# **Neuron**

## A Single-Cell Transcriptomic Atlas of Human Neocortical Development during Mid-gestation

## Graphical Abstract



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## In Brief

An extensive single-cell catalog of cell types in the mid-gestation human neocortex extends our understanding of early cortical development, including subplate neuron transcriptomes, celltype-specific regulatory networks, brain evolution, and the cellular basis of neuropsychiatric disease.

## **Highlights**

- $\bullet$  High-resolution transcriptome map of 40,000 cells from developing human brain
- Cell-type-specific transcription factor (TF) expression and TF-gene networks
- Defines intermediate cell transition states during early neurogenesis
- **.** Implicates specific cell types in neuropsychiatric disorders





## A Single-Cell Transcriptomic Atlas of Human Neocortical Development during Mid-gestation

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#### SUMMARY

We performed RNA sequencing on 40,000 cells to create a high-resolution single-cell gene expression atlas of developing human cortex, providing the first single-cell characterization of previously uncharacterized cell types, including human subplate neurons, comparisons with bulk tissue, and systematic analyses of technical factors. These data permit deconvolution of regulatory networks connecting regulatory elements and transcriptional drivers to single-cell gene expression programs, significantly extending our understanding of human neurogenesis, cortical evolution, and the cellular basis of neuropsychiatric disease. We tie cell-cycle progression with early cell fate decisions during neurogenesis, demonstrating that differentiation occurs on a transcriptomic continuum; rather than only expressing a few transcription factors that drive cell fates, differentiating cells express broad, mixed cell-type transcriptomes before telophase. By mapping neuropsychiatric disease genes to cell types, we implicate dysregulation of specific cell types in ASD, ID, and epilepsy. We developed CoDEx, an online portal to facilitate data access and browsing.

INTRODUCTION

encompass hundreds or thousands of distinct cell types, each with unique functions [\(Silbereis et al., 2016](#page-17-0)). Groundbreaking work in mouse revealed the power of single-cell transcriptomics to provide a framework for understanding the complexity and heterogeneity of cell types in the brain ([Hrvatin et al., 2018; Loo et al.,](#page-16-0) [2019; Macosko et al., 2015; Saunders et al., 2018; Shekhar et al.,](#page-16-0) [2016; Tasic et al., 2016; Zeisel et al., 2018](#page-16-0)). The availability of highquality tissue and advances in single-cell transcriptomic technologies permit us to catalog the cell-type diversity of the human cortex in a comprehensive and unbiased manner [\(Ecker et al., 2017\)](#page-16-1).

The human cortex is composed of billions of cells estimated to

Despite the enormous progress that has been made in characterizing early cortical development [\(Geschwind and Rakic, 2013;](#page-16-2) [Lui et al., 2011; Silbereis et al., 2016\)](#page-16-2), many of the molecular mechanisms underpinning the generation, differentiation, and development of the diverse types of cells remain largely unknown (Molnár, [2011\)](#page-17-1). Molecular taxonomies of cortical cell types from developing human brains enable us to understand the mechanisms of neurogenesis and how the remarkable cellular diversity found in the human cortex is achieved [\(Camp et al., 2015; Fan et al., 2018; Liu](#page-16-3) [et al., 2016; Nowakowski et al., 2017; Pollen et al., 2015; Zhong](#page-16-3) [et al., 2018](#page-16-3)). Several studies have taken a first step in this direction, analyzing several hundred or a few thousand cells from developing human brain ([Fan et al., 2018; Liu et al., 2016; Nowakowski et al.,](#page-16-4) [2017; Pollen et al., 2015; Zhong et al., 2018](#page-16-4)). Advances in technology and throughput (e.g., Drop-seq; [Macosko et al., 2015](#page-17-2)) allow us

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to analyze an order of magnitude more cells to complement and extend these studies, providing a deeper picture of human cortical development and its perturbation in disease.

#### RESULTS

#### A Catalog of Cell Types in Developing Human Neocortex Identifies Major Cell Types, Progenitor States, and Subtypes of Excitatory and Inhibitory Cells

Here we use single-cell RNA sequencing (scRNA-seq) to define cell types and compile cell-type transcriptomes in the developing human neocortex. We focus on the cortical anlage at mid-gestation (gestation week [GW] 17 to GW18) [\(Figure 1](#page-2-0)A), because this period contains the major germinal zones and the developing cortical laminae containing migrating and newly born neurons, and neurodevelopmental processes occurring during this epoch are implicated in neuropsychiatric disease [\(de la Torre-Ubieta et al., 2016; Gandal et al., 2016\)](#page-16-5). To optimize detection of distinct cell types, we separated the cortex into the germinal zones (ventricular zone [VZ] and subventricular zone [SVZ]) and developing cortex (subplate [SP] and cortical plate [CP]) before single-cell isolation. Using Drop-seq [\(Macosko](#page-17-2) [et al., 2015](#page-17-2)), we obtained and compared high-quality profiles for  $\sim$ 40,000 cells from human cortex ([Figures 1A](#page-2-0), [S1A](#page-13-0), and S1B; [Tables S1](#page-13-0), [S2,](#page-13-0) and [S3](#page-13-0)) and a small subset with microfluidic approaches (Fluidigm) for technical comparisons.

We first applied unbiased clustering based on t-distributed stochastic neighbor embedding (tSNE; see [STAR Methods\)](#page-18-0) and spectral K nearest-neighbor graph-based clustering ([But](#page-16-6)[ler et al., 2018\)](#page-16-6), identifying 16 transcriptionally distinct cell groups. Cell types originated from the expected anatomical source and clustered by biological cell type, rather than batch or technical artifacts [\(Figures 1B](#page-2-0)–1G, [S1](#page-13-0)C, and S1D). We identified multiple groups of cells at different stages of neuronal differentiation and maturation, corresponding to all known major cell types at this developmental period ([Figures 1](#page-2-0)B–1F and [S1](#page-13-0)E; [Table S4](#page-13-0)). Clusters contained between 50 and 2,000 cells. The smallest cluster captured, which belonged to microglia, was composed of  $\sim$ 50 cells. Other small clusters for oligodendrocyte precursors (OPCs), endothelia, and pericytes were composed of 306, 237, and 114 cells, respectively [\(Figure 1G](#page-2-0); [Table S4](#page-13-0)). Clusters were reproducible and robust, as ascertained by bootstrapping [\(Figure S1F](#page-13-0)). Ordering of cells by pseudo-time in an unbiased manner using Monocle 2, a computational method that performs lineage trajectory recon-struction based on single-cell transcriptomics data ([Qiu et al.,](#page-17-3) [2017; Trapnell et al., 2014\)](#page-17-3), confirmed the predicted developmental trajectory ([Figures 1H](#page-2-0) and 1I). For example, it is possible to observe the ordered transitions between different neural progenitor types and maturing glutamatergic neurons, with radial glia (RG) transitioning to intermediate progenitors (IPs) and IPs transitioning to newborn migrating neurons [\(Figure 1](#page-2-0)I).

We observed that cell-type detection appears to be more sensitive to the number of cells profiled than the sequencing depth [\(Figures S2](#page-13-0)A–S2D). Furthermore, although each cell profile is an incomplete representation of that cell type [\(Lun et al., 2016;](#page-16-7) [Fig](#page-13-0)[ures S2](#page-13-0)E and S2F), pooling transcriptomes within cells of a given type provides more complete cell-type transcriptome representations. We iteratively subsampled cells from clusters to empirically assess the completeness of cell-type signatures with different sample sizes ([Figure S2](#page-13-0)G). At a depth of 40,000 cells, we obtain stable transcriptomes representing 3,000–5,000 genes for most cell types present [\(Table S4](#page-13-0)). Comparison of these data with a method using lower throughput and higher sequencing depth (Fluidigm C1) ([Nowakowski et al., 2017;](#page-17-4) [Figure S3\)](#page-13-0) revealed that the ability to leverage an order of magnitude more cells yielded more stable mRNA transcript profiles for a given cell type [\(Figures S3A](#page-13-0)–S3C). Integration of our dataset with the largest previous study (4,000 cells; [Nowakowski et al., 2017\)](#page-17-4) using canonical correlation analysis ([Butler et al., 2018\)](#page-16-6) showed substantial alignment of cells between the two datasets ([Figures](#page-13-0) [S3](#page-13-0)D and S3E). This is the first direct comparison of different human fetal single-cell datasets, and it demonstrates the reproducibility of these expression profiles and significantly extends them by providing more stable gene expression rankings. We provide these cell-type-specific expression profiles with annotated gene expression ranking confidence measures for each cell type [\(Table S4](#page-13-0)) and a web interface, CoDEx (Cortical Development Expression Viewer), for browsing these data ([http://](http://geschwindlab.dgsom.ucla.edu/pages/codexviewer) [geschwindlab.dgsom.ucla.edu/pages/codexviewer\)](http://geschwindlab.dgsom.ucla.edu/pages/codexviewer).

Comparison of scRNA-seq datasets to bulk RNA sequencing (RNA-seq) expression profiles ([de la Torre-Ubieta et al., 2018](#page-16-8)) showed consistently that gene expression profiles generated

#### Figure 1. A Catalog of Cell Types in Developing Human Neocortex

(G) Cluster metrics. Ratio of cells derived from the germinal zone (GZ) or CP. Percentage of total cell population. Percentage of cells derived from each donor. Bar colors indicate grouping of cells by major cell class, e.g., caudal ganglionic eminence (CGE)- and MGE-derived interneurons are both blue. MP, mitotic progenitor. (H) Pseudo-time analysis of cells expected to be part of the neurogenesis-differentiation axis, colored by Monocle state or pseudo-time. Each point represents a cell. Pseudo-time represents an ordering of cells based upon the inferred trajectory, predicting the lineage trajectory.

(I) Pseudo-time trajectory colored by Seurat clusters.

