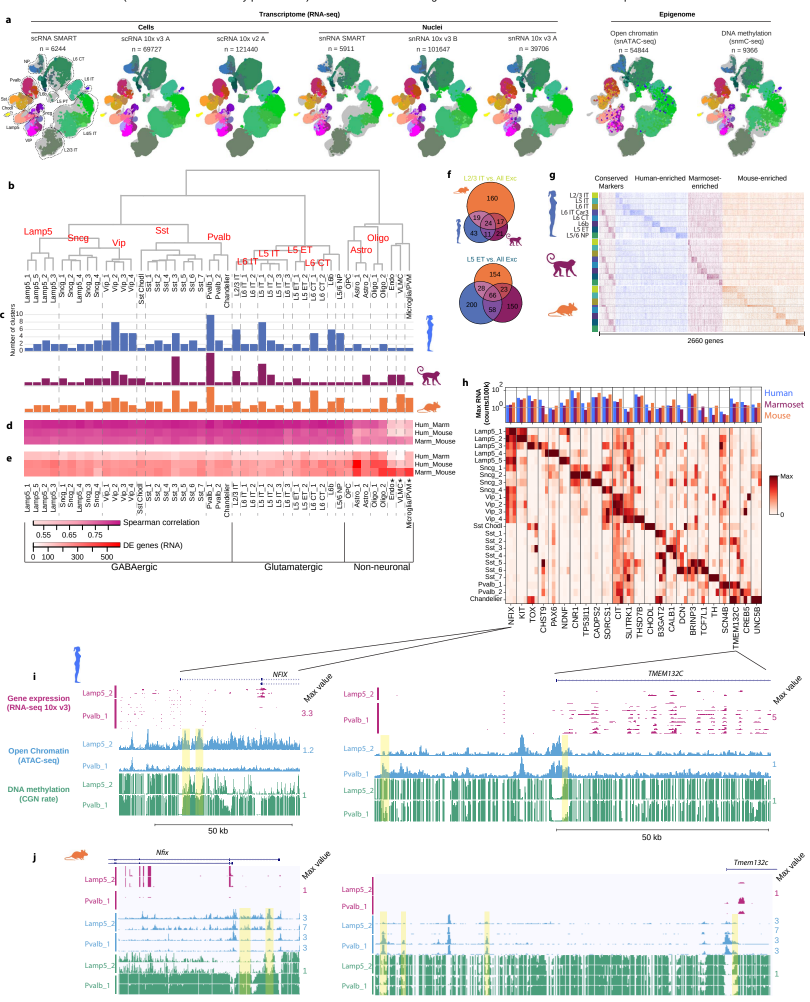
To establish a consensus classification of MOp/M1 cell types among mouse, human and marmoset, we integrated snRNA-seq datasets across species and identified 45 conserved transcriptomic types that spanned three major cell classes, including 24 GABAergic, 13 glutamatergic, and 8 non-neuronal types (Fig. 2b, Extended Data Fig. 1). These types were grouped into broader subclasses based on shared developmental origin for GABAergic inhibitory neurons [i.e., three caudal ganglionic eminence (CGE)-derived subclasses (Lamp5, Sncg and Vip) and two medial ganglionic eminence (MGE)-derived subclasses (Sst and Pvalb)], layer and projection pattern in mouse for glutamatergic excitatory neurons [i.e., intratelencephalic (IT), extratelencephalic (ET), corticothalamic (CT), near-projecting (NP) and layer 6b (L6b)], and non-neuronal functional subclass (e.g., oligodendrocytes and astrocytes) (Table 2). Note that the layer 5 extratelencephalic (L5 ET) neurons had been named as pyramidal tract (PT) neurons or subcerebral projection neurons (SCPN) in the literature 81,82; in this study we chose to use the name L5 ET for this subclass of neurons to be more representative across cortical areas and species (Supplementary Notes). The resolution of this cross-species conserved taxonomy was lower than that derived from each species alone, due to gene expression variations among species. The degree of species alignments varied across consensus types (Fig. 2c); some types could be aligned one-to-one (e.g., Lamp5 1, L6 IT 3), while others aligned several-to-several

(e.g., Pvalb_1, L2/3 IT, L5 IT_1). This may reflect over- or under-clustering, limitations in aligning highly similar cell types or species-specific expansion of cell-type diversity (companion paper ⁶⁵).

We hypothesized that cell types would share more similar gene expression profiles between human and marmoset than between either primate and mouse because primates share a more recent common ancestor. Indeed, we found that between primates, transcriptomic profiles of consensus cell types were more correlated and had 25-50% fewer differentially expressed (DE) genes than between primates and mouse (**Fig. 2d,e**). Three non-neuronal types had greater spearman correlations of overall gene expression (**Fig. 2d**, right columns) between marmoset and mouse likely because non-neuronal cells were undersampled in human M1 resulting in fewer rare types ⁴⁸. Robust conservation of cell types across mammals, including types with known specificity in electrical properties and connectivity such as chandelier cells and long-range projecting *Sst*-expressing cells (*Sst Chodl*), is strong evidence for the functional significance of these types.

Glutamatergic subclasses expressed many marker genes (using Seurat's FindAllMarkers function with test.use set to 'roc', >0.7 classification power) compared to other subclasses, and the majority of markers were species-specific (Fig. 2f,g). The evolutionary divergence of marker gene expression may reflect species adaptations or relaxed constraints on genes that can be substituted with others for related cellular functions. Subclasses also had a core set of marker genes that were conserved across all three species (Fig. 2g); these genes are candidates for consistent labeling of consensus cell types and for determining the conserved features of those cells that are central to their function. GABAergic consensus types also had conserved markers with similar absolute expression levels across species (Fig. 2h, bar plots) and relatively specific expressions compared to other cell types (Fig. 2h, heatmap). Marker genes of Lamp5_2 (NFIX) and Pvalb_1 (TMEM132C) GABAergic neurons showed evidence for cell-type-specific enhancers located in regions of open chromatin and DNA hypomethylation in both human (Fig. 2i) and mouse (Fig. 2j).

In summary, the multi-omic approach reveals a unified molecular genetic landscape of cortical cell types that integrates gene expression, chromatin state and DNA methylation maps and yields a robust molecular classification of cell types that is consistent between transcriptomic and epigenomic analyses. These studies further allow the identification of putative regulatory elements associated with cell type identity. Cell types are generally conserved between primates and rodents, and have a small number of conserved marker genes that are candidates for consistent labeling of conserved cell types.



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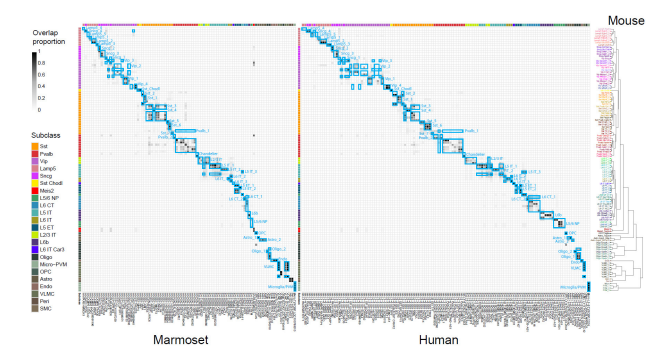
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Figure 2. MOp consensus cell type taxonomy. a, Integrated transcriptomic and epigenomic datasets using SCF show consistent molecular cell-type signatures as revealed by a lowdimensional embedding in mouse MOp. Each Uniform Manifold Approximation and Projection (UMAP) plot represents one dataset. Colors indicate different subclasses. b, Dendrogram of integrated human, marmoset, and mouse cell types based on single nucleus RNA-seq datasets (10x Chromium v3). c, Number of within-species clusters that are included in each cross-species cluster. d-e, For each consensus cluster, correlations (d) and differentially expressed (DE; Wilcoxon test, FDR < 0.01 and \log_e fold-change > 2) genes (e) between pairs of species. Asterisks denote non-neuronal populations that were under-sampled in human. f. Venn diagrams of shared DE genes between species for L2/3 IT and L5 ET glutamatergic neuron subclasses. g. Conserved and species-specific DE genes for all glutamatergic subclasses. Heatmap shows gene expression normalized by the maximum for each gene for up to 50 randomly sampled nuclei from each subclass and species. h, Conserved markers of GABAergic neuron types across three species. i-j, Genome browser showing transcriptomic and epigenetic signatures for gene markers of Lamp5 2 (NFIX) and Pvalb 1 (TMEM132C) GABAergic neurons in human (i) and mouse (j). Yellow bars highlight sites of open chromatin and DNA hypomethylation in the cell type with corresponding marker expression.



Extended Data Figure 1. Cluster overlap heatmap showing the proportion of nuclei in each pair of species clusters that are mixed in the cross-species integrated space. Cross-species consensus clusters are indicated by labeled blue boxes. Mouse clusters (rows) are ordered by the mouse MOp transcriptomic taxonomy dendrogram reproduced from ⁴⁵. Marmoset (left columns) and human (right columns) transcriptomic clusters (reproduced from ⁴⁸ are ordered to align with

mouse clusters. Color bars at top and left indicate subclasses of within-species clusters.

Spatially resolved cell atlas of the mouse MOp by MERFISH

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Sequencing-based single-cell methods require dissociation of cells from tissues, and hence the spatial organization of neuronal and non-neuronal cells, which is critical for brain function, is lost. To obtain a spatially resolved cell atlas of the mouse MOp region, we used MERFISH, a single-cell transcriptome imaging method 28,29 , to identify cell types *in situ* and map their spatial organization. We selected a panel of 258 genes (254 of which passed quality control) to image by MERFISH, on the basis of both prior knowledge of marker genes for major subclasses of cells in the cortex and marker genes differentially expressed in the neuronal clusters identified by the sn/scRNA-seq experiments, and we imaged $\sim 300,000$ individual cells across the MOp and its vicinity (companion paper 54).

Clustering analysis of the MERFISH-derived single-cell expression profiles resulted in a total of 95 cell clusters in MOp, including 42 GABAergic, 39 glutamatergic, and 14 non-neuronal clusters (**Fig. 3a,b**), as well as four distinct cell clusters observed exclusively outside the MOp (in striatum or lateral ventricle). These 95 clusters showed excellent correspondence with the 116 cell clusters identified by the sn/scRNA-seq datasets ⁵⁴. MERFISH analysis also revealed clusters not identified by scRNA-seq and vice versa, mostly in the form of refined splitting of clusters ⁵⁴.

The spatial distribution of the clusters derived from MERFISH showed a complex, laminar organization of cells in the MOp (Fig. 3c). MERFISH data divided glutamatergic neurons into IT, ET, NP, CT, and L6b subclasses, each of which were further divided into finer clusters. Many of these clusters adopted narrow distributions along the cortical depth direction that subdivided individual cortical layers, though often without discrete boundaries ⁵⁴. Notably, IT cells, the largest branch of neurons in the MOp, formed a largely continuous spectrum of cells with gradual changes both in their expression profiles and in their cortical depth positions, in a highly correlated manner ⁵⁴ (**Fig. 3d**). The five major subclasses of GABAergic neurons (Lamp5, Sncg, Vip, Sst and Pvalb) were also divided into finer clusters. Interestingly, many individual GABAergic clusters showed layered distribution as well, preferentially residing within one or two cortical layers ⁵⁴. Among the non-neuronal cell clusters, VLMCs formed the out-most layer of cells of the cortex, mature oligodendrocytes and some astrocytes were enriched in white matter, whereas other major subclasses of non-neuronal cells were largely dispersed across all layers. In addition to the laminar organization, MERFISH analysis also revealed interesting spatial distributions of cell types along the medial-lateral and anterior-posterior axes ⁵⁴. Overall, the 95 neuronal and non-neuronal cell clusters in the MOp form a complex spatial organization refining traditionally defined cortical layers.

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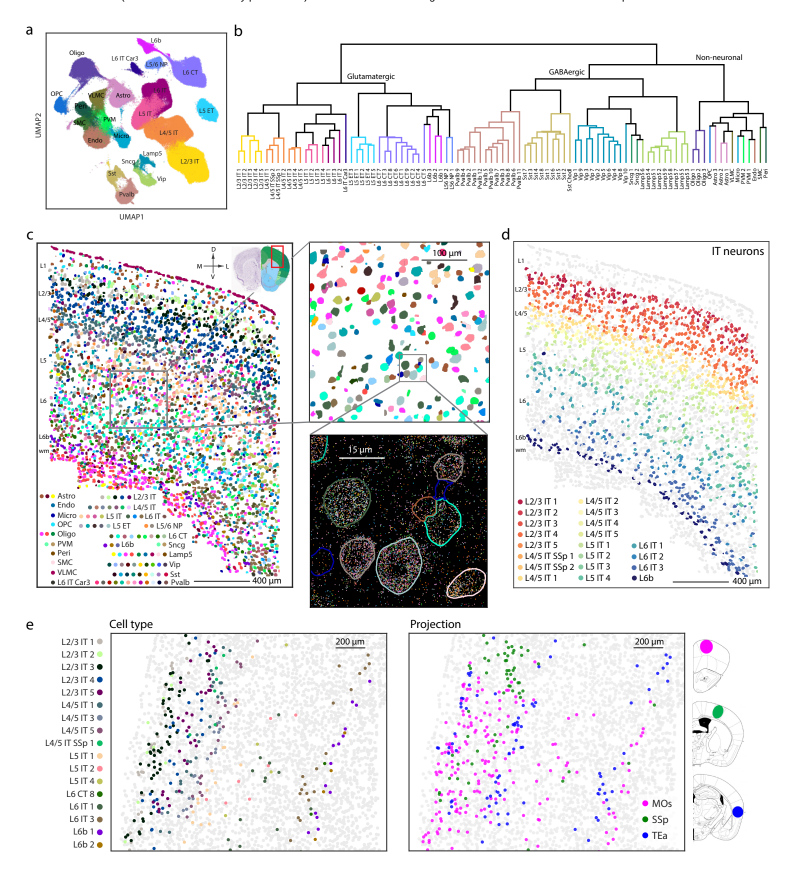
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Integration of retrograde tracing with MERFISH (Retro-MERFISH) further allowed us to map the projection targets of different neuronal cell types in the MOp. By injecting retrograde tracers into several different cortical areas (secondary motor cortex, primary somatosensory cortex, and temporal association area) and imaging retrograde labels together with the MERFISH gene panel in the MOp (**Fig. 3e**), we observed that all three examined target regions received inputs from multiple cell clusters in the MOp, primarily from IT cells. In addition, each IT cluster projected to multiple regions, with each region receiving input from a different composition of IT clusters ⁵⁴. Overall, the projection of MOp neurons does not follow a simple "one cell type to one target region" pattern, but rather forms a complex many-to-many network.

In summary, these MERFISH measurements revealed the spatial organization of neuronal and non-neuronal cell types in the MOp with an unprecedented resolution and granularity. Integration of MERFISH with retrograde tracing further allowed determination of both gene expression profiles and projection targets with single-cell resolution, revealing the compositions and spatial distributions of MOp neurons that project to several cortical regions.

Figure 3. In situ cell-type identification, spatial mapping and projection mapping of individual cells in the MOp by MERFISH. a, UMAP of the ~300,000 cells in the mouse MOp imaged by MERFISH. Cell clusters are grouped into 23 subclasses, and all cells in the same subclass are plotted in the same color. b, Dendrogram showing the hierarchical relationship among the 39 glutamatergic, 42 GABAergic, and 14 non-neuronal clusters in the mouse MOp identified by MERFISH, colored by the subclass that each cluster belongs to. c, Left: Spatial map of the cell clusters identified in a coronal slice (Bregma +0.90), with cells colored by their cluster identity as shown in the color index. Top right: Zoom-in map of the boxed region of the left panel. Bottom right: Spatial localization of individual RNA molecules in the boxed region of the top right panel, colored by their gene identity. The segmented cell boundaries are colored according to the cell clusters they belong to. d. The IT neurons in the same coronal slice as shown in c. The IT neurons are colored by their cluster identity, as shown in the color index, together with L6b cells in dark blue to mark the bottom border of the cortex. All other cells are shown in grey. e, Neuronal cluster identities of the cells projecting into three other regions of the brain, secondary motor cortex (MOs), primary somatosensory cortex (SSp), and temporal association area (TEa). Dye-labeled cholera toxin b (CTb) are used as retrograde tracers, and the CTb signals and the MERFISH gene panel are imaged in the MOp to determine both the cell cluster identities (left panel) and projection targets (right panel) of individual cells. Only clusters with 3 or more cells labeled by CTb are shown in color and the remaining cells are shown in grey.



Multimodal analysis of cell types with Patch-seq

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To characterize the electrophysiological and morphological phenotypes and laminar location of 375

376 the transcriptomically identified cell types, i.e., the t-types, we used the recently developed

Patch-seq technique ^{30,62}. We patched >1,300 neurons in MOp of adult mice, recorded their 377

378 electrophysiological responses to a set of current steps, filled them with biocytin to recover their 379

morphology (~50% of the cells) and obtained their transcriptomes using Smart-seq2 sequencing

(companion paper ⁶⁴). We mapped these cells to the mouse MOp transcriptomic taxonomy ⁴⁵.