<sup>(</sup>A) Schematic illustrating experimental design and anatomical dissections. VZ, ventricular zone; iSVZ, inner subventricular zone; oSVZ, outer subventricular zone; IZ, intermediate zone; SP, subplate; CPi, inner cortical plate; CPo, outer cortical plate; RG, radial glia; IP, intermediate progenitor; MN, newborn migrating excitatory neuron; EN, excitatory neuron; IN, interneuron; O, oligodendrocyte precursor; E, endothelial cell; P, pericyte; M, microglia.

<sup>(</sup>B) Scatterplot visualization of cells after principal-component analysis and t-distributed stochastic neighbor embedding (tSNE), colored by Seurat clustering and annotated by major cell types.

<sup>(</sup>C) Heatmap of gene expression for each cell. Cells are grouped by Seurat clustering, and the mean expression profile of enriched genes for each cluster was used to hierarchically cluster the Seurat clusters. The top 20 most enriched genes are shown per cluster with canonical marker genes noted. Color bar matches Seurat clusters in (B).

<sup>(</sup>D and E) tSNE of cells colored by anatomical source (D), or mean expression of groups of canonical marker genes of major cell types (E).

<sup>(</sup>F) Heatmap of expression profiles of canonical cell-type marker genes. Cells are grouped by Seurat clustering. Color bar matches Seurat clusters in (B).

<span id="page-4-0"></span>![](_page_4_Figure_1.jpeg)

using different scRNA-seq methodologies across different laboratories strongly correlated with bulk RNA-seq gene expression profiles (Spearman 0.69–0.83) ([Figures S4A](#page-13-0) and S4B). However, we observed that approximately 400 proteincoding genes representing longer, brain-enriched, cell adhesion molecules involved in neuronal development [\(Figures S4](#page-13-0)E–S4H) were consistently under-represented in single-cell datasets compared with bulk tissue RNA-seq ([Figures S4](#page-13-0)C and S4D). Overall comparison of expression of canonical cell-type marker genes showed similar expression levels compared with bulk tissue RNA-seq [\(Figure S4](#page-13-0)I), indicating that despite small biases in gene detection shared across scRNA-seq methods, the relative frequencies of major cell types were not over- or under-represented, further demonstrating the robustness of the scRNAseq dataset [\(Figure S4](#page-13-0)).

We next reasoned we could use the depth of our dataset to identify cell states and cell subtypes not identified in previous studies with smaller cell numbers. We performed an additional round of clustering on each major cell cluster [\(Figure 2;](#page-4-0) [Table](#page-13-0) [S5](#page-13-0); [STAR Methods](#page-18-0)), which finely resolved maturation states during neurogenesis and identified multiple cell subtypes not previously characterized in single-cell datasets in humans: SP neurons [\(Figure 2D](#page-4-0)), distinct subtypes of glutamatergic neurons [\(Figure 2D](#page-4-0)), early parvalbumin (PV) interneurons, and NPY expressing SST interneurons ([Figure 2C](#page-4-0)). A previous study profiling  $\sim$ 2,300 cells from developing human cortex did not detect PV interneurons and suggested PV interneurons may not develop until after GW26 ([Zhong et al., 2018](#page-17-5)). Here, we find that at mid-gestation (GW17–GW18), PV interneurons comprise  $\sim$ 0.1% of the total population, underscoring how necessary larger datasets are to identify rare cell types.

The provenance of human neocortical interneurons has been disputed [\(Hansen et al., 2013; Ma et al., 2013; Radonji](#page-16-9)ć [et al.,](#page-16-9) [2014; Zhong et al., 2018\)](#page-16-9). We observed no clusters of progenitors expressing markers of interneurons and no clusters of interneurons expressing mitotic or progenitor markers ([Figure S1](#page-13-0)E). In addition, subclustering of interneurons did not identify a cell population displaying characteristics of interneuron progenitors. OLIG2 is a marker of both medial ganglionic eminence progenitors and OPCs ([Miyoshi et al., 2007\)](#page-17-6). We observed OLIG2+ cells

only in the OPC cluster that express other OPC markers but do not express interneuron marker genes [\(Figure S1E](#page-13-0)). Thus, even with the order-of-magnitude-greater cell depth and increased ability to detect low-abundance cell types (e.g., 0.1%), we do not find evidence of a neocortical interneuron progenitor during mid-gestation in humans.

#### Cell-Type Enrichment of Transcription Factors and Co-factors

We next sought to gain insight into cell-type-specific regulatory programs by comparing transcription factor (TF) expression across major cell types. We found previously characterized TFs and co-factors enriched in their corresponding cell types [\(Figures 1F](#page-2-0) and [3](#page-6-0)A) and multiple TFs and co-factors that have not been associated with specific neocortical cell types [\(Fig](#page-6-0)[ure 3](#page-6-0)). These TFs also displayed laminae-specific expression in a bulk tissue laser-capture micro-dissected (LCM) expression dataset ([Miller et al., 2014](#page-17-7)) and temporal trajectories similar to canonical cell-type markers [\(Figures 3B](#page-6-0) and [S5](#page-13-0)A).

To validate predictions for these putative novel cell-type markers, we performed RNA fluorescence *in situ* hybridization (FISH), which confirmed laminae-specific expression of each of the TFs and co-factors tested: *ZFHX4* and *CARHSP1* in neural progenitors and *CSRP2* in excitatory neurons [\(Figures 3C](#page-6-0)–3H). Of particular interest was *ZFHX4*, which has been previously associated with 8q21.11 microdeletion syndrome [\(Palomares](#page-17-8) [et al., 2011](#page-17-8)). Our data localize ZFHX4 specifically to neural progenitors in the developing human neocortex for the first time [\(Figures 3A](#page-6-0)–3E), implicating specific dysregulation of neural progenitors as the mechanism underlying this syndrome.

The TF ST18 appeared to partially cluster with SP markers [\(Figures 2](#page-4-0)D, [4A](#page-8-0), and 4B; Hoerder-Suabedissen and Molnár, [2015; Oeschger et al., 2012](#page-16-10)). We found that SP markers previously defined in other species were not uniquely expressed in the SP in another fetal gene expression atlas [\(Miller et al.,](#page-17-7) [2014;](#page-17-7) [Figures 4B](#page-8-0) and 4C). We identified SP-enriched genes in this cortical laminar atlas ([Figure 4](#page-8-0)C; [STAR Methods\)](#page-18-0), which showed strong overlap with ST18 [\(Figure 4B](#page-8-0)). Subclustering separated deep-layer neurons from the ST18-expressing SP neurons ([Figure 4](#page-8-0)D). Genes enriched in the SP neuron cluster,

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Figure 2. Subclustering Analysis Identifies Progenitor States and Subtypes of Excitatory and Inhibitory Cells
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(A) Diagram of subclustering analysis workflow. An iterative approach was used; cells from each initial cluster were re-processed, clustered, and analyzed from the raw count matrix using Seurat. tSNE is colored by Seurat clustering and annotated by major cell types.

(B) Subclustering of progenitors. Progenitors separate by cell type and cell-cycle state.

<sup>(</sup>C) Subclustering of interneurons. Interneuron MGE (InMGE) subclusters by maturity and cell subtype. InMGE-7 displays enrichment of TAC1, a marker of PV interneurons, and does not express SST [\(Pfeffer et al., 2013](#page-17-9)). InMGE-6 shows strong enrichment of NPY and SST. Interneuron CGE (InCGE) subclusters by maturity. All clusters are CALB2+, with differing levels of expression likely reflecting maturity.

<sup>(</sup>D) Subclustering of excitatory neurons. Newborn excitatory neurons (ExNs) and maturing excitatory neurons (ExMs) subcluster by maturity. ExMs begin to display separation of laminae markers. The excitatory upper-layer-enriched cluster (ExM-U) shows enrichment of laminae markers for different subclusters and expression of the callosal marker LMO4 ([Molyneaux et al., 2007\)](#page-17-10). The deep-layer cluster (ExDp1) separates by layer. ExDp1-2 is enriched for the subplate marker NR4A2 (Hoerder-Suabedissen and Molnár, 2015), ExDp1-0 is enriched for lower L5 and L6 markers (CRYM, TBR1, and FOXP2) ([Molyneaux et al., 2007\)](#page-17-10), and ExDp1-1 and ExDp1-3 are enriched for L4 and upper L5 markers (RORB, FOXP1, and ETV1) [\(Ferland et al., 2003; Molyneaux et al., 2007\)](#page-16-11).

Heatmaps of expression profiles by subcluster of groups or individual marker genes (y axis). The laminae bar indicates the percentage of cells derived from the CP. Purple: 100% of cells derive from the CP, 0% GZ; green: 0% of cells derive from the CP, 100% GZ. Upper-layer and deep-layer gene groups are the top 50 most enriched genes from the excitatory upper-layer-enriched cluster and the deep-layer cluster, respectively. tSNEs of cells are colored by features of interest: subcluster, anatomical source, donor, or gene expression. Gray indicates cells with an undefined transcriptional signature. For heatmaps and tSNE, gene expression is plotted as a *Z* score for the population of cells in the plot; therefore, some cell types display differences in relative expression of cell-type markers between subclusters of the same major cell type, but all express the marker at some level (e.g., all RG express markers of RG, but some subclusters of RG have higher relative expression than other subclusters of RG). Labels "mat" and "dif" indicate inferred order of maturation or differentiation, respectively.

<span id="page-6-0"></span>![](_page_6_Figure_1.jpeg)

or highly correlated with ST18, display strong SP enrichment in the cortical laminae dataset, verifying our capture of SP neurons and identification of many additional SP neuron markers [\(Figures](#page-8-0) [4E](#page-8-0)–4H). In addition, we performed RNA FISH to confirm SPspecific expression of ST18 ([Figure 4I](#page-8-0)). This represents the first transcriptomic characterization of human SP neurons at singlecell resolution.

We next reasoned that we could begin to leverage these single-cell data to uncover some cellular and molecular mechanisms driving human cortical evolution by determining whether specific cell types were enriched with genes showing human specific expression trajectories (hSET) in bulk tissue ([Bakken](#page-16-12) [et al., 2016](#page-16-12); see [STAR Methods](#page-18-0)). We observed the strongest enrichment of hSET genes in outer RG (oRG) and the excitatory upper-layer-enriched cluster [\(Figure S5](#page-13-0)B). Both of these cell types represent processes central to both neocortical expansion ([Lui et al., 2011\)](#page-16-13) and elaboration of cortical connectivity in humans ([Fame et al., 2011\)](#page-16-14). Among the approximately 600 genes with oRG-enriched expression, we identified *LYN*, a Src tyrosine kinase previously implicated in neuronal polarization and a-amino-3-hydroxy-5-methyl-4-isoxazolepropionic acid (AMPA) signaling ([Hayashi et al., 1999; Namba et al.,](#page-16-15) [2014\)](#page-16-15) that had not been previously associated with this cell type. We used a fetal LCM atlas ([Miller et al., 2014\)](#page-17-7) and RNA FISH to validate these observations, showing that LYN localized to the germinal zones and was specifically expressed in the VZ and outer SVZ (oSVZ) ([Figures S5](#page-13-0)C and S5D).

#### Mapping of Cell-Type-Specific Gene Regulatory Networks in the Developing Human Neocortex

To deconvolute the cell-type specificity of regulatory elements, we leveraged a recently generated map of regulatory elements active in developing fetal cortex and their putative target genes [\(de la Torre-Ubieta et al., 2018\)](#page-16-8) to identify promoters and enhancers regulating the expression of genes enriched in cells defined in this study ([STAR Methods](#page-18-0); [Table S6](#page-13-0)). Enhancers associated with specific cell types were characterized by remarkable consistency in mean enhancer size, number associated with each gene, and distance to the target gene for each cell type ([Figures 5](#page-10-0)A–5F). In addition, there was no correlation between target gene length or GC content and number of associated enhancers [\(Figures 5G](#page-10-0) and 5H). We extended this map by computationally reconstructing gene regulatory networks using the single-cell regulatory network inference and clustering (SCENIC) pipeline ([Aibar et al., 2017](#page-15-0); [Figure 5](#page-10-0)I) with empirically determined regulatory elements [\(de la Torre-Ubieta et al.,](#page-16-8) [2018\)](#page-16-8), rather than standard promoter annotations. This produced 124 regulons—each representing a TF, along with a set of co-expressed and motif-enriched target genes—and the regulon activity scores for each cell [\(Table S7\)](#page-13-0). Multiple TFs previously associated with specific cell types showed enriched regulon activity in the expected cell types ([Figures 5J](#page-10-0) and 5K; [Table S7\)](#page-13-0). We also identified TFs with previously uncharacterized cell-type- or cell-subtype-specific activity, including *NFE2L2* in RG, *NHLH1* in post-mitotic IPs, *ZNF354C* in excitatory neurons, and *BACH2* in maturing excitatory neurons ([Figure 5](#page-10-0)L; [Table S7](#page-13-0)). This represents a first-generation map of cell-type-specific gene regulatory networks in the developing human neocortex.

#### Dissecting the Acquisition of a Neuronal Program

Neurons are generated from the controlled asymmetric division of neural progenitors, which prompted us to analyze the distinct transcriptional states of cycling cells during this process [\(Lui](#page-16-13) [et al., 2011\)](#page-16-13). Neural progenitors clustered by cell-cycle state in addition to cell type ([Figures 1E](#page-2-0), [6](#page-12-0)A, and [S6](#page-13-0)A–S6C), with about 30% of progenitors cycling, roughly consistent with previous observations (37% based on immunostaining) ([Hansen et al., 2010\)](#page-16-16). We also observed that many cycling progenitors individually expressed markers of several distinct major cell types, including RGs, IPs, and neurons ([Figures 6](#page-12-0)A–6C). Doublets were an insufficient explanation for the co-expression of distinct cell-type makers for multiple reasons, including that the number of cells expressing multiple major cell-type markers is twice the empirically assessed doublet rate [\(Table S2;](#page-13-0) [Figure S1B](#page-13-0); [STAR](#page-18-0) [Methods\)](#page-18-0) and the highly non-random distribution of the cell types expressing markers of two cell types ([Figure 6](#page-12-0)C).

Therefore, as an alternative explanation, we hypothesized that we were identifying an intermediate or transition state: mitotically active cells in the early stages of neurogenesis, i.e., RG producing IPs, RG producing neurons, and IPs producing neurons. Consistent with this hypothesis, mixed marker cells progressing through different stages of the cell cycle consistently displayed transcriptomes composed of multiple major cell types [\(Figures 6D](#page-12-0), [S7](#page-13-0)A and S7B). By S phase, RG+IP+ and IP+Neuron+ cells more closely resembled their presumed endpoint cell type: IP and neuron, respectively [\(Figures 6](#page-12-0)D and [S7B](#page-13-0)). The transcriptomic signature of RG+Neuron+ S phase and G2/M phase cells was closer to RG, potentially reflecting the greater dissimilarity between RG and neurons ([Figures 6D](#page-12-0), [S7A](#page-13-0), and S7B). In addition, the mixed marker cells share a high percentage of the endpoint cell-type

#### Figure 3. Cell-Type Enrichment of TFs and Co-factors

<sup>(</sup>A) Heatmap of expression of TFs, co-factors, and chromatin remodelers enriched in RG, excitatory neurons, and deep-layer excitatory neurons. Cells are grouped by cluster. Red indicates factors previously unknown to be enriched in the neocortical cell types of interest.

<sup>(</sup>B) Expression of factors of interest in bulk tissue LCM laminae from developing cortex.

<sup>(</sup>C) RNA FISH of fetal cortex probed with the newly identified cell-enriched TF ZFHX4 (neural progenitors in the VZ and SVZ) and known markers PAX6 (RG marker) and EOMES (IP marker). Insets show higher magnification of the VZ and SVZ. Scale bar, 250  $\mu$ m (left) or 100  $\mu$ m (inset).

<sup>(</sup>D) Quantification of normalized fluorescence intensity per layer for each set of probes (see [STAR Methods](#page-18-0)). Mean ± SE displayed in barplot.

<sup>(</sup>E) Quantification of the percentage of PAX6+ or EOMES+ cells co-expressing ZFHX4. ZFHX4 is expressed in both RG and IPs. Mean ± SE displayed in barplot. (F and G) RNA FISH of fetal cortex probed with the newly identified cell-enriched TFs CARHSP1 (neural progenitors in the VZ and SVZ) and CSRP2 (glutamatergic neurons in the CP). Bar graph shows quantification of normalized fluorescence intensity per layer for each set of probes (see [STAR Methods](#page-18-0)). Mean ± SE displayed in barplot. Scale bar,  $250 \mu m$  (top) or 100  $\mu m$  (inset).

<sup>(</sup>H) Schematic of cell-type-specific expression of factors of interest. Color indicates  $-\log_{10} p$  value from Fisher's test.

<span id="page-8-0"></span>![](_page_8_Figure_1.jpeg)

signature, but the magnitude of expression of the cell-typerelevant signature genes is smaller than in cells in the fully differentiated cell clusters [\(Figures 6E](#page-12-0) and [S7C](#page-13-0)–S7F). Mixed marker cells not in S, G2, or M phase may represent cells starting to cycle and differentiate, consistent with findings in mice that some RG precursors also express neuronal marker genes of both deep and superficial layers, representing transcriptionally primed cells [\(Zahr et al., 2018](#page-17-11)). Alternatively, these mixed marker cells may be newborn cells that retain some transcripts of the mother cell type, as has been previously suggested in mice ([Zahr et al., 2018; Zhong et al., 2018](#page-17-11)).

To independently validate the existence of cells in these transition states, we performed RNA FISH, observing S-phase neural progenitors in the VZ expressing both PAX6 and STMN2, indicating an induction of a neuronal program in a cell before its neurogenic division [\(Figure 6](#page-12-0)F). Indeed, 8.9% (VZ), 6.7% (inner SVZ [iSVZ]), and 7.5% (oSVZ) of these cells co-express markers of RG and neurons ([Figure 6](#page-12-0)G), confirming our scRNA-seq data (see [STAR Methods\)](#page-18-0). We were able to quantify the relative proportions of progenitors undergoing distinct differentiation divisions ([Figure 6](#page-12-0)H), finding that RG produce roughly equal numbers of RG, IPs, and neurons but that IPs produce approximately two times as many neuronal progeny as IP progeny. Altogether, these results indicate that during early neurogenesis (1) cell fate decisions occur before S phase; (2) differentiating parent cells not only express the few key TFs that drive cell fates but also express broad, mixed cell-type transcriptomes; and (3) neural cell-type differentiation occurs on a continuum and involves transcriptomic transitions tied to cell-cycle progression ([Figure 6](#page-12-0)I).

#### Cellular Determinants of Disease

We next reasoned that we could use this atlas of developing human brain cell types to identify the developmental stages and cell types in which mutations causing high risk for neuropsychiatric disease act so as to provide a reference for understanding disease mechanisms and circuits ([Figure 7\)](#page-14-0). We first examined enrichment of high-confidence risk genes for autism spectrum disorder (ASD), defined by harboring high-risk, likely protein-disrupting mutations ([Sanders et al., 2015;](#page-17-12) [Figures 7](#page-14-0)A, 7D, and [S8](#page-13-0)A). Most ASD risk genes were expressed in developing glutamatergic neurons, both deep and upper layer ([Figures 7](#page-14-0)A and 7D), consistent with previous studies [\(Amiri et al., 2018; Parik](#page-16-17)[shak et al., 2013\)](#page-16-17). However, at the individual gene level, there is substantial variability, and several genes are expressed in inhibitory neurons, as well as excitatory neurons or progenitors [\(Figures 7](#page-14-0)A and [S8](#page-13-0)A). For example, MYT1L and AKAP9 display pan-neuronal expression, whereas GRIN2B is glutamatergic subtype specific and ILF2 is expressed in cycling progenitors [\(Figures 7A](#page-14-0) and [S8](#page-13-0)A). In the adult neocortex, expression again concentrated in glutamatergic neurons, with some genes exhibiting more pan-neuronal expression patterns [\(Figure S8A](#page-13-0)).