Our dataset covered all major subclasses of glutamatergic and GABAergic neurons, with cells

assigned to 77 t-types (Fig. 4a). This allowed us to describe the electrophysiological and

morphological phenotypes of most t-types (see examples in Fig. 4b,c).

We found that the measured morpho-electrical (me) phenotype of a neuron was largely determined by its transcriptomic subclass, with different subclasses having distinct phenotypes. For example, Sst interneurons were often characterized by large membrane time constants, pronounced hyperpolarization sag, and rebound firing after stimulation offset. However, within each subclass, there was substantial variation in electrophysiological and morphological properties between t-types. This variation was not random but organized such that transcriptomically similar t-types had more similar morpho-electric properties than distant ttypes. For example, excitatory t-types from the IT subclasses with more similar transcriptomes were located also at adjacent cortical depths, suggesting that distances in t-space co-varied with distances in the me-space, even within a layer (Fig. 4g). Likewise, the electrophysiological properties of Sst interneurons varied continuously across the transcriptomic landscape ⁶⁴.

At the level of single t-types, we found that some t-types showed layer-adapting morphologies across layers (Fig. 4e,f) or even considerable within-type morpho-electric variability within a layer. For example, Vip Mybpc1 2 neurons had variable rebound firing strength after stimulation offset. Surprisingly few t-types were entirely homogeneous with regard to the measured morphoelectric properties (Fig. 4d).

In summary, we found that the morpho-electric phenotype of a neuron in MOp was primarily determined by the major subclass of neurons it belonged to, with different subclasses being transcriptomically as well as morpho-electrically distinct. Within each subclass, variation in electrophysiological and morphological properties often appeared to be continuous across the transcriptomic landscape, without clear-cut boundaries between neighbouring t-types.

Patch-seq also permits direct comparison of the morpho-electric properties of homologous cell types across species ⁴⁸. Here we focused our analysis on one of the most recognizable mammalian neuron types, the gigantocellular Betz cells found in M1 of primates and large carnivores. These neurons are predicted to be in the layer 5 ET (L5 ET) subclass ⁴⁸, which also contains the homologous corticospinal projecting neurons in the mouse. To allow cross-species

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440 441 analysis of primate Betz cells and mouse ET neurons, we first created a joint embedding of excitatory neurons in mouse, macaque and human, which showed strong homology across all three species for the L5 ET subclass (**Fig. 4h**). Patch-seq recordings were made from L5 neurons in acute and cultured slice preparations of mouse MOp and macaque M1. We also capitalized on a unique opportunity to record from neurosurgical tissue excised from the human premotor cortex, which also contains Betz cells, during an epilepsy treatment surgery. To permit visualization of cells in heavily myelinated macaque M1 and human premotor cortex, AAV viruses were used to drive fluorophore expression in glutamatergic neurons in slice culture.

Patch-seq cells in each species that mapped to the L5 ET subclass (Fig. 4h) were all large layer 5 neurons that sent apical dendrites to the pial surface (Fig. 4i, note truncation in human Betz cell). However, macaque and human L5 ET neurons were much larger, and had long "tap root" basal dendrites that are a canonical hallmark of Betz cells 83. Subthreshold membrane properties were relatively well conserved across species. For example, L5 ET neurons in all three species had a low input resistance, although it was exceptionally low in macaque and human (Fig. 4i). Conversely, suprathreshold properties of macaque and human Betz/ET neurons were highly specialized. Most notably, human and macaque neurons responded to prolonged suprathreshold current injections with a biphasic firing pattern in which a pause in firing early in the sweep was followed by a dramatic increase in firing late in the sweep (Fig. 4k). Intriguingly, we identified several genes encoding ion channels that were enriched in macague and human L5 ET neurons compared with mouse (Fig. 41). These primate specific ion channels may contribute to the distinctive suprathreshold properties of primate ET neurons. Together this indicates that primate Betz cells are homologous to mouse thick-tufted L5 ET neurons, but display phenotypic differences in their morphology, physiology and gene expression. Similar to transcriptomics, these results indicate strong conservation of cell subclasses but with significant species specializations in anatomical and functional properties.

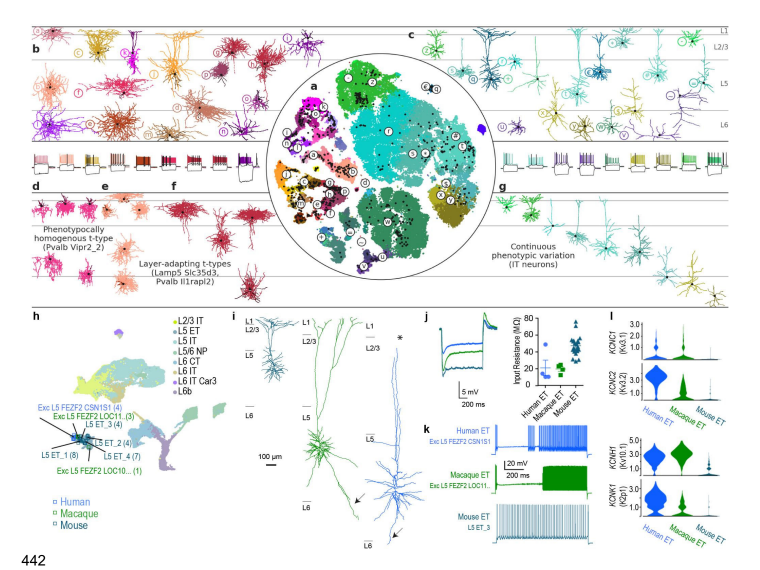


Figure 4. Correspondence between transcriptomic and morpho-electrical properties of mouse MOp neurons by Patch-seq, and cross-species comparison of L5 ET neurons. a, t-SNE of the scRNA-seq 10x v2 dataset ⁴⁵ with the Patch-seq neurons (black dots) positioned on top of it ⁸⁴. b, Examples of GABAergic interneuron morphologies and electrophysiological recordings (below). Letters refer to cells marked in a. c, Examples of glutamatergic excitatory neuron morphologies and electrophysiological recordings. d, Example of a phenotypically homogenous t-type (Pvalb Vipr2_2, chandelier neurons). e-f, Two examples of t-types showing layer-adapting morphologies (e, Lamp5 Slc35d3, neurogliaform cells; f, Pvalb Il1rapl2, fast-spiking basket cells). g, Example of a transcriptomic subclass (excitatory IT neurons) that shows continuous within-subclass co-variation between distances in transcriptomic space and morphological space (compare the color ordering in a (right) with the color ordering in g. h, UMAP visualization of cross-species integration of snRNA-seq data for glutamatergic neurons isolated from mouse, macaque and human, with colors corresponding to cell subclass. Patch-seq samples mapping to various ET neuron types are denoted by squares, color-coded by species. i,

Dendritic reconstructions of L5 ET neurons. The human (Exc L5 FEZF2 CSN1S1) and macaque (Exc L5 FEZF2 LOC114676463) neurons display classical Betz cell features, including taproot dendrites (arrows). Note, the human neuron is truncated (asterisk) before reaching the pial surface. i, Voltage response of mouse, macaque and human ET neurons to a 1 s, -300 pA current injection (left). Input resistance is low in all species, but exceptionally low in human and macaque Betz cells. Error bars represent SEM (right; macaque n=4, human n=4, mouse n=22; FDR corrected two-sided Wilcoxon ranked sum test (human vs mouse W=12, p = 0.31, d=2.09; human vs monkey W = 5, p = .49, d = .08; monkey v mouse W = 0 p = .0004., d = 2.5). k, Example spike trains in response to a 10s suprathreshold current injection. Macague and human L5 ET neurons tended to respond with a distinctive, biphasic firing pattern, I, Violin plots of enriched potassium channel gene expression in human and macaque compared to mouse L5 ET neurons.

Multimodal correspondence by Epi-Retro-Seq

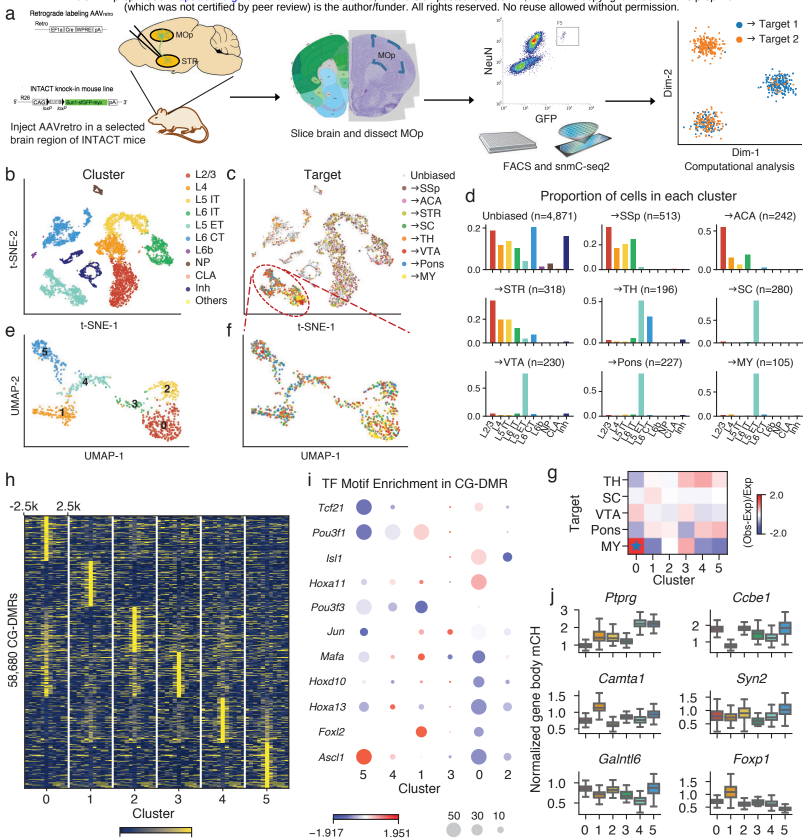
To obtain a comprehensive view of the molecular diversity among projection neurons in MOp, we developed Epi-Retro-Seq (companion paper ⁷⁹) and applied it to mouse MOp neurons that project to each of the 8 selected brain regions that receive inputs from MOp (**Fig. 5a**). The target regions included two cortical areas, SSp and anterior cingulate area (ACA), and six subcortical areas, striatum (STR), thalamus (TH), superior colliculus (SC), ventral tegmental area and substantia nigra (VTA+SN), pons, and medulla (MY). Specifically, we injected the retrograde tracer rAAV2-retro-Cre ⁷⁷ into the target region in INTACT mice ⁸⁵, which turned on Credependent GFP expression in the nuclei of MOp neurons projecting to the injected target region. Individual GFP-labeled nuclei of MOp projection neurons were then isolated using fluorescence-activated nucleus sorting (FANS). Single-nucleus methylcytosine sequencing (snmC-Seq2) ⁴⁹ was performed to profile the DNA methylation (mC) of each single nucleus.

After removing low-quality cells, potential doublets, and non-neuronal cells, we obtained high-quality methylomes for 2,111 MOp projection neurons. When co-clustering them with MOp neurons collected without enrichment of specific projections, we observed a precise agreement among all of the major cell subclasses (**Fig. 5b,c**), demonstrating the robustness of Epi-Retro-Seq to classify cell types. Although neurons projecting to different target regions were not completely separated on t-SNE, we observed the explicit enrichment of cortico-cortical and cortico-striatal projecting neurons in IT subclasses (L2/3, L4, L5 IT, L6 IT, and L6 IT Car3), and cortico-subcerebral projecting neurons in L5 ET. Many cortico-thalamic projecting neurons were also observed in L6 CT subclass (**Fig. 5d**). These observations are consistent with the known laminar distribution of the cortico-cortical and cortical-subcortical projection neurons ⁸¹, reflecting the high quality of retrograde-labeling of neuronal nuclei in our Epi-Retro-Seq dataset.

 The enrichment of L5 ET neurons in the Epi-Retro-Seq data (40.2% vs. 5.62% in unbiased profiling of MOp using snmC-seq2) allowed a more detailed investigation of the subtypes of L5 ET neurons which are known to project to multiple subcortical targets in TH, VTA+SN, pons and MY ⁸¹. The 848 L5 ET neurons further segregated into 6 clusters (**Fig. 5e,f**). MY-projecting neurons showed a clear enrichment in L5 ET cluster 0 (**Fig. 5f,g**), in agreement with scRNA-Seq data for anterolateral motor cortex (ALM), part of MOs ^{15,86}. We used gene body non-CG methylation (mCH) levels to integrate the L5 ET Epi-Retro-Seq data with the ALM Retro-seq data and also observed the enrichment of MY-projecting cells in the same cluster ⁷⁹.

A major advantage of DNA methylation profiling of neurons is its ability to obtain information for both genes and cis-regulatory elements. Specifically, mCH at gene bodies is strongly anticorrelated with gene expression in neurons, while promoter-distal differentially CG-methylated regions (CG-DMRs) are reliable markers of regulatory elements such as enhancers ²⁰. We thus identified 511 differentially CH-methylated genes (CH-DMGs) and 58,680 CG-DMRs across the L5 ET clusters (**Fig. 5h**). We also inferred transcription factors (TFs) that may contribute to defining the cell subclusters by identifying enriched TF-binding DNA sequence motifs within CG-DMRs (**Fig. 5i**). For example, *Ascl1* is a transcription factor whose motif was significantly enriched in the MY-projecting cluster. Previous studies had shown its necessity for neuronal differentiation and specification in multiple regions of the nervous system ^{87,88}. In addition, 230 hypo-CH-DMGs were identified between the MY-projecting cluster and other projection neurons. Interestingly, one of the most differentially methylated genes is *Ptprg* (**Fig. 5j**), which interacts with contactin proteins to mediate neural projection development ⁸⁹.

In summary, Epi-Retro-Seq mapping data for MOp revealed specific enrichment of MY-projecting neurons in one of the molecularly-defined subpopulations of MOp L5 ET neurons, allowing identification of regulatory elements for this unique cell type. In addition to MOp, we have performed 63 Epi-Retro-Seq mapping experiments for 7 cortical regions, comprising 26 cortico-cortical projections and 37 cortico-subcortical projections ⁷⁹. Together, these epigenomic mapping data for projection neurons facilitates the understanding of gene regulation in establishing neuronal identity and connectivity, by discovering projection-specific gene regulatory elements which can be used to target specific types of projection neurons.



-log10 P-value

mCH zscore

Cluster

Cluster

1.0 mCG 0.3

Figure 5. Epi-Retro-Seq links molecular cell type with distal projection targets. a,

Workflow of Epi-Retro-Seq. **b, c,** UMAP embedding of MOp cells profiled by Epi-Retro-Seq (n=2,115) and unbiased snmC-Seq2 (n=4,871) computed with 100kb-bin-level mCH, colored by subclasses (**b**) or projection targets (**c**). **d,** Distribution across subclasses of neurons from unbiased snmC-Seq2 and neurons projecting to each target. **e, f,** UMAP embedding of L5 ET cells in MOp profiled by Epi-Retro-Seq (n=848) computed with 100kb-bin-level mCH, colored by clusters (**e**) or projection targets (**f**). **g,** Enrichment of L5 ET neurons projecting to each target in each cluster. * represents FDR<0.05. **h,** mCG levels at CG-DMRs identified between the six clusters and their flanking 2.5k regions. Top 100 DMRs in each cluster were shown. **i,** TF motif enrichment in CG-DMRs in each cluster. Color represents z-scored gene-body mCH level of the TFs, and size represents -log10 *P* value of motif enrichment in the CG-DMRs. **j,** Boxplots of

TFs, and size represents -log10 P value of motif enrichment in the CG-DMRs. j, Boxplots of

normalized mCH levels at gene-bodies of example CH-DMGs in the six clusters. Numbers of cells represented by the boxes are 242, 165, 118, 42, 119, and 162 for the six clusters. The

elements of boxplots are defined as: center line, median; box limits, first and third quartiles;

whiskers, 1.5× interquartile range.

MOp projection neuron types and input-output wiring diagram

Building upon the molecularly defined and spatially resolved cell atlas (**Fig. 3**) and the multimodal correspondence between gene expression and morpho-electric properties of MOp neurons (**Fig. 4**), we next describe a comprehensive cellular resolution input-output MOp wiring diagram. To achieve this, we combined classic tracers, genetic viral labeling in Cre driver lines and single neuron reconstructions with high-resolution, brain-wide imaging, precise 3D registration to CCF, and computational analyses (companion paper ⁶⁹).

First, we systematically characterized the global inputs and outputs of MOp upper limb (MOpul) region using classic anterograde (PHAL) and retrograde (CTb) tract tracing ⁶⁹ (**Fig. 6a**). At the macro-scale, MOp-ul projects to more than 110 gray matter regions and cervical spinal cord, and ~60 structures in the cerebral cortex and thalamus project back to MOp-ul.