In addition, our expanded atlas of cell types identified several genes that showed distinct patterns of extra-neuronal expression, including SLC6A1, which was enriched in pericytes, and TRIO, SETD5, TCF7L2, and KAT2B, which were enriched in OPCs [\(Figures 7](#page-14-0)A and [S8](#page-13-0)A). For the first time, these data suggest that cell types involved in maintenance of the blood-brain barrier and the peri-neural environment may also mediate ASD risk. Several of these genes are expressed in different cell types in the adult neocortex, such as SLC6A1 in interneurons, highlighting the importance of broader single-cell catalogs [\(Fig](#page-13-0)[ure S8](#page-13-0)A). Expanding this analysis to high-confidence intellectual disability (ID) and epilepsy risk genes ([Figures 7B](#page-14-0)–7D, [S8B](#page-13-0) and S8C) showed that most epilepsy risk genes are expressed in glutamatergic neurons ([Figures 7B](#page-14-0), 7D, and [S8](#page-13-0)B). ID risk genes were also enriched in glutamatergic neurons but also showed enrichment in RG, which was not observed with ASD or epilepsy [\(Figures 7C](#page-14-0), 7D, and [S8C](#page-13-0)). The impact on early progenitor types in ID relative to ASD and epilepsy is consistent with the more severe disease phenotype in ID. Although the results for ID were highly significant, the ID risk gene list is smaller, making the comparisons less powered. Altogether, these results demonstrate cell-type-specific expression of ASD, epilepsy, and ID risk genes by mid-fetal development and provide a framework for the cellular and developmental context in which individual ASD, epilepsy, and ID genes should optimally be studied.

Most neuropsychiatric disease risk loci are found in the noncoding genome, in which functional interpretation is hampered by limited knowledge of the genomic location and spatiotemporal activity of regulatory elements. Leveraging our cell-type-specific map of regulatory elements active in the human neocortex ([Fig](#page-15-1)[ure 8](#page-15-1)A; [Table S6](#page-13-0); [STAR Methods;](#page-18-0) [de la Torre-Ubieta et al.,](#page-16-8) [2018\)](#page-16-8), we used a partitioned heritability approach based on linkage disequilibrium (LD) score regression [\(Finucane et al., 2015](#page-16-18)) to identify cell types enriched for variants influencing brain volume or cognition or causing risk for neuropsychiatric disease. We found that variants influencing adult intracranial volume [\(Adams et al.,](#page-15-2) [2016\)](#page-15-2) were specifically enriched in the regulatory elements of

Figure 4. Characterization of Subplate Neuron Expression Profiles

(A) tSNE colored by Seurat clustering and annotated by major cell types.

<sup>(</sup>B) tSNE of cells colored by mean expression of groups of marker genes or expression of specific genes.

<sup>(</sup>C) Expression of SP markers in bulk tissue LCM laminae from developing cortex. SP markers were derived from literature sources (left) or by differential expression of the SP versus the VZ, SVZ, CP, and marginal zone (MZ) and visual confirmation of SP specificity (right).

<sup>(</sup>D) Subclustering of deep-layer excitatory cluster 1. tSNE for the full dataset colored by subclustering (left). tSNE of cells belonging to the deep-layer excitatory cluster (right), colored by subclustering, mean expression of groups of marker genes, or expression of specific genes.

<sup>(</sup>E–G) Expression of SP cluster-enriched genes (F), ST18 co-expressed genes (G), and (E) the intersection of (F) and (G) in bulk tissue LCM laminae from developing cortex. Genes are ordered left to right by enrichment or correlation (highest left). Light blue text indicates previously identified SP markers.

<sup>(</sup>H) Eigengene of intersected ST18 co-expressed and SP cluster-enriched genes from (E) plotted in bulk tissue LCM laminae from developing cortex. Box indicates first and third quartiles; the whiskers extend from the box to the highest or lowest value that is within 1.5 x interquartile range of the box; and the line is the median. \*p < 0.05, \*\*p < 0.01, \*\*\*p < 0.001, \*\*\*\*p < 0.0001.

<sup>(</sup>I) RNA FISH of fetal cortex probed with the newly identified subplate-enriched TF ST18. Bar graph shows quantification of normalized fluorescence intensity per layer for each set of probes (see [STAR Methods\)](#page-18-0). Mean  $\pm$  SE displayed in barplot. Scale bar, 250  $\mu$ m (left) or 100  $\mu$ m (inset).

<span id="page-10-0"></span>![](_page_10_Figure_1.jpeg)

cycling progenitors (PgS and PgG2M), pinpointing a specific cell type and state likely associated with neural progenitor expansion [\(Figures 8](#page-15-1)B and 8C). By connecting causal genetic drivers to specific genes within a specific cell type, this not only identifies putative cell-type-specific mechanisms involved in cortical expansion, but also provides support for the radial unit hypothesis of cortical expansion on the human lineage [\(Lui et al., 2011; Rakic, 1995](#page-16-13)).

In contrast, common genetic variants influencing educational attainment (Edu) [\(Okbay et al., 2016\)](#page-17-13) were enriched in cycling neural progenitors, CP glutamatergic neurons, medial ganglionic eminence (MGE)-derived interneurons, and intriguingly, pericytes [\(Figures 8](#page-15-1)B and 8C). A less-powered IQ genomewide association study (GWAS) ([Sniekers et al., 2017](#page-17-14)) also found enrichment in maturing CP glutamatergic neurons but not in other cell types [\(Figures 8B](#page-15-1)). Unfortunately, most psychiatric disease GWAS remain underpowered (n =  $\sim$ 46,000 and  $\sim$ 34,000 for ASD and epilepsy, respectively). However, variants causing risk for schizophrenia (n =  $\sim$ 105,000) (Pardiñ[as et al., 2018](#page-17-15)) were enriched in multiple cell types, including neural progenitors, glutamatergic neurons, interneurons, OPCs, and microglia [\(Figures](#page-15-1) [8B](#page-15-1) and 8C). One study, using a partitioned heritability approach, found enrichment for schizophrenia variants in adult cortical glutamatergic neurons and cortical interneurons, consistent with bulk tissue analysis (Horvá[th and Mirnics, 2015\)](#page-16-19), but was unable to assess enrichment in human fetal cortical cell types, given a lack of available data [\(Skene et al., 2018\)](#page-17-16). Our results implicate neural progenitors, OPCs, and fetal microglia in schizophrenia, highlighting the importance of generating single-cell resources from multiple periods and brain regions. Given the complex etiology and phenotypic diversity of schizophrenia, it may be expected that multiple cell types are affected. These results highlight how combining DNA accessibility profiling and singlecell sequencing can facilitate interpretation of the function of variants influencing brain structure and function.

#### **DISCUSSION**

This resource of transcriptomic profiles of 40,000 single cells in human fetal cortex demonstrates the utility of single-cell analysis for characterizing human neurogenesis, identifying novel cell-type regulatory mechanisms, and understanding the cellular basis of brain phenotypes with neurodevelopmental origins. By expanding the publicly available number of human fetal brain single-cell transcriptomes by an order of magnitude, these data provide a high-resolution map of expression profiles for all known major cell types from mid-gestation human brains with more complete cell-type-specific mRNA transcript profiles than previously available. To facilitate sharing, exploration, and use of this unique and valuable resource, we developed a powerful and easy-to-use online browser that allows rapid queries to ascertain cell-type-specific expression patterns presented in an intuitive graphical interface. We leverage the breadth and robustness this high-depth catalog of neocortical cell types to characterize rare cell types and states, including PV interneurons, NPY-expressing SST interneurons, and SP neurons [\(Fan et al., 2018; Liu et al., 2016; Nowakowski et al.,](#page-16-4) [2017; Pollen et al., 2015; Zhong et al., 2018](#page-16-4)). We show that most markers for SP cells identified in other species are not specific to SP cells in humans and provide a new cadre of marker genes for this important cell class that has expanded substantially on the primate lineage [\(Hoerder-Suabedissen](#page-16-10) and Molnár, 2015).

Some rare cell states identified include transitional forms that we validate via *in situ* hybridization. Characterization of these rare cell states provides novel insight into neurodevelopmental cell dynamics. Specifically, our data implicate early decision points in cell fate trajectories that are pre-S phase, leading to transcriptomically mixed cell states before their division into two distinct cell types. An early cell fate decision point tied to cell cycle is consistent with previous work indicating cell fate decisions in neurogenesis are made in G1 ([Lange et al., 2009; Pilaz](#page-16-20) [et al., 2009\)](#page-16-20). However, previous models of asymmetric neurogenic divisions suggest that only a few key TFs of the daughter lineage are expressed in the asymmetrically dividing cell, whereas we observe early induction of more extensive cell-type transcriptional programs [\(Bertrand and Hobert, 2010; Pfeuty,](#page-16-21) [2015\)](#page-16-21). This is particularly surprising in that cells are expressing transcriptomes of two distinct cell types before telophase. In addition, the transition state dynamics during early neurogenesis show that cell-type differentiation is on a gradual continuum and involves transcriptomic transitions tied to cell-cycle progression, rather than off or on expression of a small group of TFs.

We also perform a systematic exploration of the impact of multiple technical factors, including comparisons across methods and to bulk tissue RNA-seq, enabling us to thoroughly evaluate gene and cell-type detection and coverage. These types of analyses

#### Figure 5. Transcriptional Network Discovery

<sup>(</sup>A–H) Regulatory elements for cell-type-specific genes. (A) Enhancer size by cell type. Enhancers are assigned to cell types by cell-type-enriched genes. (B) Density plot of enhancer sizes that are assigned to specific cell types. (C) Distance (base pairs) from enhancer to promoter by cell type. (D) Density plot of distance (base pairs) from enhancer end to promoter start that are assigned to specific cell types. (E) Enhancers per gene by cell type. (F) Histogram of enhancers per gene that are assigned to specific cell types. (G) Number of enhancers per gene versus coding sequence (CDS) length of the gene by cell type. (H) Number of enhancers per gene versus GC content of the gene by cell type.