Next, we generated a fine-grained areal and laminar distribution map of multiple MOp-ul projection neuron populations using retrograde pathway-tracing. Accordingly, we identified 25 distinct neuron projection types based on their unique combinations of axonal targets and laminar somatic distributions (**Fig. 6b**, top; for details see ⁶⁹). For example, IT cells (e.g. TEatargeting or contralateral MOp-targeting) are distributed throughout L2-L6b; ET cells (ponsor medulla-targeting) are distributed primarily in L5b and most CT (posterior thalamic nucleus-targeting) neurons are distributed in L6a.

In parallel with these tracer-labeled, projection- and layer-defined cell types, we quantitatively characterized the distribution patterns of neuronal subpopulations in the MOp-ul labeled in 28

Cre-expressing "driver" lines (**Fig. 6b**, bottom). These lines selectively label neurons from different IT (e.g. Cux2, Plxnd1, Tlx3), L5 ET (Rbp4, Sim1, Fezf2), and CT (Ntsr1, Tle4) subpopulations with distinct laminar distributions ^{75,90,91}.

Subsequently, we used viral tracers to systematically examine MOp-ul cell-type-specific inputs and outputs (**Fig. 6c**). First, neurons projecting *to* Cre-defined starter cells were labeled using transsynaptic rabies viral tracing methods; an example from the *Tlx3* L5 IT line is shown in **Fig. 6c** (upper left, red). Projections *from* MOp were labeled following AAV-GFP injections into C57BL6/J mice, revealing patterns consistent with PHAL tracing results (**Fig. 6a**). Projections from L2/3 IT, L4 IT, L5 IT, L5 ET, and L6 CT cells were mapped following injections of Cre-dependent viral tracers into Cre lines selective for these laminar- and projection- cell subclasses ⁷¹. Most Cre line anterograde tracing experiments revealed a component of the overall output pathway (**Fig. 6c**). For example, the L6 *Ntsr1* line revealed a typical CT projection pattern with dense projections specific to thalamic nuclei. This result is consistent with labeling from retrograde injections in various thalamic nuclei (PO, VAL, VM) and cortical areas such as MOs and SSp (**Fig. 6b**, top). Further characterization of the distinctive projection patterns of several IT, L5 ET, and CT driver lines is provided in the anatomy companion paper ⁶⁹.

To further refine the projection neuron characterization, we carried out single cell analysis by combining sparse labeling, high-resolution whole-brain imaging, complete axonal reconstruction and quantitative analysis (companion papers ^{68,69}); additional analysis was also conducted using BARseq ⁶⁹, a high-throughput projection mapping technique based on *in situ* sequencing ⁶⁷. We augmented the full morphology reconstruction dataset with publicly available single cell reconstructions in MOp from the Janelia Mouselight project ²⁶. We systematically characterized axonal projections of 151 single MOp pyramidal neurons. This analysis revealed a rich diversity of projection patterns within the IT, ET and CT subclasses (Fig. 6c,d). For example, individual L6 neurons display several distinct axonal arborization targets that likely contribute to the composite subpopulation output described for the Ntsr1 and Tle4 diver lines (Fig. 6d). Confirming and extending previous reports 86, we characterized detailed axonal trajectories and terminations of two major types of L5b ET cells, namely medulla-projecting and non-medulla projecting neurons; both types may collateralize in the thalamus and terminate in the midbrain (Fig. 6d). Individual IT cells across L2-L6 also generate richly diverse axonal trajectories (detailed in ^{68,69}. Further analyses of complete single neuron morphologies, precisely registered in the CCF, will provide the ultimate resolution toward defining anatomical cell types and clarify the anatomical heterogeneity described at the subpopulation level.

In summary, combining multiple approaches complementary in their coverage, throughput, and resolution, we provide a comprehensive identification of major projection neuron types with

correspondence to molecular markers. We further delineate their input-output patterns at the subpopulation level and describe projection patterns at single-cell resolution, deriving the first multi-scale wiring diagram of MOp. A major future goal is to link these anatomic and especially projection types with transcriptomic types (**Fig. 2b**), with precise registration to a spatial atlas (e.g. **Fig. 3e**).

a. Global inputs and outputs of the MOp-ul

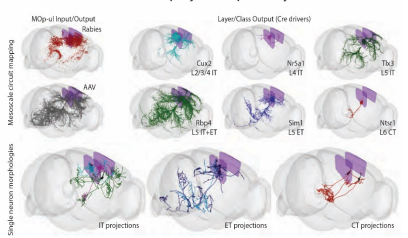
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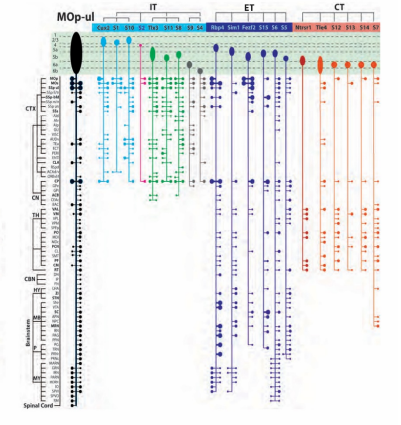
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c. 3D view of multi-scale projection pathways



d. MOp neuron type schema



b. Characterization of MOp neuron types

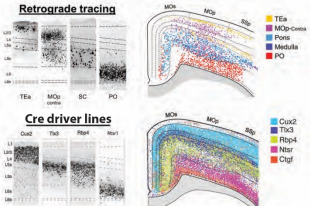


Figure 6. Global wiring diagram and anatomical characterization of MOp-ul neuron types. a. Flatmap representation of the MOp-ul input/output wiring diagram. Black lines and red dots indicate axonal projections (outputs) and retrograde labeling sources (inputs), respectively, with line thickness and dot sizes representing relative connection strengths. Most MOp-ul projection targets in the cortex and thalamus also contain input sources, suggesting bidirectional connections. The flatmap is adapted from the Swanson Brainmap 4.0 and the mouse brain flatmap ⁹². **b,** MOp-ul neurons classified by projection targets or transgenic *Cre* expression. (Top) Retrograde tracing using CTB revealed layer-specific distributions of MOpul neurons with respect to their major projection targets. Representative images (left) show neurons labeled by CTB injections into cortical areas (TEa, contralateral MOp), superior colliculus (SC) in the midbrain, and posterior complex (PO) of the thalamus. Detected cells were pseudo-colored and overlaid onto a schematic coronal section near the center of MOp-ul (right). MOp neurons that project to TEa are distributed in L2 and L5 (yellow), to the contralateral MOp in L2-L6b (purple), to targets in the pons and medulla in L5b (blue), and to thalamus in L6a (red). (Bottom) The distribution of neurons labeled in 28 transgenic Cre lines was mapped in MOp and across the whole cortex. Images (left) show laminar patterns of Cre+ nuclei in MOp-ul from four driver lines (Cux2, Tlx3, Rbp4, and Ntsr1). Detected nuclei from these lines, plus the Ctgf-Cre line, were pseudo-colored and overlaid onto a schematic coronal section near the center of MOp-ul (right). Cre+ nuclei are found in L2-4 in Cux2; L5a and superficial L5b in Tlx3; L5a and L5b in Rbp4; L6a in Ntsr1, and L6b in Ctgf. c, 3D views show brain-wide MOp input-output patterns at the population and single cell resolution. (Top left) Regional MOp inputs and outputs were mapped using retrograde (in red, example shows rabies tracing from the *Tlx3-Cre* driver line) and anterograde (in black, example shows AAV-EGFP) tracing methods. (Top right) Whole-brain axonal trajectories from 6 Cre line-defined subpopulations labeled with Cre-dependent AAV tracer injections at the same MOp-ul location. (Bottom) Individual projection neurons were fully reconstructed following highresolution whole-brain imaging of sparsely labeled cells. Representative examples of IT, ET, and CT neurons are shown in each panel. The two ET examples represent distinct projectiontypes; medulla (dark blue)- and non-medulla-projecting (light blue). 3D renderings were generated following registration of projection and reconstruction data into CCFv3 using BrainRender ⁹³. **d,** Projection patterns arising from major cell types, IT, ET and CT, with corresponding Cre-line assignment and somatic laminar location, compared with the overall projection pattern from the MOp-ul region (left, black). Along each vertical output pathway, horizontal bars on the right and left sides represent ipsilateral and contralateral collaterals, respectively, with dot sizes indicating the strength of axonal terminals in different targets. Brain structure nomenclature adopted from ARA 94.

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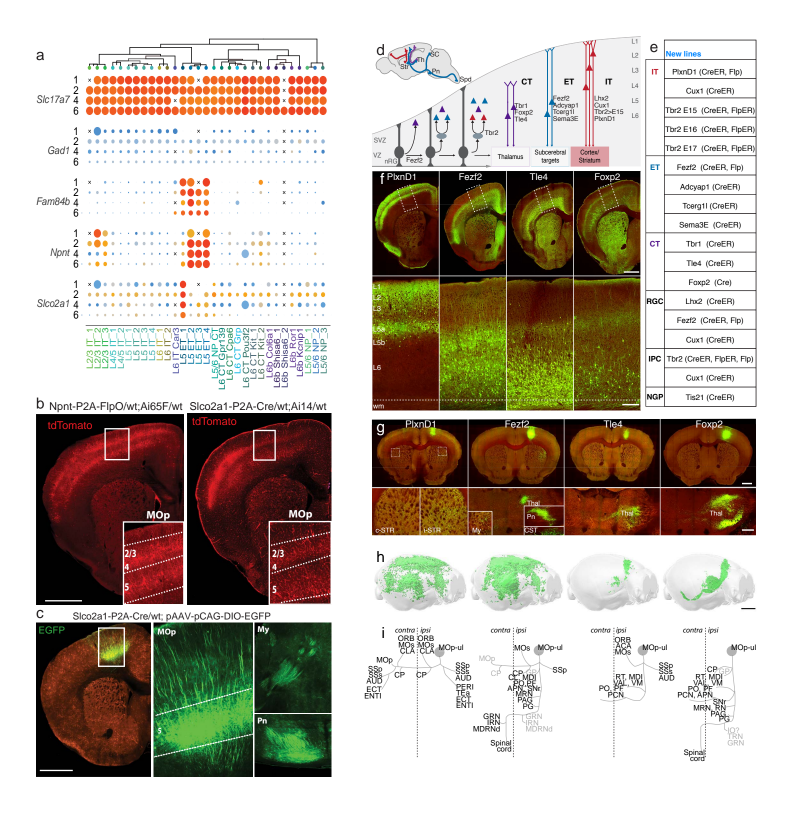
The identification and classification of MOp cell types based on single-cell integration of transcriptomes and epigenomes (**Fig. 2**), spatially resolved single-cell transcriptomics (**Fig. 3**) and anatomical and physiological analysis (**Fig. 4-6**) provides deep insights into the molecular basis of cellular diversity. In addition to establishing a principled basis for a taxonomy of brain cell types, knowledge of cellular gene expression also provides information to create mouse models in which genetically encoded reporters and actuators are targeted to these molecularly defined cell types ³³.

As an embodiment of this approach, we used CRISPR/Cas-9-mediated homologous recombination in ES cells to generate genetically modified mice (Stafford, Daigle, Chance et al., companion manuscript in preparation) in which sequences encoding FlpO and Cre recombinases were targeted respectively to Npnt and Slco2a1, genes whose differential expression discriminates between two types of L5 ET neurons with distinct subcortical projection target specificities ^{15,86}. Confirming the assignment of *Npnt*- and *Slco2a1*-expressing cells to subsets of L2/3 IT and L5 ET neurons in the consensus transcriptomic taxonomy (Fig. 7a), FlpO- and Cre-dependent tdTomato reporter expression in Npnt-P2A-FlpO:Ai65F and Slco2a1-P2A-Cre:Ai14 mice localized to these cortical cell layers in MOp (Fig. 7b). In Npnt mice, both L2/3 and L5 neurons were labeled. In Slco2a1 mice, predominantly L5 neurons were labeled. It is noteworthy that Slco2a1 labeled cells occupying a deeper sub-lamina of L5 than those targeted by Npnt, in accord with a previous report describing the two types of L5 ET neurons ⁸⁶ (see also **Fig. 9** below). To test the projection specificity of neurons labeled by these novel genetic tools, we injected a recombinant AAV encoding a Cre-dependent EGFP reporter into deep L5 in MOp of a Slco2a1-P2A-Cre mouse (Fig. 7c). Consistent with previous studies ⁸⁶ as well as those described in **Figures 5, 6 and 9** (below), GFP-labeled axon terminals were found in pontine gray and medulla, indicating that this mouse line labels the medullaprojecting L5 ET cell type.

To expand on cell type driver lines, we further built a genetic toolkit for cortical pyramidal neurons (PyNs) with more comprehensive coverage of projection types and with combinatorial strategies for improved specificity (companion paper ⁷⁵). First, we generated and characterized a set of 15 Cre and Flp gene knockin mouse driver lines for targeting major PyN subpopulations and progenitor types, guided by knowledge in their gene expression as well as developmental genetic programs (**Fig. 7d,e**). These include the broad CT (*Tbr1*, *Tle4*, *Foxp2*), ET (*Fezf2*, *Adcyap1*, *Tcerg1l*, *Sema3e*) and IT (*Plxnd1*, *Cux1*, and *Tbr1* late embryonic inductions) subclasses as well as subpopulations within these subclasses. When crossed with reporter alleles, these driver lines activated reporter expression that precisely recapitulated endogenous expression patterns highlighted here with 4 representative lines (**Fig. 7f**): L2/3 and L5a for IT-*Plxnd1* (IT^{*Plxnd1*}), L5b and L6 for ET- *Fezf2* (ET^{*Fezf2*}), L6 for CT-*Tle4* and CT- *Foxp2* (CT^{*Tle4*}, CT^{*Foxp2*}). To examine the projection pattern of these driver-defined subpopulations, we converted inducible CreER expression to constitutive Flp expression

 followed by MOp injection of a Flp-dependent AAV reporter vector (**Fig. 7g-i**). Largely as expected, IT^{Plxndl} projected to multiple ipsi- and contra-lateral cortical areas and the striatum/caudate putamen; ET^{Fezf2} projected robustly to several ipsi-lateral cortical sites, striatum, and numerous subcortical targets including thalamus, medulla and the corticospinal tract; CT^{Tle4} projected to a set of highly specific thalamic nuclei. Surprisingly, CT^{Foxp2} projected to a set of specific thalamic nuclei as well as to midbrain, brainstem and corticospinal tract. Further characterization of this set of new driver lines (**Fig. 7e**) is presented in ⁷⁵.

Together, these tools and strategies establish an experimental approach for accessing hierarchically organized neuronal cell types at progressively finer resolution. Such genetic access will enable an integrated multi-modal analysis to further validate and characterize these cell populations as well as to explore their multi-faceted function in neural circuit operation and behavior.--



709 Figure 7. Genetic tools for targeting cortical glutamatergic projection neuron types. a. 710 Dendrogram of excitatory cells types within the MOp transcriptomic taxonomy followed by the 711 proportion of cells (dot size) expressing pan-excitatory or -inhibitory marker genes (Slc17a7 or 712 Gad1), or L5 ET marker genes (Fam84b, Npnt, and Slco2a1). Expression results from four 713 different single cell RNA-seq platforms are shown: 1: scRNA-seq SMART-Seq; 2: scRNA-seq 10X v3 A; 4: snRNA-seq SMART-Seq; 6: snRNA-seq 10X v3 B 45. b, Representative images of 714 715 native tdTomato fluorescence from MOp of Npnt-P2A-FlpO; Ai65F and Slco2a1-P2A-Cre; Ai14 716 animals. Reporter expression was observed in L2/3 and L5 neurons with both driver lines and in 717 the vasculature with only the Slco2a1 line. c. Representative images of native EGFP 718 fluorescence from MOp, My (medulla), and Pn (pons) in the brain of an Slco2a1-P2A-Cre 719 animal injected in MOp with a Cre-dependent reporter AAV (pCAG-FLEX-EGFP-WPRE). 720 Robust reporter expression in L5 neurons was observed at the injection site (MOp) and in fibers 721 terminating in My and Pn. d, Schematic (upper left panel) depicting several major pyramidal 722 neuron (PyN) projection classes that mediate intra-telencephalic streams (IT-red; cortical and 723 striatal) and cortical output channels (ET-blue, CT-purple). Str. striatum; Th, thalamus; SC, 724 superior colliculus; Spd, spinal cord. Developmental trajectory of PyNs (lower panel) depicting lineage progression from progenitors to mature PyNs across major laminar and projection types. 725 726 Genes used to target progenitor and PyN subpopulations are listed according to their cellular 727 expression patterns. VZ, ventricular zone; SVZ, subventricular zone. e, Table presenting new 728 gene knockin driver mouse lines targeting PvN progenitors and projection types. RGC, radial 729 glia cell; IPC, intermediate progenitor cell; NGP, neurogenic progenitor. f, Cre recombination 730 patterns visualized through reporter expression (green) and background autofluorescence (red) 731 from four driver/reporter lines PlexinD1-2A-CreER (PlxnD1); Snap25-LSL-EGFP, Fezf2-2A-732 CreER (Fezf2); Ai14, Tle4-2A-CreER (Tle4); Snap25-LSL-EGFP and Foxp2-IRES-Cre 733 (Foxp2);AAV9-CAG-FLEX-EGFP (systemic injection). Top row: coronal hemisections 734 containing MOp. Bottom row: a segment of MOp (dashed lines, top row) with laminar 735 delineations. CreER Tamoxifen (TM) inductions were at P21 and P28. g, Anterograde tracing 736 from PvN subpopulations in MOp. CreER drivers were crossed with a Rosa26-CAG-LSL-Flp 737 mouse, and postnatal TM induction to convert to constitutive Flp expression for anterograde 738 tracing with a Flp-dependent AAV vector expressing EGFP (AAV8-CAG-fDIO-TVA-EGFP). 739 Representative images of native EGFP fluorescence from the MOp injection site (top row) from 740 cell-type-specific viral vector (green) and background autofluorescence (red) at selected 741 subcortical projection targets for four driver lines: Th; Str; cerebral peduncle (cp), Pn, My and 742 corticospinal tract (CST). h, Whole-brain three dimensional renderings of axon projections 743 registered to the CCFv3 for each PvN subpopulation in the MOp cortex (parasagittal view). i. 744 Schematics of main projection targets for each PyN subpopulation. Vertical dashed line indicates midline; filled circle indicates MOp injection site. Scale bars: hemisections (f & g) and h, 1mm; 745 bottom row in f, 200µm; bottom row in g, 500µm; h, 2 mm. 746 747

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Integrated multimodal characterization reveals L4 IT neurons in MOp

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To investigate if our collective multimodal characterization can lead to an integrated understanding of cell types in MOp, we selected two case studies to demonstrate convergence of multiple corresponding properties onto specific cell types.