<sup>(</sup>I) Schematic showing the computational approach used for transcriptional network discovery with the SCENIC pipeline (see [STAR Methods\)](#page-18-0). 1) Co-expression modules between transcription factors and candidate genes are constructed. 2) Genes in co-expression modules are then pruned to genes, which are inferred to be direct targets of the transcription factor, making a regulon. Direct targets are determined by the presence of the transcription factor binding motif in the regulatory elements associated with that gene. 3) The activity of each regulon is then assessed in each cell.

<sup>(</sup>J) Cell-type enrichment of regulon activity. Each regulon was scored as active or inactive for each cell, and cluster enrichment was then determined by Fisher's test. Color indicates false discovery rate (FDR)-corrected  $-\log_{10} p$  value.

<sup>(</sup>K) SCENIC regulon activity in each cell (AUCell) for the indicated TF plotted on tSNE.

<sup>(</sup>L) TFs with previously uncharacterized cell-type or cell-subtype-specific activity. Regulon activity in each cell (AUCell) for the indicated TF (top panels) or expression of the TF plotted on tSNE (bottom panels).

<span id="page-12-0"></span>![](_page_12_Figure_1.jpeg)

have not been performed in most published papers to date. By direct comparison with bulk tissue data, we show that there is a high level of correspondence between the transcriptomes identified in both single-cell and bulk tissue data. However, we find that single-cell transcriptomes miss a small number of certain genes, biased toward long neuronal-enriched transcripts such as cell adhesion molecules, an observation that has not been noted in previous studies [\(Nowakowski et al., 2017; Pollen et al., 2015;](#page-17-4) [Zhong et al., 2018\)](#page-17-4). In addition, we evaluate and provide high-confidence cell-type transcriptomes relative to previously published methods with lower throughput and higher sequencing depth. Finally, we provide the first direct comparison and integration of different human fetal brain single-cell datasets, demonstrating the reproducibility of these methods for identifying cell clusters.

By integrating these data with tissue-specific regulatory information, we provide a map of TF gene regulatory networks for specific cell types in developing human brain. We highlight how this can be used to identify critical cell types in monogenic disorders (e.g., ZFHX4 and 8q21.11 deletion), as well as in ASD, expanding the implicated cell landscape in this disorder to include inhibitory neurons and, in a few cases, non-neural cells, in addition to glutamatergic neurons. These results emphasize the importance of expanding single-cell taxonomies to include single-cell epigenetic analysis [\(Luo et al., 2017\)](#page-16-22). Lastly, we show that genes with human-specific expression patterns act preferentially in oRG and upper-layer cortical neurons, which is consistent with the expansion of these zones during brain evolution. These data provide a molecular context for cortical expansion and increased cortical-cortical connectivity in humans, and they extend our understanding of developmental dynamics and the origin of neuropsychiatric disease risk in human neocortex.

#### <span id="page-13-0"></span>STAR+METHODS

Detailed methods are provided in the online version of this paper and include the following:

- **O [KEY RESOURCES TABLE](#page-18-1)**
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	- $\circ$  Partitioned heritability analysis
	- $\circ$  Comparisons to adult brain single-nuclei expression profiles
	- $\circ$  Gene list enrichment analysis
- **. [DATA AND CODE AVAILABILITY](#page-25-0)**

#### SUPPLEMENTAL INFORMATION

Supplemental Information can be found online at [https://doi.org/10.1016/j.](https://doi.org/10.1016/j.neuron.2019.06.011) [neuron.2019.06.011](https://doi.org/10.1016/j.neuron.2019.06.011).

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Figure 6. Dissecting the Acquisition of a Neuronal Program

(A) tSNE colored by mean expression of cell-cycle phase markers.

(B) tSNE colored by co-expression of groups of canonical cell-type markers. Yellow indicates co-expression.

(C) Percentage of cells in each Seurat cluster displaying co-expression of major cell-type markers.

(D) Mixed transcriptomic signatures of mixed marker cells in S phase corresponding to the expression of markers from multiple cell types. For the RG-to-IP comparison, RG and IP eigengenes were derived from differentially expressed genes between RG and IP cells and similarly for the RG-to-Neuron comparison and IP-to-Neuron comparison. Boxplots: the boxes indicate first and third quartiles, the whiskers extend from the box to the highest or lowest value that is within 1.5 x interquartile range of the box, and the line is the median.

(E) Shared gene signatures between major cell types and mixed cell types. Overlap of gene signatures from major cell types (y axis) and genes differentially expressed between major cell types and mixed marker cells (labeled on the gray bar). x axis: percentage of genes differentially expressed between major cell types that are also differentially expressed between corresponding major cell type and mixed marker cells. For example,  $\sim$ 85% of IP signature genes are more highly expressed in RG+IP+ cells than in RG+ cells.

(F) RNA FISH of fetal cortex probed with the S-phase marker PCNA (green), the RG marker PAX6 (red), and the neuron marker STMN2 (magenta) and stained with DAPI (blue). Panels on the right show high-magnification single-plane confocal images of individual cells expressing all three markers. Scale bar, 100 µm (left) or  $10 \text{ µm}$  (right).

(G) Quantification of the percentage of cells co-expressing the S-phase marker PCNA, the RG marker PAX6, and the neuron marker STMN2. Mean ± SE displayed in barplot.

(H) Quantification of relative amounts of mitotic RG and relative amounts of IPs undergoing different differentiation events.

(I) Diagram of mixed cell-type transcriptomic states that is characteristic of neurogenic differentiation trajectories in human neocortex. \*p < 0.05, \*\*p < 0.01, \*\*\*p < 0.001, \*\*\*\*p < 0.0001.

<span id="page-14-0"></span>![](_page_14_Figure_1.jpeg)

#### Figure 7. Cellular Determinants of Disease

(A–C) Cell-type expression of ASD, epilepsy, or ID risk genes, respectively. Expression of ASD risk genes is enriched in fetal glutamatergic neurons, with some genes specifically expressed in other cell types. Red: gene is discussed in text. Cells are ordered by cluster.

(D) Cell-type enrichment of ASD, epilepsy, or ID risk genes. Numbers indicate the log<sub>2</sub> odds ratio; the red line indicates the FDR-significance threshold (p value 0.05).

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#### AUTHOR CONTRIBUTIONS

L.d.l.T.-U., J.L.S., and D.H.G. designed the study. D.H.G. oversaw all analyses and supervised the study, with help from K.P., M.B.G., and W.E.L. L.d.l.T.-U., J.L., S.N., M.B.G., C.K.O., and D.L. performed the single-cell

<span id="page-15-1"></span>![](_page_15_Figure_1.jpeg)

#### Figure 8. Partitioned Heritability Analysis Demonstrates Enrichment of Heritability in Specific Brain Traits and Neuropsychiatric Diseases in Diverse Cell Types

(A) Schematic showing the approach to identifying regulatory elements (REs) for specific cell types and assessing enrichment for specific brain traits. REs of genes enriched in specific cell types are identified by chromatin accessibility correlation between the promoter of the gene and the other accessible peaks within 1 Mb. The set of promoter and distal RE peaks are then tested for enrichment in SNPs associated with brain traits and neuropsychiatric disease using partitioned heritability by LD score regression.

(B) Heatmap showing significant partitioned heritability enrichment for specific brain traits and neuropsychiatric disorders in different cell populations. Color indicates the partitioned heritability enrichment. Numbers are the FDR-corrected p values. References for each GWAS are in [Table S8.](#page-13-0) We did not observe enrichment of irritable bowel syndrome (IBD) or finger whorl variants in the regulatory elements of any cortical-derived cell types, supporting the cell-type specificity of gene regulation.

(C) For selected GWAS, bar plots indicate the FDR-corrected significance or the enrichment (right) of partitioned heritability. Red vertical line indicates the FDRsignificance threshold (p value 0.05). Error bars represent SE. N, GWAS sample size.

L.d.l.T.-U., A.G.E., X.S., and W.C. analyzed the data, with input from S.S. L.d.l.T.-U., D.L., and C.K.V. performed the validation experiments. T.H. and J.K.L. performed copy number variation analyses. D.P., L.d.l.T.-U., and D.H.G. wrote and prepared the manuscript, with feedback from all authors.

#### SUPPORTING CITATIONS

The following references appear in the Supplemental Information: [Gawad](#page-16-23) [et al. \(2016\)](#page-16-23).

#### DECLARATION OF INTERESTS

The authors declare no competing interests.

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### <span id="page-18-0"></span>STAR+METHODS

### <span id="page-18-1"></span>KEY RESOURCES TABLE

![](_page_18_Picture_218.jpeg)

![](_page_19_Picture_285.jpeg)

(*Continued on next page*)

![](_page_20_Picture_198.jpeg)

#### <span id="page-20-0"></span>LEAD CONTACT AND MATERIALS AVAILABILITY

Further information and requests for reagents should be directed to and will be fulfilled by the Lead Contact, Daniel H. Geschwind ([dhg@mednet.ucla.edu\)](mailto:dhg@mednet.ucla.edu).

#### <span id="page-20-1"></span>EXPERIMENTAL MODEL AND SUBJECT DETAILS

#### Developing human brain tissue samples

De-identified fetal tissue samples were obtained from the UCLA Gene and Cell Therapy Core according to IRB guidelines or from the University of Maryland Brain and Tissue Bank (RNA FISH). For all donors profiled for single-cell RNA-seq (Drop-seq), DNA was acquired from fetal brain tissue and donors were genotyped with Illumina HumanOmni2.5 chips. Sex was determined based on homozygosity in X chromosome SNPs (3 male, 1 female) and expression of XIST and Y chromosome genes. High confidence CNVs were called with plumbCNV [\(Cooper et al., 2015](#page-16-28)). No known major pathogenic CNVs implicated in neuropsychiatric disorders were found in these donors. The largest CNV called and confirmed visually was 380kb. Samples processed for Drop-seq were obtained from 4 donors [female 17,17,18 gestation weeks (GW); male: 18 GW]. Samples processed for Fluidigm were obtained from 2 donors [female 17, 17.5 GW]. Samples processed for RNA fluorescent *in situ* hybridization (RNA FISH) were obtained from the UCLA Gene and Cell Therapy Core or from the University of Maryland Brain and Tissue Bank according to IRB guidelines from five donors aged GW15.5-18. This study was performed according to the legal and institutional ethical regulations of the UCLA Office of Human Research Protection. Full informed consent was obtained from all of the parent donors.

#### <span id="page-20-2"></span>METHOD DETAILS

#### Tissue dissection and single-cell isolation

Coronal sections were prepared from fetal cortices using a razor blade under a dissection microscope in ice-cold Hank's Balanced Salt Solution (HBSS). The coronal sections were then further dissected at the intermediate zone (IZ) to divide them into two regions: 1) consisting of the germinal zones (GZ) [ventricular zone (VZ), and subventricular zone (SVZ], and 2) consisting of the developing cortex (CP) [subplate (SP), cortical plate (CP), and marginal zone (MZ)]. The majority of the IZ was included as part of the "CP" dissection, but there is likely a small amount included in the GZ. Following dissection, GZ and CP sections were separately gently dissociated via enzymatic digestion with papain (Worthington) and filtered into a pure homogeneous cell suspension through dual filtering with a 40um strainer followed by an ovomucoid gradient (Worthington). Cell survival (90%–95%) and yield were quantified with Trypan blue staining, before immediately proceeding with Drop-seq or Fluidigm single-cell isolation. To assess doublet rates by human-mouse cell mixing experiments, mouse E15 cortical cultures were prepared in parallel. Briefly, mouse cortices were dissected in ice-cold HBSS and enzymatically dissociated with trypsin into a homogeneous cell suspension. Survival (90%–95%) and yield were quantified with Trypan blue staining. Mouse and human cells were mixed in a 1:10 ratio immediately prior to single-cell isolation by Drop-seq.

#### Single-cell RNA-seq

Drop-seq was run on single cells according to the online Drop-seq protocol v.3.1 (<http://mccarrolllab.com/download/905/>) and the methods published in [Macosko et al. \(2015\)](#page-17-2). Cells were maintained on ice and diluted to 125,000/mL in PBS + 0.01% BSA immediately prior to isolation. Barcoded beads were obtained from Chemgenes and cells were isolated in a Polydimethylsiloxane (PDMS) microfluidics device. Libraries were prepared with the Nextera XT DNA Library Preparation Kit (Illumina) according to the manufacturer's instructions. Libraries were then sequenced to an average of 57,814 reads/cell in an Illumina HiSeq2500 instrument with a modified 100bp paired-end protocol where R1 = 25bp and R2 = 75bp to maximize mapping. This read depth was empirically determined to yield the best per cell gene detection versus sequencing depth.

Fluidigm C1 scRNA-seq was run using the Fluidigm low-throughput small IFC (96 cells) or the high-throughput small IFC (800 cells) according to the manufacturer's instructions. Libraries were then prepared with the Nextera XT DNA Library Preparation Kit (Illumina) according to the manufacturer's instructions and sequenced to an average of 414,411 (high-throughput) and 682,569 (lowthroughput) reads/cell in an Illumina HiSeq2500 instrument with a modified 100bp paired-end protocol where R1 = 25bp and R2 = 75bp to maximize mapping. The low-throughput libraries were sequenced in an Illumina HiSeq2500 where R1 = 50bp and R2 = 50bp. The high-throughput libraries were sequenced in an Illumina HiSeq3000 where R1 = 12bp and R2 = 126bp.

#### RNA fluorescent in situ hybridization (RNA FISH)

In order to independently validate single-cell expression profiles, we used RNAscope ([Wang et al., 2012](#page-17-19)), an RNA FISH technique capable of single-molecule RNA detection with minimal off-target signal. Fetal tissue samples were obtained from the UCLA Gene and Cell Therapy Core or from the University of Maryland Brain and Tissue Bank according to IRB guidelines from five donors aged GW15.5-18. Developing human cortices were flash-frozen, embedded in OCT and cryosectioned in the coronal plane (15µm section thickness). Sections were then subjected to RNA FISH following the manufacturer's protocol for fresh frozen tissues using the Fluorescent Multiplex Assay Kit v1 (Advanced Cell Diagnostics Cat# 320850) with the following probes: CARHSP1-C1, CSRP2-C2, CRYAB (Cat# 426271-C2), EOMES (Cat# 429691-C3), LYN-C3, PAX6 (Cat# 588881-C1), PAX6 (Cat# 588881-C2), PCNA (Cat# 553071-C1), SATB2 (Cat# 420981-C1), ST18-C2, STMN2-C3, and ZFHX4-C2.

To determine the expression pattern across cortical layers for cell-enriched TFs [\(Figures 3](#page-6-0)C–3G and [4I](#page-8-0)), tiled images of multiple coronal sections from three independent donors spanning the entire cortex were acquired using a Leica DMi8 epifluorescence microscope at 40X magnification. Fluorescence analyses were performed in ImageJ version 2.0.0. For each image, regions of interest (ROIs) outlining individual cortical layers were manually created based on nuclear packing as visualized by DAPI staining. Background fluorescence was subtracted using a mask based on an empirically determined thresholded value for each image. To account for changes in cell density across cortical layers, the background-corrected signal for each channel was normalized to the DAPI intensity. The normalized fluorescence intensity is the mean gray value of each RNAFISH channel for each cortical layer divided by the mean DAPI gray value for the corresponding layer. DAPI-corrected values outside of three standard deviations of the mean were removed.

To quantify the co-expression of RG and neuronal markers in S-phase cells, images of coronal sections probed with the RG marker PAX6, the S-phase marker PCNA and the neuronal marker STMN2 were acquired on a Zeiss LSM780 confocal using a 63X magnification objective. Confocal tiled images of developing human cortex encompassing the VZ and SVZ were acquired for the entire thickness of the section at a 0.29µm Z-step size. For each cell quantified, the presence of puncta for each marker gene was determined across the entire Z-plane. Only cells expressing all three markers in the same Z-plane overlapping the DAPI staining were considered to be an RG-Neuron transition state. A total of 5,692 cells were quantified from two donors. Similar image acquisition and analysis was performed to quantify cells co-expressing PAX6 or EOMES, and ZFHX4. A total of 4,723 cells were quantified from three donors.

#### <span id="page-21-0"></span>QUANTIFICATION AND STATISTICAL ANALYSIS

#### Alignment and processing

The raw Drop-seq data was processed using the Drop-seq tools v1.12 pipeline from the McCarroll Laboratory [\(http://mccarrolllab.](http://mccarrolllab.org/wp-content/uploads/2016/03/Drop-seqAlignmentCookbookv1.2Jan2016.pdf) [org/wp-content/uploads/2016/03/Drop-seqAlignmentCookbookv1.2Jan2016.pdf\)](http://mccarrolllab.org/wp-content/uploads/2016/03/Drop-seqAlignmentCookbookv1.2Jan2016.pdf). Reads were aligned to the Ensembl release 87 *Homo sapiens*genome. We calculated unique molecular identifier (UMI) counts for each gene of each cell by collapsing UMI reads using Drop-seq tools.

The raw Fluidigm C1 data was processed using a custom pipeline. Cell barcode demultiplexing and initial processing was performed with Fluidigm mRNASeqHT\_demultiplex.pl v1.0.2. Raw reads were aligned to the Ensembl release 75 *Homo sapiens* genome with RNA STAR [\(Dobin et al., 2013](#page-16-26)). Aligned reads were sorted and alignments mapping to different chromosomes were removed from the BAM file using samtools [\(Li et al., 2009](#page-16-30)). Gene expression levels were quantified using HTSeq with a union exon model [\(Anders et al., 2015](#page-16-31)).

Quality control statistics were collected using RNA STAR statistics, Drop-seq tools metrics, and PicardTools (commands ReorderSam, CollectAlignmentSummaryMetrics, CollectRnaSeqMetrics, CollectGcBiasMetrics) and samtools (duplication metrics).

#### Assessment of doublet rate

Doublet rate, i.e., the frequency of which more than one cell was captured in a single Drop-seq droplet, was assessed by species mixing experiments [\(Figure S1;](#page-13-0) [Table S2\)](#page-13-0). Overall doublet rate (mouse+mouse, human+human, human+mouse) is derived based on the frequency of beads associating to both mouse and human cells in a single drop or well. For the species mixing experiments, cells from mouse E15 cortical cultures were added to human cells at a concentration of 1:10 as described above. The raw Drop-seq data was processed using the Drop-seq tools v1.12 pipeline from the McCarroll Laboratory ([http://mccarrolllab.org/wp-content/](http://mccarrolllab.org/wp-content/uploads/2016/03/Drop-seqAlignmentCookbookv1.2Jan2016.pdf) [uploads/2016/03/Drop-seqAlignmentCookbookv1.2Jan2016.pdf](http://mccarrolllab.org/wp-content/uploads/2016/03/Drop-seqAlignmentCookbookv1.2Jan2016.pdf)). Reads were aligned to a mixed species reference genome (*Homo sapiens* and *Mus musculus*) obtained from GEO GSE63269. The BAMs were then filtered into two organism specific BAMs using the Drop-seq tools command 'FilterBAM'. For each species-specific BAM, UMI counts were then calculated for each gene of each cell by collapsing UMI reads using Drop-seq tools.