Traditionally MOp has been considered an agranular cortical area, defined by the lack of a cytoarchitectonic layer 4 which usually contains spiny stellate or star pyramid excitatory neurons. However, a previous study challenged this view and presented evidence that L4 neurons similar to those typically found in sensory cortical areas also are present in MOp ⁹⁵. Here as the first case study, we used multimodal evidence to confirm the presence of L4-like neurons in mouse MOp and possibly in primate M1 as well (**Fig. 8**).

We first performed a joint clustering (see Methods) and UMAP embedding of all IT cells (excluding the highly distinct L6 IT Car3 cells) from 11 different mouse molecular datasets, including 6 sc/snRNA-seq datasets, and the snmC-Seq2, snATAC-Seq, Epi-Retro-Seq, MERFISH and Patch-seq data (Fig. 8a). This resulted in 5 joint clusters, mostly along a continuous variation axis moving from L2/3 to L4/5 to L5 to L6. The joint clustering enabled linkage of the cells independently profiled by each individual modality into types transcriptomic, epigenomic, spatially resolved transcriptomic, and morpho-electrictranscriptomic, and cross-correlation of these disparate properties. Consequently, we identified epigenomic peaks linked to cluster-specific marker genes - Cux2 for L2/3 IT and L4/5 IT (1), Rspo1 for L4/5 IT (1), Htr2c for L4/5 IT (2-3), and Rorb for L4/5 IT and L5 IT (Fig. 8b, cluster names from SingleCellFusion). MERFISH data also showed that L4/5 IT and L5 IT cells occupied distinct layers, and the L4/5 IT type expressed Rspo1 (Fig. 8c), a L4 cell type marker in sensory cortical areas identified in previous studies ¹⁵. Transcriptomic IT types from mouse corresponded well with those from human and marmoset, but such correspondence was mostly at main branches or subclass level while significant confusions existed at single cluster level (Fig. 8d), likely due to the substantial gene expression variation between rodents and primates (Fig. 2). In particular, mouse L4/5 IT 1 and 2 transcriptomic clusters together corresponded to a set of 5-7 L3-5 IT clusters in human and marmoset.

We further compared the L4-like cells in mouse MOp with those from mouse primary visual cortex (VISp) ¹⁵ after co-clustering all the SMART-Seq glutamatergic neurons from both regions (**Fig. 8e**). In the UMAP representation, L4/5 IT cells in MOp occupied a subspace of the L4 IT co-cluster defined by the intersection of marker genes *Cux2* and *Rorb*, suggesting that L4-like cells in MOp are similar to a subset of L4 cells in VISp while the L4 cells in VISp have additional diversity and specificity.

L4-like IT cells in MOp also exhibited morphological features characteristic of traditionally defined L4 excitatory neurons. From the Patch-seq study ⁶⁴, cells from the L4/5 IT 1 type had no

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813 814 or minimal apical dendrites without tufts in layer 1, in contrast to cells from the neighboring L2/3 IT, L4/5 IT 2 and L5 IT types which had tufted apical dendrites (Fig. 8f). We also obtained full morphological reconstructions of excitatory neurons with their somas located in L2, L3 or L4 in MOp or the neighboring secondary motor area (MOs) from fMOST imaging of Cux2-CreERT2; Ai166 mice ^{68,80}. As shown in Fig. 8b, Cux2 is a specific marker gene for L2/3 IT and L4/5 IT 1 types. These full reconstructions allowed us to examine, in addition to dendritic morphologies, the full extent of both local and long-range axon projections. The MOp/MOs neurons with somas in putative L4 (between L2/3 and L5) exhibited two local morphological features characteristic of L4 neurons found in sensory cortical areas (Fig. 8g). First, the dendrites of the L4 neurons were simple and untufted whereas those of the L2 and L3 neurons all had extensive tufts. Second, the local axons of L4 neurons mostly projected upward into L2/3 in addition to collateral projections; on the contrary, the local axons of L2 and L3 neurons mostly projected downward, reaching into L5. These local projection patterns are consistent with the canonical feedforward projections within a cortical column observed in somatosensory and visual cortices, with the first feedforward step being from L4 to L2/3 and the second feedforward step from L2/3 to L5 ⁹⁶. We also found that the MOp/MOs L4 neurons had intracortical longrange projections like the L2 and L3 neurons (Fig. 6d).

Taken together, our multimodal characterization demonstrates that mouse MOp indeed has excitatory neurons with L4 characteristics, namely, occupying a specific layer between L2/3 and L5, having simple and untufted dendrites and upward-projecting local axons, belonging to a transcriptomic type (L4/5 IT_1) marked by a L4-specific gene *Rspo1* as well as the intersection of a L2/3/4-specific gene *Cux2* and a L4/5-specific gene *Rorb*, and having corresponding epigenomic regulatory elements. L4-like neurons may also exist in human and marmoset M1.

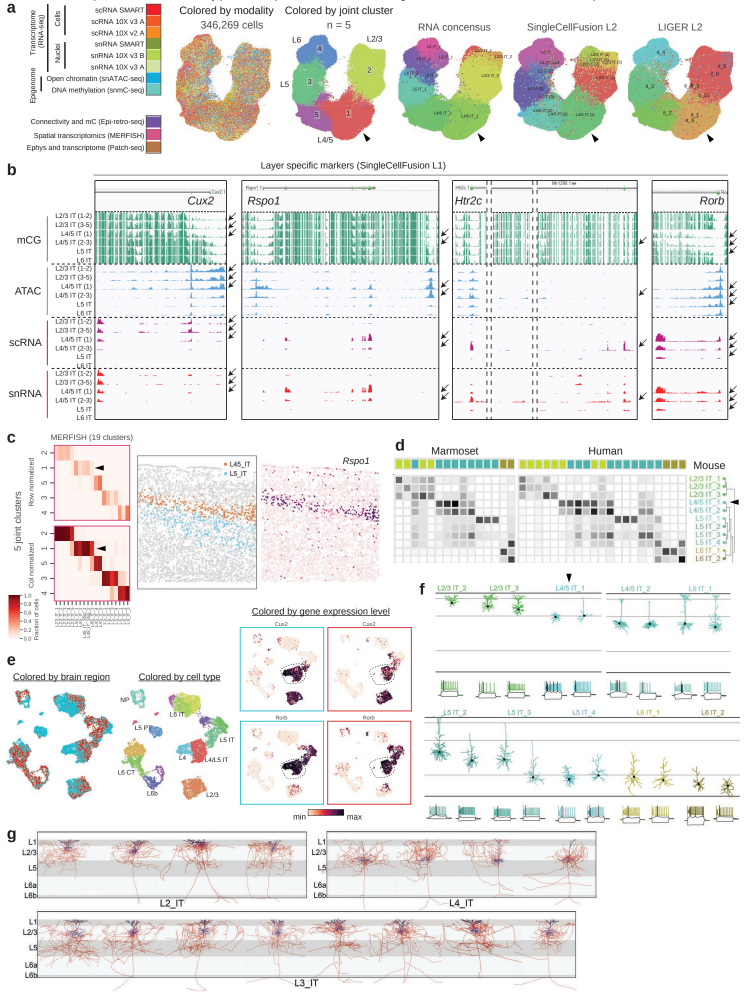


Figure 8. Existence of L4 excitatory neurons in MOp. a, UMAP embedding of IT cells from 11 datasets. Cells are colored by modalities, by cluster identities from the 11-dataset joint clustering, and by cluster identities generated from other consensus clustering methods in ⁴⁵. **b**, Genome browser of layer-specific gene markers - from L2/3 to L5 - across IT cell types as defined in ⁴⁵ (SingleCellFusion L1). Arrows indicate cell types with correlated transcription and epigenomic signatures of the specific marker gene. c, MERFISH IT clusters correspond well with the above joint clusters from a (confusion matrices, left panel) and reveal a L4 specific cluster (L45 IT) distinctly separated from the L5 IT cluster (middle panel) and marked by gene Rspo1 (right panel). d, Correspondence between mouse and human or marmoset transcriptomic IT types, e, UMAP embedding of excitatory cells from MOp (scRNA SMART) 45 and VISp 15 show that L4 excitatory cells from MOp correspond to a subset of L4 excitatory cells from VISp. Cells are colored by brain regions (MOp, red; VISp, blue), by cell types, and by expression levels (log10(TPM+1)) of marker genes Cux2 and Rorb. f, Dendritic morphologies and spiking patterns of representative Patch-seq cells from all IT types (L2/3 to L6). Arrow heads in a, c, d and f indicate the L4/5 IT 1 type. g, Local dendritic and axonal morphologies of fully reconstructed IT neurons with somas located in L2, L3 and L4. Black, apical dendrites. Blue, basal dendrites. Red, axons.

Integrated multimodal characterization of two L5 ET projection neuron types in MOp Previous studies had shown that in the mouse anterolateral motor (ALM) cortex, part of MOs, L5 ET neurons have two transcriptomically distinct projection types that may carry out different motor-control functions; the thalamus projecting type may be involved in movement planning whereas the medulla (MY) projecting type may be involved in initiation of the movement ^{15,86}. Here as the second case study, through integrated multimodal characterization we demonstrate that L5 ET neurons in MOp can also be divided into MY-projecting and non-MY-projecting types.

As shown in the companion paper ⁴⁵, we mapped the mouse MOp L5 ET transcriptomic types to the previous VISp-ALM transcriptomic taxonomy ¹⁵. From this mapping we found that the MOp L5 ET 1 type corresponded to the ALM MY-projecting type marked by *Slco2a1*, whereas MOp L5 ET 2-4 types corresponded to the ALM thalamus-projecting types with L5 ET 2/3 marked by *Hpgd* and L5 ET 4 by *Npsr1*. Here we show such distinction is consistent across all molecular datasets (**Fig. 9a-b**). Mouse transcriptomic type L5 ET 1 corresponded well with both integrated molecular type SCF L5 ET (1) and MERFISH clusters L5_ET_5, as well as with a L5 ET transcriptomic type from human and marmoset. Mouse transcriptomic types L5 ET 2-4 corresponded with integrated molecular types SCF L5 ET (2-3), MERFISH clusters L5_ET_1-4, and two L5 ET transcriptomic types from human and marmoset. The laminar distribution of these two groups was revealed by MERFISH, with cells in L5_ET_1-4 clusters intermingled in the upper part of L5 and cells in L5_ET_5 located distinctly in lower L5 (**Fig. 9c**). The two

groups were further distinguished by epigenomic peaks associated with specific marker genes,

Slco2a1 for SCF L5 ET (1) type and Npnt for SCF L5 ET (2-3) types (**Fig. 9d**), providing

validity to the two novel transgenic driver lines we generated, Slco2a1-P2A-Cre and Npnt-P2A
FlpO (**Fig. 7**).

Epi-Retro-Seq study (see above) revealed more complex long-range projection patterns among the 6 epigenetic L5-ET clusters identified, with cluster 0 predominantly projecting to MY while other clusters having variable and less specific projection patterns (clusters 2 and 3 also containing MY-projecting cells) (**Fig. 5g**). We co-clustered the L5 ET cells from the Epi-Retro-Seq data and the snRNA-seq 10x v3 B data ⁴⁵ to investigate the correspondence of Epi-Retro-Seq clusters and projection targets with transcriptomic clusters (**Fig. 9e**). We found that the consensus transcriptomic cluster L5 ET 1 corresponds to Epi-Retro-Seq clusters 0, 2 and 3, all of which contain MY-projecting neurons. On the other hand, transcriptomic clusters L5 ET 2-4 correspond to Epi-Retro-Seq clusters 1, 4 and 5, which do not contain MY-projecting neurons. Thus, all MY-projecting neurons are mapped to transcriptomic type L5 ET 1, while neurons in the L5 ET 2-4 types do not project to MY.

Anterograde tracing in *Slco2a1-P2A-Cre* mice demonstrated predominant projection from *Slco2a1*-labeled neurons in MOp to MY (**Fig. 7**). We identified multiple full morphology reconstructions of MOp L5 ET neurons from fMOST imaging of *Fezf2-CreER;Ai166* and *Pvalb-T2A-CreERT2;Ai166* transgenic mice ⁶⁸. These reconstructions could be clearly separated into a MY-projecting group and a non-MY-projecting group (**Fig. 9g**), though they were not directly linked to transcriptomic types yet. Both groups of cells had thick-tufted dendrites that were similar to each other (**Fig. 9g**). Consistent with this, Patch-seq cells corresponding to transcriptomic types L5 ET 1-4 also were indistinguishable from each other by their dendritic morphologies (**Fig. 9f**).

Altogether, our integrated multimodal characterization identified two major types of mouse L5 ET projection neurons, MY-projecting and non-MY-projecting, with distinct gene markers, epigenomic elements, laminar distribution, and corresponding types in human and marmoset.

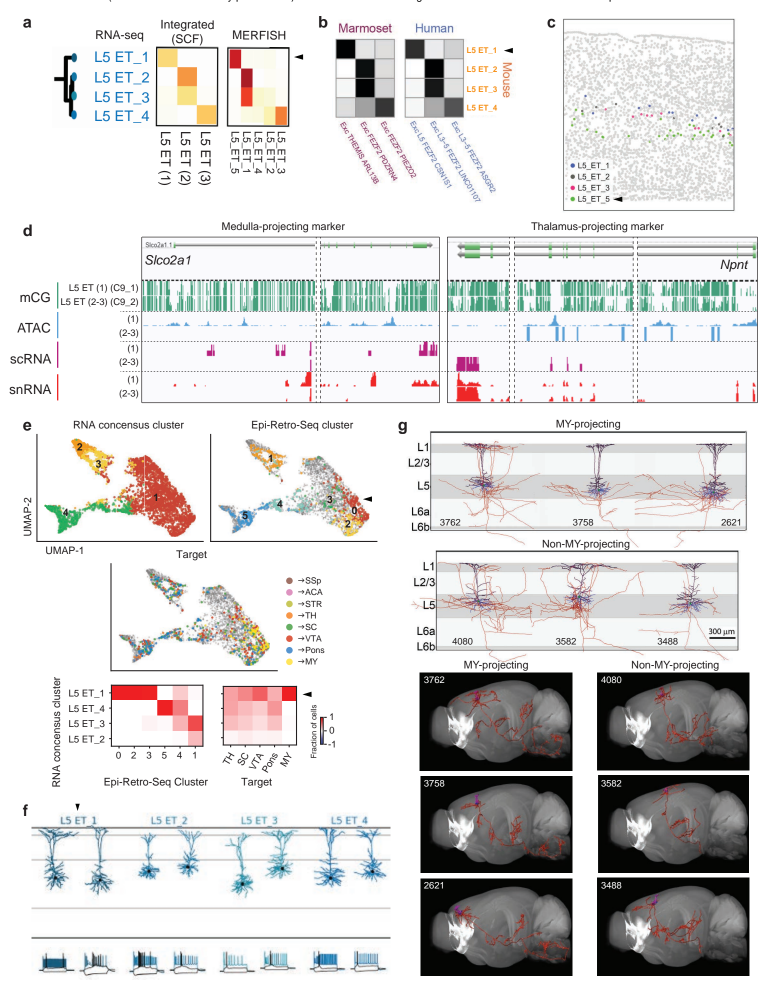


Figure 9. Two distinct L5 ET projection neuron types in MOp. a, Within the mouse L5 ET subclass, good correspondence is observed between the 4 transcriptomic clusters and the 3 integrated molecular clusters (SingleCellFusion) or the 5 MERFISH clusters. b, Within the L5 ET subclass, the 4 mouse transcriptomic clusters correspond well with the 3 transcriptomic clusters in either human or marmoset. c, In MERFISH, cells belonging to the L5 ET 1-4 clusters co-occupy the upper L5, whereas L5 ET 5 cells are distinctly located in lower L5. d, Genome browser of gene markers between the MY-projecting (Slco2a1) and the non-MYprojecting (Npnt) L5 ET neurons. e, Integration panels between L5 ET Epi-Retro-Seq clusters and consensus transcriptomic clusters. The transcriptomic dataset used here is snRNA 10x v3 B, which has the largest number of L5 ET cells (>4k). Top panels, UMAP plots colored by consensus transcriptomic clusters, Epi-Retro-Seq clusters and projection targets (retrograde tracer injection sites). Bottom panels, confusion matrices between consensus transcriptomic clusters and Epi-Retro-Seq clusters or major projection targets. The heatmaps are column-wise normalized rather than row-wise to avoid misleading interpretation, since the number of cells sampled from each projection may differ a lot in Epi-Retro-Seq. f, Dendritic morphologies and spiking patterns of representative Patch-seq cells corresponding to the 4 mouse transcriptomic L5 ET types, g. Local dendritic and axonal morphologies (upper panels) and brain-wide axon projections (lower panels) of representative fully-reconstructed L5 ET neurons, separated into MY-projecting and non-MY-projecting types. Black, apical dendrites. Blue, basal dendrites. Red, axons.

An integrated synthesis of multimodal features of cell types in the primary motor cortex

As the conclusion of this series of studies from BICCN, we present an overview and integrated synthesis of the knowledge gained in constructing a multimodal census and atlas of cell types in the primary motor cortex of mouse, marmoset and human (**Fig. 10**). A critical aspect of our studies is that this synthesis is only made possible by the systematic integrative computational analyses across multiple transcriptomic and epigenomic data types that connect a diverse range of cellular features together at cell subclass or type level to allow mutual correlation.

This integrated synthesis uses the mouse MOp consensus transcriptomic taxonomy (containing 18 subclasses and 116 clusters/types) ⁴⁵ as the anchor (**Fig. 10**) because it was derived from the largest and deepest datasets and was the reference taxonomy for nearly all the cross-modality and cross-species comparisons. Correspondence matrices between the different molecular modalities show that the mouse MERFISH-based spatial transcriptomic taxonomy (95 clusters) ⁵⁴, the integrated mouse molecular taxonomies combining transcriptomic and epigenomic data using either SingleCellFusion (SCF, 56 neuronal clusters) or LIGER (71 clusters) approach ⁴⁵, and the human and marmoset transcriptomic taxonomies (127 and 94 clusters, respectively) ⁴⁸ all aligned largely consistently with the mouse consensus transcriptomic taxonomy. Such alignment convincingly demonstrates that cell types in a given brain region can be consistently described

by different types of characterization. At the same time, it should also be noted that the alignments are not perfect and disagreements do exist at the individual cluster level (which is most pronounced in cross-species comparisons), suggesting that differential variations exist in different data types and consistency, in particular that across species, may be more appropriately described at an intermediate level of granularity.

In this integrated synthesis, we can further assign additional attributes to the molecularly defined cell types (**Fig. 10**). Based on Patch-seq ⁶⁴, Retro-seq (e.g. Epi-Retro-Seq ⁷⁹), Retro-MERFISH ⁵⁴, and axon projection ^{68,69} studies, we relate many transcriptomic neuronal types or subclasses to cortical neuron types traditionally defined by electrophysiological, morphological and connectional properties, thus bridging our cell type census with historical and community knowledge. We provide the relative proportion of each cell type within the mouse MOp using either snRNA-seq or MERFISH data. The MERFISH data also identify the spatial distribution pattern of each cell type ⁵⁴. For example, we found that excitatory or inhibitory neuron types are distributed along the cortical depth, with many individual types adopting narrow cortical-depth distributions, often occupying predominantly a single layer or a sublayer, and related types (e.g. the L2/3-6 IT excitatory types) can display a gradual transitioning across cortical depths/layers. On the other hand, non-neuronal cell types are either distributed across all layers or specific to layer 1 or the white matter (WM). Patch-seq data also provided the cortical depth positions of a variety of neuronal cell types.

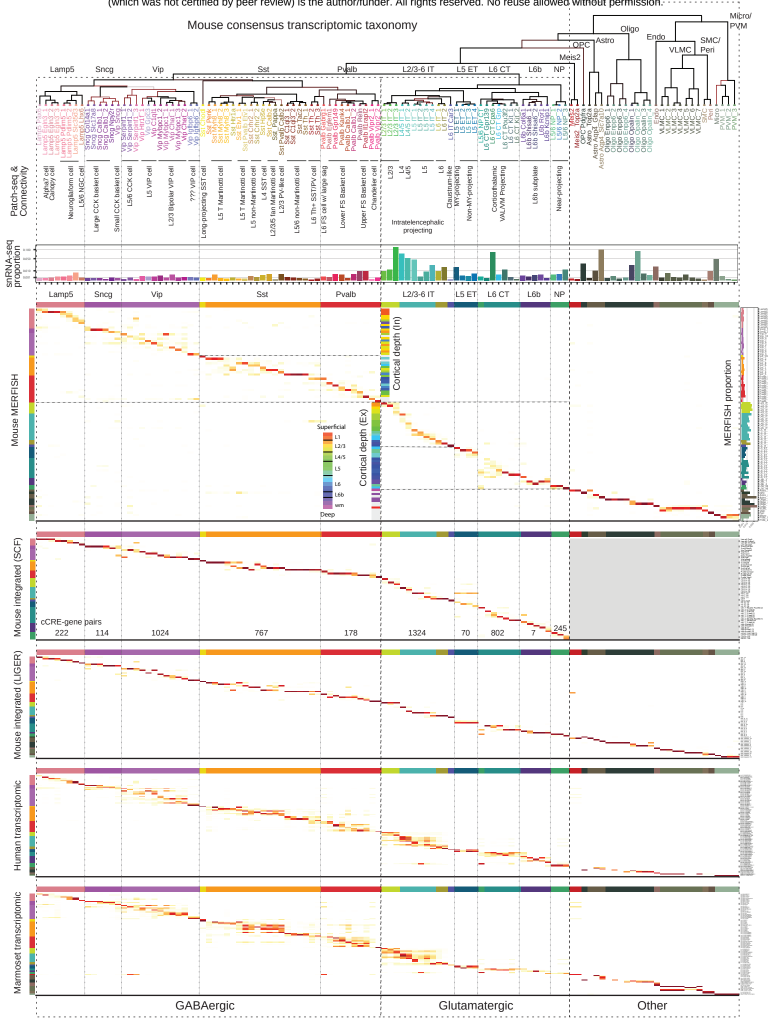
Finally, we demonstrate the possibility to elucidate gene regulatory mechanisms by discovering candidate cis-regulatory elements (cCREs) as well as master transcription factors (TFs) specific to neuronal subclasses by mining the combined mouse MOp transcriptomic and epigenomic datasets to access both RNA expression of genes and accessibility or DNA methylation of cCREs from the same cell clusters.

For example, we found 7,245 distal cCRE (>1 kbp from transcriptional start site)-gene pairs in neuronal cells in MOp that showed positive correlation between accessibility at the 6,280 cCREs and expression levels of 2,490 putative target genes (see Methods, and companion papers ^{45,52}). We grouped these putative enhancers into modules based on accessibility across cell clusters (Extended Data Fig. 2). 76% of putative enhancers showed remarkable sub-type specific chromatin accessibility and were enriched for lineage-specific transcription factors, while 24% of putative enhancers (1,527) were widely accessible and linked to genes expressed across neuronal cell clusters with highest expression levels in Glutamatergic neurons (module M1, Extended Data Fig. 2b). Putative enhancers in this module showed enrichment of sequence motifs recognized by transcription factors CTCF, MEF2 indicating a more general rule of these factors in establishing neuronal gene regulatory programs (Extended Data Fig. 2c). Meanwhile, other modules (M2 to M14) of enhancer-gene pairs were active in a subclass-specific manner (Extended Data Fig. 2b-d). Thus, using this approach we have identified a large number of

 enhancer-gene pairs for each subclass of neurons (Fig. 10). These enhancers can be potentially used to generate cell type-targeting viral tools.

Similarly, we identified transcription factors showing cell-type specificity supported by both RNA expression and DNA binding motif enrichment in hypo-CG-DMR of MOp subclasses (see Methods, and companion papers ^{45,50}) (**Extended Data Fig. 3**). Combining these two orthologous pieces of evidence identified many well-studied TFs in embryonic precursors, such as the Dlx family members for pan-inhibitory neurons, and Lhx6 and Mafb for MGE derived inhibitory neurons. We further identified many additional TFs with more restricted patterns in specific subclasses, such as Rfx3 and Rreb1 (in L2/3 IT), Atoh7 and Rorb (in L4/5 IT), Pou3 family members (in L5 ET), Etv1 (in L5/6 NP), Esrr family members (in Pvalb), and Arid5a (in Lamp5). The agreement of these two modalities suggests a requirement of TF regulatory activity in mature neurons to maintain aspects of cell phenotypes and identity.

In summary, our comprehensive multimodal characterization of cell types from the MOp region demonstrates that transcriptomic, epigenomic, spatial, physiological, morphological and connectional properties can be all correlated and integrated together, to reveal organizational principles of brain cell types and bridge molecular, structural and functional studies in different modalities and across species.



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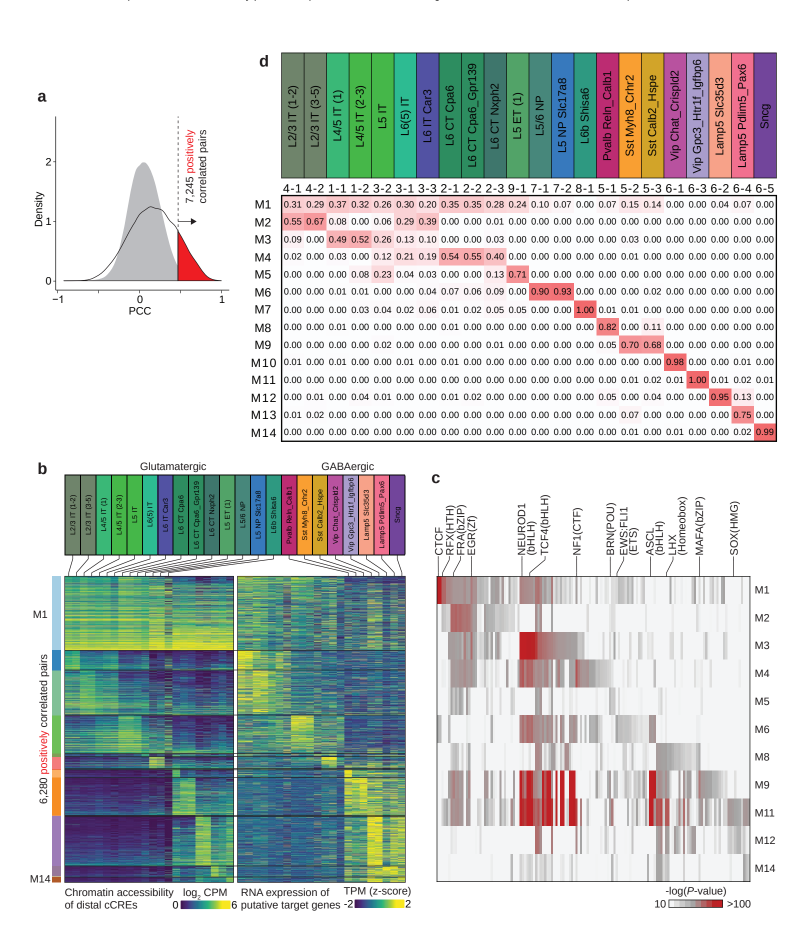
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Figure 10. An integrated multimodal census and atlas of cell types in the primary motor cortex of mouse, marmoset and human. The mouse MOp consensus transcriptomic taxonomy at the top is used to anchor cell type features in all the other modalities. Subclass labels are shown above major branches and cluster labels are shown below each leaf node. Confusion matrices show the correspondence between the mouse MOp transcriptomic taxonomy with those derived from other molecular datasets, including mouse MERFISH, the integrated mouse molecular taxonomies by SingleCellFusion (SCF) or LIGER, and the human and marmoset transcriptomic taxonomies. Using Patch-seq and connectivity studies, many transcriptomic neuronal types or subclasses are annotated and correlated with known cortical neuron types traditionally defined by electrophysiological, morphological and connectional properties. (Note: no Patch-seq data were collected for the Vip cells labeled by question marks.) The relative proportions of all cell types within the mouse MOp are calculated from either the snRNA-seq 10x v3 B (horizontal bar graph) or MERFISH (vertical bar graph to the right of the MERFISH matrix) dataset. Median cortical depth position of each cell type derived from MERFISH is shown as color-coded bar graphs at the center of the MERFISH matrix, colored according to the rainbow scheme from superficial (red) to deep (purple) layers as shown on the left. Cell types with dispersed distributions spanning relatively large ranges of cortical depth are colored in grey. The numbers of cCRE-gene pairs in modules corresponding to neuronal subclasses identified by Cicero from the scRNA-seg and snATAC-seg datasets are shown at the bottom of the SCF matrix.



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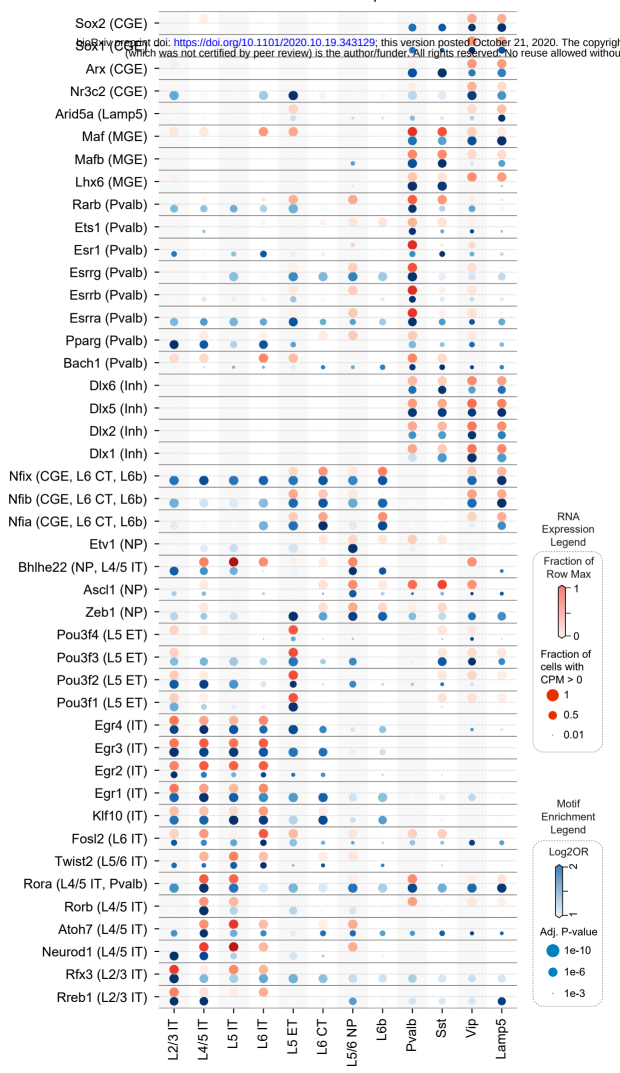
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Extended Data Figure 2. Characterization of putative enhancer-gene pairs. a, Detection of putative enhancer-gene pairs. 7,245 pairs of positively correlated cCRE and genes (highlighted in red) were identified using an empirically defined significance threshold of FDR<0.01. Grey filled curve shows the distribution of PCC for randomly shuffled cCRE-gene pairs. b, Heatmap of chromatin accessibility of 6,280 putative enhancers across joint cell clusters (left) and expression of 2,490 target genes (right). Note genes are displayed for each putative enhancer separately. CPM: counts per million, TPM: transcripts per million. c, Enrichment of known transcription factor motifs in distinct enhancer-gene modules. Displayed are known motifs from HOMER with enrichment -log p-value >5. In module M1, de novo motif analysis of putative enhancers in this module showed enrichment of sequence motif recognized by transcription factors CTCF, MEF2. CTCF is a widely expressed DNA binding protein with a well-established role in transcriptional insulation and chromatin organization, but recently it was also reported that CTCF can promote neurogenesis by binding to promoters and enhancers of related genes. In the L2/3 IT selective module M2, the putative enhancers were enriched for the binding motif for Zinc-finger transcription factor EGR, a known master transcriptional regulator of excitatory neurons ⁹⁷. In the Pvalb selective module M8, the putative enhancers were enriched for sequence motifs recognized by the MADS factor MEF2, which is associated with regulating cortical inhibitory and excitatory synapses and behaviors relevant to neurodevelopmental disorders 98. d. Heatmap shows the weights of each joint cell cluster in each module, which were derived from the coefficient matrix. The values of each column are scaled (0-1).