#### Filtering and normalization

To select Drop-seq cells for downstream analysis: 1) Cells were selected for downstream analysis using the cell barcodes associated with the most UMIs. We estimated the number of cells captured as 5% of the input beads and retained this many cell barcodes for downstream analysis. 2) For samples with mouse cells spiked in, mouse cells were removed by filtering all cells with > 250 UMIs mapping to the mouse genome. 3) Removed cells with < 200 unique genes detected (gene detection:  $^3$ 1 count). 4) Removed cells with > 3

standard deviations above the mean number of genes detected (3152). 5) Removed cells with > 5% of their counts mapping to MT genes. 6) Removed genes detected in < 3 cells.

Normalization was performed using Seurat (v2.3.4; [Butler et al., 2018](#page-16-6)). Briefly, raw counts are read depth normalized by dividing by the total number of UMIs per cell, then multiplying by 10,000, adding a value of 1, and log transforming (ln (transcripts-per-10,000 + 1)) using the Seurat function 'CreateSeuratObject'. Raw UMI counts data were assessed for the effects from biological covariates (anatomical region, donor, age, sex), and technical covariates (library batch, sequencing batch, number of UMI, number of genes detected, CDS length, GC content) [\(Figure S1](#page-13-0)). The effects of number of UMI (sequencing depth), donor, and library preparation batch were removed using a linear model from the read depth normalized expression values (custom R scripts, lm(expression  $\sim$ number\_of\_UMI + donor + lab\_batch), and Seurat function 'ScaleData').

#### Single-cell clustering and visualization

Clustering was performed using Seurat (v2.3.4) [\(Butler et al., 2018](#page-16-6)). Read depth normalized expression values were mean centered and variance scaled for each gene, and the effects of number of UMI (sequencing depth), donor, and library preparation batch were removed using a linear model with Seurat ('ScaleData' function). Highly variable genes were then identified and used for the subsequent analysis (Seurat 'MeanVarPlot' function). Briefly, average expression and dispersion are calculated for each gene, genes are placed into bins, and then a z-score for dispersion within each bin is determined. Principal component analysis (PCA) was then used to reduce dimensionality of the dataset to the top 40 PCs (Seurat 'RunPCA' function). Clustering was then performed using graph based clustering implemented by Seurat ('FindClusters' function). Briefly, a K-nearest neighbor graph based on Euclidean distance in PCA space is constructed from the PC scores for each cell. Edges between cells are weighted based on shared overlap in neighborhoods determined by Jaccard distance. Cells are then iteratively grouped together with the goal of optimizing the density of links inside communities as compared to links between communities. Cell clusters with fewer than 30 cells were omitted from further analysis.

For visualization, t-distributed stochastic neighbor embedding (tSNE) coordinates were calculated in PCA space, independent of the clustering, using Seurat ('RunTSNE' function). tSNE plots were then colored by the cluster assignments derived above, gene expression values, or other features of interest. Gene expression values are mean centered and variance scaled unless otherwise noted.

For sub-clustering analysis an iterative approach was used [\(Figure 2\)](#page-4-0), cells from each initial cluster were re-processed, clustered, and analyzed from the raw counts matrix using Seurat as described above. For larger clusters (> 1,000 cells) the top 10 PCs were used, and for smaller clusters (< 1,000 cells) the top 5 PCs were used.

#### Cluster stability

To assess cluster stability, we adapted the approach from [Hennig \(2007\)](#page-16-32) using bootstrapping and the Jaccard index. Briefly, a bootstrap sample of cells is drawn with replacement from the original dataset, then re-quality filtered, normalized, analyzed, and clustered, then this process is repeated over 100 iterations. For each iteration, the maximum Jaccard index is computed between the new clustering and the original clustering. The mean Jaccard index of 100 iterations of bootstrapping is reported [\(Figure S1](#page-13-0)F). Jaccard Index values range from 0-1, with  $> 0.5$  indicating stable clustering.

#### Differential gene expression analysis and cell type enrichment

In general, differentially expressed genes between different cell groups were determined using a linear model implemented in R as follows: Im(expression ~number\_of\_UMI + donor + lab\_batch). P values were then Benjamini-Hochberg corrected. To identify cell type enriched genes, differential expression analysis was performed for each cluster individually versus all other cells in the dataset for genes detected in at least 10% of cells in the cluster. Genes were considered enriched if they were detected in at least 10% of cells in the cluster,  $0.2 \log_2 6$  denriched, and Benjamini-Hochberg corrected p value  $< 0.05$  [\(Table S6\)](#page-13-0).

#### Pseudo-time analysis

Monocle 2 was used to construct single-cell pseudo-time trajectories ([Qiu et al., 2017; Trapnell et al., 2014\)](#page-17-3). First, the dataset was subset to cells in Seurat clusters inferred to be part of the neurogenesis differentiation axis (progenitor and excitatory neuron clusters) ([Figure 1](#page-2-0)I). The subset dataset was then run through the Monocle 2 pipeline beginning with raw counts. Dispersed genes to use for pseudo-time ordering were calculated using the 'estimateDispersions' function and required to be expressed in at least 10 cells. DDRTree was used to reduce dimensions and the effects of number of UMI (sequencing depth), donor, and library preparation batch were corrected for (Monocle function: reduceDimension(mo\_filtered, max\_components = 30, residualModelFormulaStr = ''number\_of\_UMI+donor+lab\_batch''). The visualization function 'plot\_cell\_trajectory' was used to plot the minimum spanning tree on cells.

#### Stability of cluster gene expression signatures

Cluster gene expression signatures were evaluated by the stability in mean gene expression level ranking. Gene expression ranking was determined by mean expression level for each gene across all cells in the cluster. A bootstrapping approach was then used to evaluate stability of the gene expression rankings. Cells from each respective cluster were sampled with replacement over

1,000 iterations. At each iteration, a sample population was drawn pseudo-randomly and the mean expression level across the population was calculated and then genes were ranked by mean expression level, with a ranking of 1 being the most highly expressed. The mean of the rankings over all iterations and the standard deviation were plotted ([Figures S2](#page-13-0)G, [S5B](#page-13-0), and S5C).

#### Alignment of single-cell datasets

Datasets were aligned using Seurat canonical correlation analysis ([Butler et al., 2018](#page-16-6)). First, read depth normalized expression values were mean centered and variance scaled for each gene. Then the 2,000 most highly variable genes were identified for each dataset using the Seurat 'FindVariableGenes' function. Next, canonical correlation analysis was run using the union of the variable gene sets as described in [Butler et al. \(2018\).](#page-16-6) The analysis returns canonical correlation vectors (CCV) across both datasets, which are then used to align the datasets. We used the top 20 CCVs to run the alignment procedure. The datasets were then aligned using the Seurat 'AlignSubspace' function, which utilizes CCVs and a nonlinear time warping algorithm to align metagenes between datasets. We then applied t-SNE to reduce dimensionality to plot the aligned datasets.

#### Comparison to bulk tissue RNA-seq

Bulk tissue RNA-seq samples from human fetal neocortex (GW17-19) GZ (n = 9) and CP (n = 9) were obtained from [de la Torre-Ubieta](#page-16-8) [et al. \(2018\).](#page-16-8) GZ and CP dissections were carried out as described above. Bulk tissue RNA-seq samples were read depth normalized to counts per million (CPM). For comparison to single-cell RNA-seq data from a different laboratory, we obtained Fluidigm C1 gener-ated raw sequencing data from human fetal brain from [Pollen et al. \(2015\).](#page-17-20) For all single-cell datasets, single-cell expression profiles were pooled by aggregating gene expression counts across groups of cells to simulate bulk tissue RNA-seq samples. To aggregate or pool gene expression counts, groups of cells were randomly drawn from the single-cell dataset, and raw counts were summed across each group of cells for each gene to pool the expression profiles. Each pooled expression profile was then read depth normalized to CPM.

To compare pooled samples of different sizes to bulk tissue RNA-seq by correlation in gene expression values, 1) Samples of different numbers of cells were drawn, and counts were summed across the cells for each gene to make pooled samples of single-cells. 2) Gene expression levels (summed counts) of the pooled single-cell datasets were then correlated to the mean CPM of the bulk tissue RNA-seq dataset [\(Figure S4](#page-13-0)B).

To identify genes under-represented in single-cell RNA-seq compared to bulk tissue RNA-seq, we identified genes with higher or lower relative expression in the pooled single-cell expression profiles compared to bulk tissue RNA-seq. Genes greater than two standard deviations from the mean relative expression level of pooled versus bulk were labeled as under or over-represented in the respective single-cell RNA-seq dataset. To assess biases in capture of different cell types with Drop-seq, the expression of groups of cell type marker genes in the pooled Drop-seq dataset were compared to expression in the bulk tissue RNA-seq dataset. The expression ratios of pooled Drop-seq versus bulk tissue RNA-seq were converted to a z-score for plotting [\(Figure S4I](#page-13-0)).

Gene Ontology enrichment analysis was performed using g:Profiler [\(Reimand et al., 2016](#page-17-18)).

#### Cell type enrichment of TFs and co-factors

TFs, co-factors, and chromatin remodelers were obtained from AnimalTFDB 2.0 ([Zhang et al., 2015\)](#page-17-17). Genes were considered enriched in a major cell type if they were  $> 0.4$  log<sub>2</sub>fold enriched for any cluster corresponding to cells of that type, and were  $< 0.25$ log<sub>2</sub>fold enriched for any other cluster [\(Figure 3A](#page-6-0)). For example, radial glia (RG) enriched genes are > 0.4 log<sub>2</sub>fold enriched in either or both the ventricular radial glia (vRG) and outer radial glia (oRG) cluster, and < 0.25 log<sub>2</sub>fold enriched in any other cluster.

#### Subplate markers

To derive a human SP set of markers across mid-gestation, we used the fetal LCM laminae dataset ([Miller et al., 2014](#page-17-7)) to identify SP enriched genes ([Figures 4](#page-8-0)B–4D). Genes were sorted by fold change of SP versus the VZ, IZ, CP, and MZ, using the Brainspan online tool (<http://www.brainspan.org/lcm/search/index.html>), and then manually curated for SP specificity.

#### Cell cycle analysis

Cell cycle state was determined by mean expression of groups of cell cycle stage marker genes obtained from [Macosko et al. \(2015\)](#page-17-2) [\(Figures S6A](#page-13-0)–S6C). Two methods for cell cycle normalization were tested: 1) All cell cycle stage marker genes were excluded from the highly variable genes used for PCA, tSNE, and Seurat clustering described above. 2) Cell cycle correction by removing the effects of cell cycle state using a linear model via Seurat. First, each cell is assigned a S-phase and G2/M phase score using Seurat's 'Cell-CycleScoring' function. Then the cell cycle score is regressed out along with number of UMI (sequencing depth), donor, and library preparation batch using a linear model as described above.

#### Transition state analysis

Cells were considered positive for markers of a cell type if the mean expression of a group of cell type marker genes was > 0.5 log normalized expression, e.g., RG+Neuron+ cells express RG marker genes at a mean expression level > 0.5 log normalized expression and neuronal marker genes at a mean expression level > 0.5 log normalized expression.

Cell type gene signatures were determined using two methods: 1) Differential expression of cells in a type versus cells in another type, e.g.to determine an RG signature and newborn neuron signature differentiating the two cell types, differential expression of RG cells from RG clusters (oRG and vRG cluster) versus newborn migrating excitatory neurons (ExN cluster). 2) Enrichment of genes in a cell type. For each cluster, genes enriched in the cluster were determined as described above. Then the union of enriched genes for all clusters in a cell type was taken. The RG signature is the union of genes enriched in vRG and oRG clusters, the intermediate progenitor (IP) signature is genes enriched in the IP cluster, and the Neuron signature is genes enriched in the migrating excitatory neuron cluster.

Transcriptomic analysis of cycling mixed marker cells used cells dual positive for RG, IP, or neuronal markers specifically in the Sphase or G2/M phase clusters, e.g., RG+Neuron+ S-phase cells are cells from the S-phase cluster and dual positive for RG and neuronal markers. Dual positive cells were then compared to single marker type positive cells, e.g., RG+Neuron+ S-phase cells were compared to RG+ Neuron- IP- and RG- Neuron+ IP- cells to ensure comparison to cells of a clear transcriptomic type. The eigengene of the cell type signatures was then calculated for each cell positive for cell type markers in an expected differentiation trajectory, e.g., to explore the RG to neuronal transition the Neuron eigengene was calculated using the neuron signature across RG+ Neuron- IP- negative cells, RG+ Neuron+ cells, and RG- Neuron+ IP- cells. Cell type signatures were determined as described above.

The amount of overlap and the magnitude of expression of the cell type gene signature of cycling mixed marker cells was compared to the end point cell type. The gene signatures of the cell types involved in a differentiation trajectory were compared by fold change of the gene signature in the beginning state cell type to the endpoint cell type. For example, for the RG to neuron transition, the RG and neuronal gene signatures are determined by differential expression of cells from the RG to the newborn neuronal cluster. The expression of the neuronal gene signature genes is then compared in RG+ cells to RG+Neuron+ cells. The percent of the neuronal genes that are more highly expressed in RG+Neuron+ cells versus RG+ cells is ascertained, this is the percent of shared genes. The mean fold change of the neuronal genes in RG+Neuron+ cells versus RG+ cells is also determined and compared to the magnitude of the mean fold change in RG+ versus Neuron+ cells, this is the percent of fold change.

#### Cell-type specific regulatory elements

A map of regulatory elements active in developing fetal cortex generated from chromatin accessibility data (ATAC-seq) ([Buenrostro](#page-16-33) [et al., 2013\)](#page-16-33) was obtained from [\(de la Torre-Ubieta et al. \(2018\)](#page-16-8). Promoter elements were identified as accessible chromatin peaks within annotated gene promoters (within 2kb upstream and 1kb downstream of the transcription start site). Distal regulatory elements were then linked to genes by correlation between the promoter accessible peak and distal ATAC-seq peaks as described [\(de la](#page-16-8) [Torre-Ubieta et al., 2018\)](#page-16-8). Cell-type specific regulatory elements comprise the union of accessible chromatin within the promoter of a given gene and associated distal regulatory elements (enhancers) for each set of genes enriched within specific cell types. Regulatory elements associated with genes enriched in specific cell types were then used for regulatory element metrics ([Figures 5](#page-10-0)A–5H) and partitioned heritability analyses ([Figure 8\)](#page-15-1).

#### Gene regulatory networks

Cell-type specific regulatory networks were identified based on genes enriched in cell clusters using the SCENIC pipeline ([Aibar et al.,](#page-15-0) [2017](#page-15-0)). First, co-expressed modules between TFs and genes were identified from the single-cell expression data using GRNBoost. The set of TFs used for the co-expression analysis consisted of 782 well known TFs from RcisTarget database. In total, 2,862,989 TF and gene linkages were identified. To obtain more reliable TF and gene linkages, the top 10% of the linkages with highest scores were kept for further analysis. Then the regulatory elements for each module were extracted from the active regulatory elements obtained from [de la Torre-Ubieta et al. \(2018\)](#page-16-8). A motif enrichment analysis was done for the regulatory elements using Homer. If the co-expressed TF has a motif enriched in the regulatory elements (p value < 0.01), it will be the regulon for this gene module. The motif database consists of known motifs from Homer ([Heinz et al., 2010](#page-16-27)) and novel motifs from JASPAR ([Khan et al., 2018\)](#page-16-24). The JASPAR motifs were formatted by MEME [\(Bailey et al., 2009](#page-16-29)). Finally, the regulon activity (AUC score) for each module in each cell was scored by AUCell ([Aibar et al., 2017\)](#page-15-0). Given the distribution of activity scores, the cells that have the regulon enriched were identified. A Fisher's exact test was performed to evaluate if a regulon is significantly enriched (p value < 0.05) in specific cell types based on the enriched cell clusters [\(Table S7\)](#page-13-0).

#### Partitioned heritability analysis

Partitioned heritability was assessed using LD score regression (v1.0.0) ([Finucane et al., 2015](#page-16-18)). Heritability was calculated by comparing the association statistics for common genetic variants falling within regulatory elements associated with specific cell types, with the LD-score, a measure of the extent of the LD block. First, an annotation file was created, which marked all HapMap3 SNPs that fell within the regulatory elements for each cell type. LD-scores were calculated for these SNPs within 1 cM windows using the 1000 Genomes EUR data. These LD-scores were included simultaneously with the baseline distributed annotation file from [Finucane et al. \(2015\)](#page-16-18). Subsequently, the heritability explained by these annotated regions of the genome was assessed from phenotypes for 18 GWAS (see [Table S8](#page-13-0) for references and sample sizes). The enrichment was calculated as the heritability explained for each phenotype within a given annotation divided by the proportion of SNPs in the genome and FDR correction within each GWAS was used to correct for multiple comparisons.

#### Comparisons to adult brain single-nuclei expression profiles

For comparison to single-nuclei RNA-seq data from human adult brain we obtained the gene expression counts matrix and cluster enrichment scores from [Lake et al. \(2018\)](#page-16-25) (GEO: GSE92942). The single-nuclei RNA-seq raw counts were read depth normalized by dividing by the total number of UMIs per cell, then multiplying by 10,000, adding a value of 1, and log transforming (ln(transcripts-per- $10,000 + 1$ ).

#### Gene list enrichment analysis

Genes with human specific expression patterns across cortical development were obtained from [Bakken et al. \(2016\).](#page-16-12) High confidence ASD risk genes, defined by harboring high risk likely protein-disrupting mutations, were obtained from Sanders et al. combined *de novo* and TADA analysis (66 genes FDR < 0.1) [\(Sanders et al., 2015\)](#page-17-12). Intellectual disability (ID) risk genes were obtained from previous exome sequencing of patients with idiopathic and non-syndromic ([de Ligt et al., 2012; Rauch et al., 2012\)](#page-16-34) and subset to mutations presenting *de novo* in patients and likely to be gene-disrupting (frameshift, nonsense or splicing sites).

In order to obtain a list of high-confidence epilepsy risk genes, we curated ClinVar and OMIM (access date: May 2018). We first searched ClinVar for ''epilepsy'' associated variants which resulted in 7,011 variant-phenotype entries. Variants identified as having a clinical significance of uncertain, benign, likely benign, not provided, or drug response were removed while variants with a clinical significance of risk factor, likely pathogenic, pathogenic, or conflicting interpretations of pathogenicity were retained. Large structural variants disrupting more than one gene were also removed. This resulted in 1,227 entries with an identical gene-condition-clinical significance (> = 1 variant listed per gene). Genes lacking a single variant with a clinical significance of pathogenic were removed and the remaining entries were then manually curated using OMIM. OMIM genes for which the molecular basis was known were then evaluated for the strength of evidence from the reported clinical features and the molecular genetics. If the clinical features revealed the patients never met the criteria for epilepsy (*e.g*., two or more unprovoked seizures) or if seizures were a variable clinical feature rather than a defining feature of the syndrome these genes were excluded from the high-confidence list. Similarly, if the molecular genetic evidence was not sufficient (e.g., not a large enough sample size) these genes were excluded from the highconfidence list. Finally, we cross checked that all high-confidence genes from OMIM's Phenotypic Series for Epileptic encephalopathy early infantile were included on our list, resulting in the addition of 14 genes. This resulted in a final list of 109 high-confidence epilepsy risk genes.

Enrichment log<sub>2</sub>odds ratios were calculated using a general linear model (binomial distribution).

#### <span id="page-25-0"></span>DATA AND CODE AVAILABILITY

The accession number for the transcriptomic dataset reported in this paper is dbGaP: phs001836.

An online interface to facilitate sharing, exploration, and use of the dataset can be found at [http://geschwindlab.dgsom.ucla.edu/](http://geschwindlab.dgsom.ucla.edu/pages/codexviewer) [pages/codexviewer](http://geschwindlab.dgsom.ucla.edu/pages/codexviewer).