Extended Data Figure 3. Dot plot illustrating the RNA expression levels (red) and hypo-CG-DMR motif enrichments (blue) of transcription factors (TFs) in mouse MOp subclasses. The size and color of red dots indicate the proportion of expressing cells and the average expression level in each subclass, respectively. The size and color of blue dots indicate adjusted P-value and log2(Odds Ratio) of motif enrichment analysis, respectively.

Transcription Factors RNA Expression And Motif Enrichment in MOp Subclasses



DISCUSSION

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A cell census and atlas of motor cortex

Understanding the principles of brain circuit organization requires a detailed understanding of its basic components, but the cellular diversity and complexity of the brain, including the neocortex, have defied a comprehensive and quantitative description. Single-cell transcriptomics and epigenomics, as well as spatially resolved single-cell transcriptomics, are accelerating efforts to classify all molecular cell types in many organ systems ^{40,99}, including the brain ^{5,6,100}. The current effort combines these technologies to derive a robust and comprehensive molecular cell type classification and census of the primary motor cortex of mouse, marmoset and human, coupled with a spatial atlas of cell types and an anatomical input/output wiring diagram in mouse. We demonstrate the robustness and validity of this classification through strong correlations across cellular phenotypes, and strong conservation across species. Together these data comprise a cell atlas of primary motor cortex that encompasses a comprehensive reference catalog of cell types, their proportions, spatial distributions and anatomical and physiological characteristics, and molecular genetic profiles, registered into a Common Coordinate Framework ⁴¹. This cell atlas establishes a foundation for an integrative study of the architecture and function of cortical circuits akin to reference genomes for studying gene function and genome regulatory architecture. Furthermore, it provides a comprehensive map of the genes that contribute to cellular phenotypes and their epigenetic regulation. These data resources and associated tools enabling genetic access for manipulative experimentation are publicly available and provide a roadmap for exploring cellular diversity and organization across brain regions, organ systems, and species.

The molecular classification presented here is overall consistent with prior literature and synthesizes a wide body of existing and new information into a coherent quantitative framework that provides metrics for the robustness of, and the similarities and distinctions between, cell types. For motor cortex, as for other cortical regions ^{15,18}, this cellular organization is hierarchical, with different branches comprising major cell classes, subclasses, and types representing the finest resolution clusters afforded by each method. This classification provides strong evidence for the existence of hitherto poorly studied but molecularly distinct subclasses such as the near-projecting (NP) pyramidal neurons, and many more novel cell types. At the level of cell class and subclass (and some highly distinctive types like chandelier cells and longrange projecting Sst Chodl interneurons), we find remarkable concordance across transcriptomics, epigenomics, spatial patterning, physiology and connectivity, as well as strong homology across species. The class and subclass branches clearly represent different developmental programs, such as GABAergic neuron derivatives of different zones of the ganglionic eminences 101,102 or the layer-selective glutamatergic neurons derived sequentially from progenitors of the cortical plate, and the hierarchical organization generates new hypotheses about developmental origins of highly distinctive cell types. This quantitative

hierarchy also challenges well-established nomenclature systems. For example, the term glia is typically used to encapsulate astrocytes, oligodendrocytes and OPCs, and microglia. However, microglia are not closely related to these neuroectoderm-derived populations based on transcriptomics or developmental origins ¹⁰³ and should be grouped with other more similar non-neuronal cell types such as endothelial cells, VLMCs and pericytes. Substantial challenges remain for redefining data-driven cell ontologies and nomenclature systems ^{100,104}.

Comparisons of the MOp results described here to other regions also help to understand what makes the motor cortex functionally distinct. Previous transcriptomic studies suggested that GABAergic interneuron types are shared among cortical regions whereas glutamatergic projection neuron types exhibit gradient-like distribution across the cortical sheet and are more distinct between distant regions but more similar between neighboring regions ^{15,44}. Thus the projection neurons in MOp are more similar to those of nearby regions, yet our anatomical tracing study defines a MOp-specific input-output wiring diagram. This result suggests that differential axonal projections of similar molecular types among different cortical areas may be the major feature defining regional functional specificity. We also find substantial variation in the proportion of specific cell types between cortical areas. For example, we identify two glutamatergic neuron types that distinguish MOp from its neighboring primary somatosensory (SSp) region, the L4 IT neurons that are present in MOp at lower abundance level than in SSp and the *Slco2a1*-expressing, medulla-projecting L5 ET neurons that are more abundant in MOp than in SSp ^{54,68}. These regional differences in cellular makeup may contribute to the functional specialization of MOp as well.

Cell type discreteness, variation and phenotypic concordance

The concordance of transcriptomic and epigenomic results and their overall correlation with other cellular phenotypes, including spatial distributions, morphological properties, electrophysiological properties, and projection/connectivity, strongly argues for a unifying molecular genetic framework for understanding cortical cell types, particularly at the level of subclasses and distinctive cell types. At the same time, substantial multimodal variations at finer granularity appear to preclude a fully discretized representation of cell types with consistency across all cellular phenotypes. One source of variation is differences in granularity with different molecular data modalities, with transcriptomics providing the highest granularity at present. This may reflect true biology or differences in technological information content, for example sparse genome coverage in epigenetic methods. A second source involves continuous rather than discrete variation. For example, while some highly granular cell types are highly distinct from others (e.g. L6 IT Car3, Sst Chodl and Pvalb chandelier cells), many other types exhibit continuous variation in their properties both within types and among closely related types with no clear boundaries between them. However, even at this fine-grained level of continuous variation, spatial, morphological and physiological properties often co-vary with transcriptomic profiles as shown by MERFISH and Patch-seq. Similar findings on continuous as well as unitary

variations have been reported for hippocampal interneurons ¹⁶. These results suggest that continuous phenotypic variation may represent a general organizing principle underlying the diversification of brain cell types.

As shown in our mouse Epi-Retro-Seq, MERFISH, and single-neuron full morphology and projection studies there is a strong correlation between molecular phenotype and axonal target specificity at the subclass level (e.g., IT, L5 ET, L6 CT). This was also the case for medulla-projecting L5 ET type. However, a strict correlation between molecular cell types and specific axonal projection targets was not generally observed. It is possible that axon pathfinding during development involves stochastic decisions and subsequent activity-dependent pruning that mature cell transcriptomes do not represent. Furthermore, individual projection neurons typically have collaterals to many different target regions which complicates understanding these relationships. Comprehensive datasets on the complete axonal projections of individual neurons whose molecular identity is clearly established will be needed to address this issue.

Cell type conservation and divergence

Evolutionary conservation is strong evidence of functional significance. The demonstrated conservation of cell types from mouse, marmoset, macaque and human strongly suggests that these conserved types play important roles in cortical circuitry and contribute to a common blueprint essential for cortical function in mammals and even more distantly related species. We also find that similarity of cell types varies as a function of evolutionary distance, with substantial species differences that either represent adaptive specializations or genetic drift. For the most part species specializations tend to appear at the finer branches or leaves of the hierarchical taxonomy. This result is consistent with a recent hypothesis in which cell types are defined by common evolutionary descent and evolve independently, such that new cell types are generally derived from existing genetic programs and appear as specializations at the finer levels of the taxonomic tree ¹⁰⁵.

A surprising finding across all homologous cell types was the relatively high degree of divergence for genes with highly cell type-specific expression in a given species. This observation provides a clear path to identify the core conserved genes underlying the canonical identity and features of those cell types. Furthermore, it highlights the need to understand species adaptations superimposed on the conserved program, as many specific cellular phenotypes may vary across species including gene expression, epigenetic regulation, morphology and connectivity, and physiological functional properties. As we illustrate in the Betz cells, there is clear homology across species in the layer 5 ET subclass, but variation in many measurable properties across species.

A framework for linking model organisms to human biology and disease

The results presented have major utility and implications for the consideration of model organisms to understand human brain function and disease. Despite major investments, animal models of neuropsychiatric disorders have often been characterized by "loss of translation," fueling heated debates about the utility of model organisms in the search for therapeutic targets for treating human diseases. The molecular genetic framework of cell type organization established by the current study will provide a robust cellular metric system for cross-species translation of knowledge and insight that bridges levels of organization based on their inherent biological and evolutionary relationships. For example, the characterization of cell types and their properties shown in Figure 10 can be used to infer the main characteristics of homologous cell types in humans and other mammalian species, despite the often extreme difficulty in measuring their specific properties in those species. On the other hand, they also reveal the potential limitations of model organisms and the necessity to study human and closely related primate species to understand the specific features of cell types as they contribute to human brain function and susceptibility to human-specific diseases. Having cell census information aligned across species as illustrated here should be highly valuable for making rational choices about the best models for each disease and therapeutic target. This reductionist dissection of the cellular components provides a foundation for understanding the general principles of neural circuit organization and computation that underlie mental activities and brain disorders.

Future directions

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The success of the current strategy to systematically and comprehensively dissect cell types and generate a cell census and atlas opens up numerous avenues for future work. This census and atlas form the foundation for the larger community to study specific features of cell types and aggregate information about cell types across species much as genomic databases aggregate information about genes. Classification of cell types and description of their molecular, spatial, and connectional properties in the adult sets the stage for developmental studies to understand the molecular genetic programs underlying cell type specification, maturation and circuit connectivity. The molecular classification and the utility of combined single cell transcriptomics and epigenomics to identify functional enhancers promises to deliver tools for genetic access to the great majority of brain cell types via transgenic and viral strategies. A combination of some of the approaches, such as imaging-based single-cell transcriptomics, with behavior stimulation and functional imaging can further elucidate the functional roles of distinct cell types in circuit computation. This systematic, multi-modal strategy described here is extensible to the whole brain, and major efforts are underway in the BICCN to generate a brain-wide cell census and atlas in the mouse with increasing coverage of human and non-human primates.

METHODS

Integrating 10x v3 snRNA-seq datasets across species

- To identify homologous cell types across species, human, marmoset, and mouse 10x v3 snRNA-
- seq datasets were integrated using Seurat's SCTransform workflow. Each major cell class
- 1200 (glutamatergic, GABAergic, and non-neuronal cells) was integrated separately across species.
- Expression matrices were reduced to 14,870 one-to-one orthologs across the three species (NCBI
- Homologene, 11/22/2019). Nuclei were downsampled to have approximately equivalent
- numbers at the subclass level across species. Marker genes were identified for each species'
- 1204 cluster and used as input to guide alignment and anchor-finding during integration steps. For full
- 1205 methods see 48 .

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Estimation of cell type homology

- To establish a robust cross-species cell type taxonomy, we applied a tree-based clustering
- method on integrated class-level datasets (https://github.com/AllenInstitute/BICCN M1 Evo).
- 1210 The integrated space (from the above mentioned Seurat integration) was over-clustering into
- small sets of highly similar nuclei for each class (~500 clusters per class). Clusters were
- aggregated into metacells, then hierarchical clustering was performed based on the metacell gene
- 1213 expression matrix using Ward's method. Hierarchical trees were then assessed for cluster size,
- species mixing, and branch stability by subsampling the dataset 100 times with 95% of nuclei.
- 1215 Finally, we recursively searched every node of the tree, and if certain heuristic criteria were not
- sufficed for a node below the upper node, all nodes below the upper node were pruned and nuclei
- belonging to this subtree were merged into one homologous group. We identified 24
- 1218 GABAergic, 13 glutamatergic, and 8 non-neuronal cross-species consensus clusters that were
- highly mixed across species and robust. For full methods see ⁴⁸.

Cross-species differential gene expression and correlations

- 1222 Expression matrices for each species, for each major cell class (GABAergic, glutamatergic, and
- 1223 non-neuronal cells) were normalized using Seurat's SCTransform function with default
- parameters to generate a 'corrected UMI' matrix and remove technical variation within each
- species. SCTransform normalized counts matrices were then counts per 100,000 UMI (CP100K)
- normalized to account for variable sequencing depths between species. CP100K normalization
- was performed by multiplying each value in the 'corrected UMI' (SCTransform normalized)
- matrix by 100,000 and dividing by the column sums (total UMIs from each nuclei).
- 1229 SCTransform-CP100K normalized matrices were then used to find DE genes and correlations
- between species for each cross-species cluster.
- DE gene analysis was performed with Seurat's FindAllMarkers function, using the Wilcoxon
- rank sum test, between each pair of species for a given cross-species cluster (e.g. human
- Lamp5 1 vs. marmoset Lamp5 1, human Lamp5 1 vs. mouse Lamp5 1, and marmoset
- Lamp5 1 vs. mouse Lamp5 1). Marker genes (FDR < 0.01, log fold-change > 2, expressed in at
- least 10% of nuclei) from each pairwise species comparison were identified for each cross-
- species cluster. We report the sum of marker genes between each species comparison as a

heatmap in Figure 2e and show that human and marmoset have fewer DE genes between each other than with mouse across all cross-species clusters.

To visualize the correspondence of a given cross-species cluster between each pair of species, we first found the average SCTransform-CP100K expression for each cross-species cluster for each species. Average expression was then log-transformed and the spearman correlations between each species pair were identified and reported in the Figure 2d heatmap, which shows human and marmoset have higher correlations than either primate with mouse for all clusters except Endo, VLMC, and Microglia/PVM clusters (likely due to differences in sampling).

Integrating mouse transcriptomic, spatially resolved transcriptomic, and epigenomic datasets To integrate IT cell types from different mouse datasets, we first take all cells that are labeled as IT, except for L6_IT_Car3, from the 11 datasets as listed in Figure 8a. These cell labels are either from dataset-specific analyses ^{54,64,79}, or from the integrated clustering of multiple datasets ⁴⁵. The integrated clustering and embedding of the 11 datasets are then generated by projecting all datasets into the 10x v2 scRNA-seq dataset using SingleCellFusion ^{45,59}. Genome browser views of IT and ET cell types (Figure 8b and Figure 9d) are taken from the corresponding cell types of the brainome portal (brainome.ucsd.edu/BICCN MOp) ⁴⁵.

Integration of L5 ET cells from Epi-Retro-Seq and 10x snRNA-Seq

For snRNA-Seq, the 4,515 cells from 10x v3 B dataset labeled as L5 ET by SCF were selected ⁴⁵. The read counts were normalized by the total read counts per cell and log transformed. Top 5,000 highly variable genes were identified with Scanpy ¹⁰⁶ and z-score scaled across all the cells. For Epi-Retro-Seq, the posterior methylation levels of 12,261 genes in the 848 L5 ET cells were computed ⁷⁹. Top 5,000 highly variable genes were identified with AllCools ⁵⁹ and z-score scaled across all the cells. The 1,512 genes as the intersection between the two highly variable gene lists were used in Scanorama ¹⁰⁷ to integrate the z-scored expression matrix and minus z-scored methylation matrix with sigma equal to 100.

Identification of candidate cis-regulatory elements

For peak calling in the snATAC-seq data, we extracted all the fragments for each cluster, and then performed peak calling on each aggregate profile using MACS2 ¹⁰⁸ with parameter: "--nomodel --shift –100 --ext 200 --qval 1e-2 –B --SPMR". First, we extended peak summits by 250 bp on either side to a final width of 501 bp. Then, to account for differences in performance of MACS2 based on read depth and/or number of nuclei in individual clusters, we converted MACS2 peak scores (-log10(q-value)) to "score per million" ¹⁰⁹. Next, a union peak set was obtained by applying an iterative overlap peak merging procedure, which avoids daisy-chaining and still allows for use of fixed-width peaks. Finally, we filtered peaks by choosing a "score per million" cut-off of 5 as candidate cis-regulatory elements (cCREs) for downstream analysis.

Predicting enhancer-promoter interactions

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First, co-accessible cCREs are identified for all open regions in all neurons types (cell clusters with less than 100 nuclei from snATAC-seq are excluded), using Cicero 110 with following parameters: aggregation k = 50, window size = 500 kb, distance constraint = 250 kb. In order to find an optimal co-accessibility threshold, we generated a random shuffled cCRE-by-cell matrix as background and calculated co-accessible scores from this shuffled matrix. We fitted the distribution of co-accessibility scores from random shuffled background into a normal distribution model by using R package fitdistrplus ¹¹¹. Next, we tested every co-accessible cCRE pair and set the cut-off at co-accessibility score with an empirically defined significance threshold of FDR<0.01. The cCREs outside of \pm 1 kb of transcriptional start sites (TSS) in GENCODE mm10 (v16) were considered distal. Next, we assigned co-accessibility pairs to three groups: proximal-to-proximal, distal-to-distal, and distal-to-proximal. In this study, we focus only on distal-to-proximal pairs. We calculated the Pearson's correlation coefficient (PCC) between gene expression (scRNA SMART-seq) and cCRE accessibility across the joint clusters to examine the relationships between the distal cCREs and target genes as predicted by the coaccessibility pairs. To do so, we first aggregated all nuclei/cells from scRNA-seq and snATACseq for every joint cluster to calculate accessibility scores (log2 CPM) and relative expression levels (log2 TPM). Then, PCC was calculated for every gene-cCRE pair within a 1 Mbp window centered on the TSS for every gene. We also generated a set of background pairs by randomly selecting regions from different chromosomes and shuffling of cluster labels. Finally, we fit a normal distribution model and defined a cut-off at PCC score with an empirically defined significance threshold of FDR<0.01, in order to select significant positively correlated cCREgene pairs.

Identification of cis-regulatory modules

We used Nonnegative Matrix Factorization (NMF) to group cCREs into cis-regulatory modules based on their relative accessibility across cell clusters. We adapted NMF (Python package: sklearn) to decompose the cell-by-cCRE matrix V (N×M, N rows: cCRE, M columns: cell clusters) into a coefficient matrix H (R×M, R rows: number of modules) and a basis matrix W (N×R), with a given rank R:

V≈WH.

The basis matrix defines module related accessible cCREs, and the coefficient matrix defines the cell cluster components and their weights in each module. The key issue to decompose the occupancy profile matrix was to find a reasonable value for the rank R (i.e., the number of modules). Several criteria have been proposed to decide whether a given rank R decomposes the occupancy profile matrix into meaningful clusters. Here we applied a measurement called "Sparseness" ¹¹² to evaluate the clustering result. Median values were calculated from 100 times for NMF runs at each given rank with a random seed, which will ensure the measurements are stable. Next, we used the coefficient matrix to associate modules with distinct cell clusters. In the coefficient matrix, each row represents a module and each column represents a cell cluster. The

- values in the matrix indicate the weights of clusters in their corresponding module. The
- 1319 coefficient matrix was then scaled by column (cluster) from 0 to 1. Subsequently, we used a
- coefficient > 0.1 (~95th percentile of the whole matrix) as a threshold to associate a cluster with
- a module. Similarly, we associated each module with accessible elements using the basis matrix.
- For each element and each module, we derived a basis coefficient score, which represents the
- accessible signal contributed by all clusters in the defined module.

Identification of subclass-selective TFs by both RNA expression and motif enrichment

- All analyses for this section were at the subclass level. For RNA expression, we used the sc
- 1327 SMART-seq dataset and compared each subclass with the rest of the population through a one-
- tailed Wilcoxon test and FDR correction to select significantly differentially-expressed
- transcription factors (adjusted P-value < 0.05, cluster average fold change > 2). To perform the
- motif enrichment analysis, we used known motifs from the JASPAR 2020 database ¹¹³ and the
- subclass specific hypo-CG-DMR identified in Yao et al ⁴⁵. The AME software from the MEME
- suite (v5.1.1) 114 was used to identify significant motif enrichment (adjusted P-value < 1e-3, odds
- ratio > 1.3) using default parameters and the same background region set as described in Yao et
- 1334 al 45. All genes in **Extended Data Figure 3** were both significantly expressed and had their motif
- enriched in at least one of the subclasses.

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Generation and use of transgenic mouse lines

- 1338 Npnt-P2A-FlpO and Slco2a1-P2A-Cre mouse driver lines were generated by CRISPR/Cas9-
- mediated homologous recombination (Stafford et al., BICCN companion manuscript in
- preparation). Details are provided in the Supplementary Methods.
- All experimental procedures were approved by the Institutional Animal Care and Use
- 1343 Committees (IACUC) of Cold Spring Harbor Laboratory, University of California, Berkeley and
- Allen Institute, in accordance with NIH guidelines. Mouse knock-in driver lines are being
- deposited at the Jackson Laboratory for wide distribution.

Data and code availability

Figure 1. Summary of experimental and computational approaches taken as well as community resources generated by the BICCN

All primary data available through the BICCN portal, data archives, and supporting tools.

- Brain Cell Data Center (BCDC), www.biccn.org
- Neuroscience Multi-Omics Archive (NeMO), www.nemoarchive.org
- Brain Image Library (BIL), www.brainimagelibrary.org
- Neurophysiology (DANDI), dandiarchive.org

- Allen Transcriptomics Explorer, https://portal.brain-map.org/atlases-and-data/rnaseq
- NeMO Analytics, www.nemoanalytics.org
- Morphological reconstructions, NeuroMorpho, www.neuromorpho.org

Figure 2. MOp consensus cell type taxonomy

Primary Data

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		http://data.nemoarchive.org/publication_release/M
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b-g	10x V3 macaque	criptomic/sncell/raw/
	10x V3 human (10X159-1	http://data.nemoarchive.org/biccn/lab/linnarsson/tra
	through 10x160-8)	nscriptome/sncell/10X/raw/10X159-1/
	10x V3 marmoset (bi005_m1,	http://data.nemoarchive.org/biccn/lab/feng/transcrip
	bi006_m1)	tome/sncell/raw/
	10x V3 mouse broad data (files	http://data.nemoarchive.org/biccn/grant/huang/mac
	with prefix pBICCNsMMrMOp)	osko/transcriptome/sncell/raw
Panel h:		

Intermediate analyses

	https://github.com/mukamel-lab/BICCN-Mouse-			
Panel a:	MOp/tree/master/flagship_fig2a			
Panel b:	dendrogram from companion paper (Bakken et al. 2020)			
	http://data.nemoarchive.org/brain/biccn/lab/lein/2020_M1_study_analysis/T			
Panel c:	ranscriptomics/flagship/dendrogram_barplots			
	http://data.nemoarchive.org/brain/biccn/lab/lein/2020_M1_study_analysis/T			
Panels d, e:	ranscriptomics/flagship/dendrogram_heatmaps			
Panels f, g:	from companion paper (Bakken et al. 2020)			
	http://data.nemoarchive.org/biccn/lab/lein/2020_M1_study_analysis/Transc			
	riptomics/flagship/cross species heatmap/GABAergic avg CP100K expr			
Panel h:	.csv.gz			
	Custom UCSC browser of all M1 tracks			
Panel i:	https://genome.ucsc.edu/s/sarojas/hg38-mop-dense			

Extended Data

Panel j:	Browser https://brainome.ucsd.edu/annoj/BICCN_MOp/				
Extended Data	http://data.nemoarchive.org/brain/biccn/lab/lein/2020_M1_study_analysis/				
Figure 1:	Transcriptomics/flagship/cluster_overlap_plots				

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Figure 3: In situ cell-type identification, spatial mapping and projection mapping of individual cells in the MOp by MERFISH

Primary Data

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Figure 4. Correspondence between transcriptomic and morpho-electrical properties of mouse MOp neurons by Patch-seq, and cross-species comparison of L5 ET neurons.

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Panel		
b,c,j,k	Electrophysiology data	https://dandiarchive.org/dandiset/000008
		http://data.nemoarchive.org/biccn/lab/lein/lein/tran
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	10x V3 human (10X159-1	http://data.nemoarchive.org/biccn/lab/linnarsson/tr
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	10x V3 marmoset (bi005_m1,	http://data.nemoarchive.org/biccn/lab/feng/transcri
	bi006_m1)	ptome/sncell/raw/
	10x V3 mouse broad data (files	http://data.nemoarchive.org/biccn/grant/huang/mac
	with prefix pBICCNsMMrMOp)	osko/transcriptome/sncell/raw

Intermediate analyses

		http://data.nemoarchive.org/biccn/lab/lein/2020_M1_study_a	
	10x 4 species	nalysis/Transcriptomics/cross species integration/sample.co	
Panel h:	integration	mbined_exc_4_species_integration.RDS	
		http://data.nemoarchive.org/brain/biccn/lab/lein/2020_M1_st	
		udy analysis/Transcriptomics/flagship/projecting patch seq	
Panel h:	Code:	onto_umap	

Figure 5: Epi-Retro-Seq links molecular cell type with distal projection targets

Intermediate analyses

https://github.com/zhoujt1994/BICCN2020Flagship.git

Figure 6: Global wiring diagram and anatomical characterization of MOp-ul neuron types

Primary Data

Label	Link to registered swc (single cells) or 25 um	
in Fig	grid file (tracer)	high resolution image data

	http://download.alleninstitute.org/publications/cell	ftn://download.hrainimagalihrary.org
Dahias	ular anatomy of the mouse primary motor cort	
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	http://download.alleninstitute.org/publications/cell	-
	ular anatomy of the mouse primary motor cort	
AAV	ex/Viral Tracer Data in MOp 25 um nrrd/	84296
Cux2	$\underline{http://download.alleninstitute.org/publications/cell}$	
L2/3/4	ular anatomy of the mouse primary motor cort	
IT	ex/Viral Tracer Data in MOp 25 um nrrd/	n/a
	http://download.alleninstitute.org/publications/cell	
Nr5a1	ular_anatomy_of_the_mouse_primary_motor_cort	
L4 IT	ex/Viral Tracer Data in MOp 25 um nrrd/	n/a
	http://download.alleninstitute.org/publications/cell	
Tlx3	ular anatomy of the mouse primary motor cort	
L5 IT	ex/Viral Tracer Data in MOp 25 um nrrd/	n/a
Rbp4	http://download.alleninstitute.org/publications/cell	
L5	ular anatomy of the mouse primary motor cort	
IT+ET	ex/Viral Tracer Data in MOp 25 um nrrd/	n/a
	http://download.alleninstitute.org/publications/cell	http://connectivity.brain-
Sim1	ular anatomy of the mouse primary motor cort	map.org/projection/experiment/2977
L5 ET	ex/Viral Tracer Data in MOp 25 um nrrd/	<u>11339</u>
	http://download.alleninstitute.org/publications/cell	http://connectivity.brain-
Ntsr1	ular anatomy of the mouse primary motor cort	map.org/projection/experiment/1596
L6 CT	ex/Viral Tracer Data in MOp 25 um nrrd/	<u>51060</u>
IT	http://download.alleninstitute.org/publications/cell	/bil/proj/u19zeng/fMOST raw data/
projec	ular anatomy of the mouse primary motor cort	mouseID 374712-18453 CH2
tions	ex/Single Cell Reconstructions in MOp/	(pending upload)
IT	http://download.alleninstitute.org/publications/cell	
projec	ular anatomy of the mouse primary motor cort	
tions	ex/Single Cell Reconstructions in MOp/	<u>n/a</u>
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tions	http://ml-neuronbrowser.janelia.org/	<u>n/a</u>
ET	http://download.alleninstitute.org/publications/cell	/bil/data/2b/da/2bdaf9e66a246844/m
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PlxnD	PlxnD1-CreER;LSL-			http://brainarchitecture.org/viewer4/m
1	Flp	180722	Huang	ouse/map/8401F
PlxnD	PlxnD1-CreER;LSL-			http://brainarchitecture.org/viewer4/m
1	Flp	180730	Huang	ouse/map/28819F
	Tle4-CreER;LSL-			http://brainarchitecture.org/viewer4/m
Tle4	Flp	180605	Huang	ouse/map/28814F
	Tle4-CreER;LSL-			http://brainarchitecture.org/viewer4/m
Tle4	Flp	180816	Huang	ouse/map/8421F

Intermediate analyses

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	https://github.com/AllenInstitute/MOp_	code to reproduce rendering of
panel c	anatomy_rendering	registered data in 3D

Figure 7: Genetic tools for targeting cortical glutamatergic projection neuron types

Primary Data

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Panels					
d-i					
	Full	experi	Origi		
Label	Descrip	ment	nating	Link to Brain	
in Fig	tive ID	id	Lab	Architecture viewer	BIL link
	PlxnD1-			http://brainarchitectu	https://download.brainimagelibrary.org/
PlxnD	CreER;			re.org/viewer4/mous	84/aa/84aa97d12a6c17ba/180722_WG_
1	LSL-Flp	180722	Huang	<u>e/map/8401F</u>	PlxnD1lslFlpCFA1female_processed/
	PlxnD1-			http://brainarchitectu	https://download.brainimagelibrary.org/
PlxnD	CreER;			re.org/viewer4/mous	e9/2a/e92aa2dc0e14ad4d/180730_WG0
1	LSL-Flp	180730	Huang	e/map/28819F	10 PlxnD1 CFA female processed/
Fezf2	Fezf2-	180830	Huang	http://brainarchitectu	https://download.brainimagelibrary.org

	CreER;			re.org/viewer4/mous	/db/b8/dbb827c84942c557/180830_JH
	LSL-Flp			e/map/28827F	WG_Fezf2LSLflp_CFA_female/
					https://download.brainimagelibrary.org
	Fezf2-			http://brainarchitectu	/2b/6e/2b6e48dc425d16db/190903_JH
	CreER;			re.org/viewer4/mous	WG0006 Fezf2LSLflp MOp CFA f
Fezf2	LSL-Flp	190903	Huang	e/map/28917F	emale_processed/
	Tle4-			http://brainarchitectu	https://download.brainimagelibrary.org
	CreER;			re.org/viewer4/mous	/84/aa/84aa97d12a6c17ba/180605_WG
Tle4	LSL-Flp	180605	Huang	e/map/28814F	Tle4lslFlpRPCFA female processed/
	Tle4-			http://brainarchitectu	https://download.brainimagelibrary.org
	CreER;			re.org/viewer4/mous	/c8/1f/c81fe306a97b33e8/180816_JH_
Tle4	LSL-Flp	180816	Huang	e/map/8421F	WG_Tle4LSLFlpNPCfa_female/

Figure 8: Existence of L4 excitatory neurons in MOp.

Intermediate analyses

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Panels a,c,e	https://github.com/mukamel-lab/BICCN-Mouse-MOp/tree/master/flagship_fig8
Panel b	https://brainome.ucsd.edu/annoj/BICCN_MOp/

Figure 9: Two distinct L5 ET projection neuron types in MOp

Intermediate analyses

Panel d	https://brainome.ucsd.edu/annoj/BICCN_MOp/
Panel e	https://github.com/zhoujt1994/BICCN2020Flagship.git

Figure 10: An integrated multimodal census and atlas of cell types in the primary motor cortex of mouse, marmoset and human.

Intermediate analyses

1411 https://github.com/yal054/snATACutils

Extended Data

https://github.com/yal054/snATACutils
https://github.com/lhqing/flagship_tf_figure (code and data for Extended data figure 3)

SUPPLEMENTARY NOTES

Nomenclature of the L5 ET subclass of glutamatergic neurons

In this manuscript we have adopted a nomenclature for major subclasses of cortical glutamatergic excitatory neurons, which have long-range projections both within and outside of the cortex, following a long tradition of naming conventions that often classify neurons based on their projection targets. This nomenclature is based on our *de novo* transcriptomic taxonomy (**Fig. 10**) that organizes cell types hierarchically and validates the naming of the primary branches of glutamatergic neurons by their major long-range projection targets. At these levels, glutamatergic neurons are clearly divided into several subclasses, the cortico-cortical and cortico-striatal projecting intratelencephalic (IT) neurons that are distributed across nearly all layers (L2/3 IT, L4/5 IT, L5 IT, L6 IT and L6 IT Car3), the layer 5 neurons projecting to extratelencephalic targets (L5 ET), the cortico-thalamic (CT) projecting neurons in layer 6 (L6 CT), the near-projecting (NP) neurons found in layers 5 and 6, and the L6b neurons whose projection patterns remain largely unknown.

While the IT, CT, NP and L6b neurons have been consistently labeled as such in the field, the L5 ET neurons have not been named consistently in the literature, largely due to their large variety of projection targets and other phenotypic features that vary depending on cortical areas and species. Here we use the term L5 ET (layer 5 extratelencephalic) to refer to this prominent and distinct subclass of neurons as a standard name that can be accurately used across cortical regions and across species, and we provide our rationale below.

It has long been appreciated that cortical layer 5 contains two distinct populations of neurons that can be distinguished, not only based on the presence or absence of projections to ET targets (ET and IT cells), but also based on their predominant soma locations, dendritic morphologies and intrinsic physiology ⁸¹. Accordingly, various names incorporating these features have been adopted to refer to L5 ET versus L5 IT cells, such as L5b versus L5a, thick- versus thin-tufted and burst-firing versus regular-firing. And the most common term used to refer to L5 ET cells residing in motor cortical areas has been PT, which refers to neurons projecting to the pyramidal tract. As accurately stated in Wikipedia, "The **pyramidal tracts** include both the <u>corticobulbar tract</u> and the <u>corticospinal tract</u>. These are aggregations of <u>efferent nerve fibers</u> from the <u>upper motor neurons</u> that travel from the <u>cerebral cortex</u> and terminate either in the <u>brainstem</u> (*corticobulbar*) or <u>spinal cord</u> (*corticospinal*) and are involved in the control of motor functions of the body."

Due to the past wide use of the term PT, we do not take the decision to use L5 ET rather than PT lightly. However, in the face of multiple lines of evidence that have accumulated over the last several years ^{115,116} and prominently highlighted in this manuscript, it is now clear that PT represents only a subset of L5 ET cells and is thus unable to accurately encompass the entire L5 ET subclass. This realization is informed by comparisons across species and cortical areas, and

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by single-cell transcriptomics and descriptions of the projections of single neurons, as well as studies linking transcriptional clusters to projection targets.

As noted above, the overall transcriptomic relationships between cortical neurons are well-described by a hierarchical tree that closely matches developmental lineage relationships as neurons become progressively restricted in their adult fates ^{45,48} (**Fig. 10**). The cortical excitatory neurons are a major branch, distinct from inhibitory, glial, and epithelial cells. Subsequent splitting of the excitatory neurons reveals several major excitatory neuron subclasses – IT, L5 ET, L6 CT, NP and L6b. These major subclasses are conserved across mammalian species ^{15,18}, as well as across all cortical areas as shown in mouse ⁴⁴. It is therefore clear that names are needed that both accurately incorporate and accurately distinguish between neurons in these subclasses, and which are applicable across all cortical areas.

Also as noted above, a widely used alternative to L5 ET is PT. Further, this term is traditionally used along with CT to distinguish between cells with these different projections. The two main observations that make these alternative nomenclatures untenable are: 1) PT refers to motor neurons that project into medulla or spinal cord, but in many cortical areas (e.g. visual and auditory areas) none of the L5 ET cells are motor neurons; and 2) even in the motor cortex many cells in the L5 ET subclass do not project to the pyramidal tract and instead project solely to the thalamus (or to thalamus and other non-PT targets). This is revealed by single neuron reconstructions ^{26,68,86} (Fig. 6 and 9), BARseq ⁶⁷, projections from neuron populations with known gene expression and anatomical position in mouse lines ⁷¹, and studies directly linking projections to transcriptomics ^{15,54} and epigenetics ⁷⁹ (Fig. 5 and 9). The term PT therefore fails to be inclusive of the entire L5 ET subclass. Furthermore, the L5 CT cells within the L5 ET subclass are largely continuous with PT cells (or "PT-like" cells), not only genetically but also anatomically ^{54,64} (Fig. 3-4), as a majority of L5 ET cells project to multiple targets, typically including both the thalamus and the PT structures (e.g., medulla and spinal cord), as well as the midbrain (Fig. 6 and 9) ⁶⁸. Thus, the L5 ET subclass should neither be split into PT and CT, nor should the CT-only cells be omitted by use of the term PT. These facts also inform us that it is important to maintain a distinction between L5 CT (a type of L5 ET) and L6 CT (a major subclass of cortical excitatory neurons that is highly distinct from L5 ET, despite the presence of some L6 CT cells at the bottom of layer 5) 54. CT can be accurately used as a generic term, but CT neurons do not belong to a single subclass of cortical excitatory neurons.

We recognize that another name that has been used to describe L5 ET cells is SCPN (subcerebral projection neuron) ⁸². Given that the telencephalon is equivalent to the cerebrum, ET and subcerebral have the same meaning and the term L5-SCPN would be an accurate and equivalent alternative. But the "L5" qualifier is crucial in either case in order to distinguish these cells from the L6 CT subclass. We favor the use of ET because SCPN has not been widely adopted and due to symmetry with the widely used "IT" nomenclature. Alternatively, given their evidence that

"unlike pyramidal tract neurons in the motor cortex, these neurons in the auditory cortex do not project to the spinal cord", Chen et al ⁶⁷ used the term "pyramidal tract-like (PT-1)." We also favor L5 ET over L5 PT-l which clings to an inaccurate and now outdated nomenclature.

Supplementary Methods

Generation of Npnt-P2A-FlpO and Slco2a1-P2A-Cre mouse lines

To generate lines bearing in-frame genomic insertions of P2A-FlpO or P2A-Cre, we engineered

double-strand breaks at the stop codons of *Npnt* and *Slco2a1*, respectively, using

1509 ribonucleoprotein (RNP) complexes composed of SpCas9-NLS protein and in vitro transcribed

- 1510 sgRNA (Npnt: GATGATGTGAGCTTGAAAAG and Slco2a1: CAGTCTGCAGGAGAATGCCT).
- 1511 These RNP complexes were nucleofected into 10^6 v6.5 mouse embryonic stem cells
- 1512 (C57/BL6;129/sv; a gift from R. Jaenisch) along with repair constructs in which *P2A-FlpO* or
- 1513 P2A-Cre was flanked with the following sequences homologous to the target site, thereby
- enabling homology-directed repair.
- 1516 Npnt-P2A-FlpO:

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- 1517 TGGCCCTTGAGCTCTAGTGTTCCCACTTGCCATAGAAATCTGATCTTCGGTTTGGGGG
- 1518 AAGGGTTGCCTTACCATGCTCCATGAGTGAGCACTGGGAAAAGGGGCAGAGGAGGC
- 1519 CTGACCAGTGTATACGTTCTCCCTAGGTCATCTTCAAAGGTGAAAAAAGGCGTGG
- 1520 TCACACGGGGAGATTGGATTGGATGATGTGAGCTTGAAGCGCGGAAGATGTGGAA
- 1521 GCGGAGCTACTAACTTCAGCCTGCTGAAGCAGGCTGGAGACCGTGGAGGAGAACCCT
- 1522 GGACCTATGGCTCCTAAGAAGAAGAAGGAAGGTGATGAGCCAGTTCGACATCCTGTG
- 1523 CAAGACCCCGCCGAAGGTGCTGGTGCGGCAGTTCGTGGAGAGATTCGAGAGGCCCA
- 1524 GCGGCGAAAAGATCGCCAGCTGTGCCGAGCTGACCTGTGCTGGATGATC
- 1525 ACCCACAACGGCACCGCGATCAAGAGGGCCCACCTTCATGAGTTATAACACCATCAT
 1526 CAGCAACAGCCTGAGTTTTGACATCGTGAACAAGAGCCTGCAGTTCAAGTACAAGA
- 1527 CCCAGAAGGCCACCATCCTGGAGGCCAGCCTGAAGAAGCTGATCCCCGCATGGGAG
- 1528 TTCACGATTATCCCTTACAACGGCCAGAAGCACCAGAGCGACATCACCGACATCGT
- 1529 GTCCAGCCTGCAGCTGCAGTTCGAAAGCAGCGAGGAGGCCGACAAGGGGAATAGCC
- 1530 ACAGCAAGAAGATGCTGAAGGCCCTGCTGTCCGAAGGCGAGAGCATCTGGGAGATT
- 1531 ACCGAGAAGATCCTGAACAGCTTCGAGTACACCAGCAGATTTACCAAAACGAAGAC
- 1532 CCTGTACCAGTTCCTGTCCTGGCCACATTCATCAACTGCGGCAGGTTCAGCGACAT
- 1533 CAAGAACGTGGACCCGAAGAGCTTCAAGCTCGTCCAGAACAAGTATCTGGGCGTGA
- 1534 TCATTCAGTGCCTGGTCACGGAGACCAAGACAAGCGTGTCCAGGCACATCTACTTTT
- 1535 TCAGCGCCAGAGGCAGGATCGACCCCCTGGTGTACCTGGACGAGTTCCTGAGGAAC
- 1536 AGCGAGCCCGTGCTGAAGAGAGTGAACAGGACCGGCAACAGCAGCAGCAACAAGC
- 1537 AGGAGTACCAGCTGCTGAAGGACAACCTGGTGCGCAGCTACAACAAGGCCCTGAAG
- 1538 AAGAACGCCCCTACCCCATCTTCGCTATTAAAAACGGCCCTAAGAGCCACATCGGC

- AGGCACCTGATGACCAGCTTTCTGAGCATGAAGGGCCTGACCGAGCTGACAAACGT 1539 GGTGGGCAACTGGAGCGACAAGAGGGCCTCCGCCGTGGCCAGGACCACCTACACCC 1540
- 1541 ACCAGATCACCGCCATCCCGACCACTACTTCGCCCTGGTGTCCAGGTACTACGCCT
- 1542 ACGACCCCATCAGTAAGGAGATGATCGCCCTGAAGGACGAGACCAACCCCATCGAG
- 1543 GAGTGGCAGCACATCGAGCAGCTGAAGGGCAGCGCCGAGGGCAGCATCAGATACC
- 1544 CCGCCTGGAACGGCATTATAAGCCAGGAGGTGCTGGACTACCTGAGCAGCTACATC
- 1545 AACAGGCGGATCTGAAAGAGGTCGCTGCTGAGAAGACCCCTGGCAGCTCCCGAGCT
- 1546 AGCAGTGAATTTGTCGCTCTCCTCATTTCCCAATGCTTGCCCTCTTGTCTCCCTCTTA
- 1547 TCAGGCCTAGGGCAGGAGTGGGTCAGGAGGAAGGTTGCTTGGTGACTCGGGTCTCG
- 1548 GTGGCCTGTTTTGGTGCAATCCCAGTGAACAGTGACACTCTCGAAGTACAGGAGCAT
- 1549 CTGGAGACACCTCCGGGCCCTTCTG
- 1551 Slco2a1-P2A-Cre:

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- TGCCCCTGGGCCTCACCATACCTGTCTCTTCCTGCCTCATAGGTACCTGGGCCTACAG 1552
- GTAATCTACAAGGTCTTGGGCACACTGCTGCTCTTCTTCATCAGCTGGAGGGTGAAG 1553
- 1554
- 1555 TACTAACTTCTCCCTGTTGAAACAAGCAGGGGATGTCGAAGAGAATCCTGGACCTAT
- GGCTCCTAAGAAGAAGAAGGTGATGAGCCAGTTCGACATCCTGTGCAAGACTC 1556
- 1557
- AAGATCGCCAGCTGTGCCGCCGAGCTGACCTACCTGTGCTGGATGATCACCCACAAC 1558
- GGCACCGCCATCAAGAGGGCCACCTTCATGAGCTACAACACCATCATCAGCAACAG 1559
- 1560 CCTGAGCTTCGACATCGTGAACAAGAGCCTGCAGTTCAAGTACAAGACCCAGAAGG
- ${\sf CCACCATCCTGGAGGCCAGCCTGAAGAAGCTGATCCCCGCCTGGGAGTTCACCATC}$ 1561
- ATCCCTTACAACGCCCAGAGCACCAGAGCGACATCACCGACATCGTGTCCAGCCT 1562
- GCAGCTGCAGTTCGAGAGCAGCGAGGAGGCCGACAAGGCCAACAGCCACAGCAAG 1563
- 1564 AAGATGCTGAAGGCCCTGCTGTCCGAGGGCGAGAGCATCTGGGAGATCACCGAGAA
- 1565 GATCCTGAACAGCTTCGAGTACACCAGCAGGTTCACCAAGACCAAGACCCTGTACC
- 1566 AGTTCCTGTTCCTGGCCACATTCATCAACTGCGGCAGGTTCAGCGACATCAAGAACG
- 1567 TGGACCCCAAGAGCTTCAAGCTGGTGCAGAACAAGTACCTGGGCGTGATCATTCAG
- 1568 TGCCTGGTGACCGAGACCAAGACAAGCGTGTCCAGGCACATCTACTTTTTCAGCGCC
- CGTGCTGAAGAGAGTGAACAGGACCGGCAACAGCAGCAGCAACAAGCAGGAGTAC

AGAGGCAGGATCGACCCCCTGGTGTACCTGGACGAGTTCCTGAGGAACAGCGAGCC

- 1570
- CAGCTGCTGAAGGACAACCTGGTGCGCAGCTACAACAAGGCCCTGAAGAAGAACGC 1571
- 1572
- GATGACCAGCTTTCTGAGCATGAAGGGCCTGACCGAGCTGACAAACGTGGTGGGCA 1573
- 1574
- ACCGCCATCCCGACCACTACTTCGCCCTGGTGTCCAGGTACTACGCCTACGACCCC 1575
- ATCAGCAAGGAGATGATCGCCCTGAAGGACGAGACCAACCCCATCGAGGAGTGGCA 1576
- 1577 GCACATCGAGCAGCTGAAGGGCAGCGCCGAGGGCAGCATCAGATACCCCGCCTGGA
- 1578 ACGGCATCATCAGCCAGGAGGTGCTGGACTACCTGAGCAGCTACATCAACAGGCGG

1579 ATCTGACCTTCAGCTGGGACTACTGCCCTGCCCCAGAGACTGGATATCCTACCCCTC 1580 CACACCTACCTATATTAACTAATGTTAGCATGCCTTCCTCCTCCTTCC

Transfected cells were cultured and resulting colonies directly screened by PCR for correct integration using the following genotyping primers:

Genotyping primers

	Flanking Primer	Internal recombinase Primer
Npnt-P2A-FlpO Left homology arm	ATGCATTGCTTCATGCCATA	CCTTCAGCAGCTGGTACTCC
Npnt-P2A-FlpO right homology arm	GATTGAGGTCAGGCCAGAAG	TCGACATCGTGAACAAGAGC
Slco2a1-P2A-Cre Left homology arm	CTGGTGAAAGGGGAACTCTTGCT	GATCCCTGAACATGTCCATCAGG
Slco2a1-P2A-Cre Right homology arm	TACAGCATCCCTGACAAACACCA	TAGCACCGCAGGTGTAGAGAAGG

The inserted transgenes were fully sequenced and candidate lines were analyzed for normal karyotype. Lines passing quality control were aggregated with albino morulae and implanted into pseudopregnant females, producing germline-competent chimeric founders which in turn were crossed with the appropriate reporter lines on the C57/BL6 background.

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Brain Image Library (BIL) archive

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- 1971 imaging analysis: X.A., G.A.A., S.B., L.D., H.D., Z.F., N.N.F., W.G., H.G., J.A.H., J.T.H.,
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- transcriptomics (MERFISH): H.D., S.W.E., Z.Y., H.Z., M.Z., X.Z., B.Z. Multimodal profiling
- 1975 (PATCH-seq): P.B., J.B., M.B., Y.B., C.R.C., J.R.C., R.D., L.H., G.D.H., X.J., B.E.K., C.D.K.,
- 1976 A.L.K., D.K., S.L., E.S.L., E.M., S.M., P.R.N., S.F.O., R.S., F.S., K.S., S.A.S., Z.H.T., J.T.T.,
- 1977 A.S.T., H.Z. Transgenic tools: S.A., X.A., H.S.B., R.K.C., T.L.D., W.G., J.T.H., D.H., Z.J.H.,
- 1978 G.K., D.J.K., A.Y.L., K.S.M., J.N., D.A.S., B.T., M.B.V., X.W.Y., Z.Y., H.Z. NeMO archive
- 1979 and analytics: R.S.A., S.A.A., H.C.B., R.C., A.C., C.C., J.C., H.C., V.F., M.G., B.R.H., R.H.,
- 1980 J.K., A.M., C.M., L.N., D.O., J.O., M.S., O.W. Brain Image Library (BIL) archive: G.H., A.J.R.
- 1981 DANDI archive: B.D., S.S.G., M.G., Y.O.H., B.H. Brain Cell Data Center (BCDC): A.B., N.B.,
- 1982 B.C., F.D.D., K.D., J.C.G., T.H.G., M.H., F.K., K.K., M.E.M., L.N., C.T., T.L.T. Project
- 1983 management: P.B., F.D.D., H.G., H.H., K.K., D.K., B.B.L., K.S.M., S.M., M.N., F.S., S.S., C.T.

Competing interests

- 1987 A.B. is a cofounder of SciCrunch, a company devoted to improving scientific communication.
- 1988 J.R.E. is a member of Zymo Research SAB. J.A.H. is currently employed by Cajal Neuroscience.
- 1989 K.E.H. is currently employed by Cajal Neuroscience. P.V.K. serves on the Scientific Advisory
- 1990 Board to Celsius Therapeutics Inc. M.E.M. is a founder and CSO of SciCrunch Inc., a UCSD
- tech start up that produces tools in support of reproducibility including RRIDs. P.R.N. is
- 1992 currently employed by Cajal Neuroscience. A.R. is a founder and equity holder of Celsius
- 1993 Therapeutics, an equity holder in Immunitas Therapeutics and until August 31, 2020 was an SAB
- member of Syros Pharmaceuticals, Neogene Therapeutics, Asimov and ThermoFisher Scientific.
- 1995 From August 1, 2020, A.R. is an employee of Genentech. B.R. is shareholder of Arima
- 1996 Genomics, Inc. and Epigenome Technologies, Inc. K.Z is a co-founder, equity holder and serves
- on the Scientific Advisor Board of Singlera Genomics. X.Z. is a co-founder and consultant of
- 1998 Vizgen.

1999 2000 2001

1984 1985

1986

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