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Journal

The Scientific monthly, 384(6698)

Authors

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Publication Date

2024-05-24

DOI

10.1126/science.adh1938

Peer reviewed



HHS Public Access

Author manuscript

Science. Author manuscript; available in PMC 2024 September 13.

Published in final edited form as:

Science. 2024 May 24; 384(6698): eadh1938. doi:10.1126/science.adh1938.

A data-driven single cell and spatial transcriptomic map of the human prefrontal cortex

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Competing Interests

AB was a consultant for Third Rock Ventures and a shareholder in Alphabet, Inc. Joel E. Kleinman is a consultant on a Data Monitoring Committee for an antipsychotic drug trial for Merck & Co., Inc.

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Abstract

The molecular organization of the human neocortex has been historically studied in the context of its histological layers. However, emerging spatial transcriptomic technologies have enabled unbiased identification of transcriptionally-defined spatial domains that move beyond classic cytoarchitecture. Here we used the Visium spatial gene expression platform to generate a data-driven molecular neuroanatomical atlas across the anterior-posterior axis of the human dorsolateral prefrontal cortex (DLPFC). Integration with paired single nucleus RNA-sequencing data revealed distinct cell type compositions and cell-cell interactions across spatial domains. Using PsychENCODE and publicly available data, we map the enrichment of cell types and genes associated with neuropsychiatric disorders to discrete spatial domains.

Summary:

Generation of a molecular neuroanatomical map of the human prefrontal cortex reveals features relevant for psychiatric disease.

The emergence of single cell and spatially-resolved transcriptomics has facilitated the generation of integrated molecular anatomical maps across a variety of tissues in both rodents and humans (1–4). Data-driven unsupervised approaches to identify spatial domains in these large datasets, particularly in the rodent brain, have refined our understanding of the spatial organization of tissues beyond classical cytoarchitectural boundaries (2), and computational models have helped reveal new insights into cellular diversification during development (5). However, efforts to generate spatially-resolved transcriptomics data in the human brain at the scale and size necessary to employ these approaches with sufficient statistical power, have lagged behind.

We previously characterized the spatial topography of gene expression in the human dorsolateral prefrontal cortex (DLPFC), a brain region associated with several neurodevelopmental and psychiatric disorders (6–11), by manually annotating the six histological layers and white matter of the neocortex in a small cohort of 3 neurotypical adult donors (12). Although we identified robust layer-enriched gene expression with this

approach, recent computational tools for unsupervised clustering (13–17) and cell to cell communication (18) combined with the ability to expand the generation of molecular neuroanatomical maps across a larger donor pool has enabled data-driven identification of higher resolution spatial domains. Applying these data-driven approaches to larger-scale studies might facilitate the demarcation of fine cortical sublayers, which currently lack molecular annotations. They also could enable identification of non-laminar spatial domains associated with anatomical or topographical features in the human brain, including vasculature. Here we sought to develop and validate a framework for application of unsupervised spatial clustering approaches in human brain tissue across different anterior-posterior positions of the DLPFC and generate an analytical roadmap that can be extended to other brain areas with less well-characterized neuroanatomical architecture in the future.

Recent single nucleus RNA-sequencing (snRNA-seq) analyses are defining transcriptionally distinct DLPFC cell types and revealing cell type-specific changes associated with neurodevelopmental and neuropsychiatric disorders, such as schizophrenia (SCZ), autism spectrum disorder (ASD), major depressive disorder (MDD) and post-traumatic stress disorder (PTSD) (19–23). However, snRNA-seq data lack spatial context, which when retained during molecular profiling, can provide important insights into cell-cell communication and disease pathogenesis. To facilitate this type of analysis, we generated large-scale, unsupervised spatial transcriptomic molecular maps of the human DLPFC from ten neurotypical brain donors, which we integrated with snRNA-seq data across a variety of brain disorders. This spatially-resolved, molecular atlas of gene expression architecture in the human brain is provided as an interactive data resource for the scientific community to help reveal molecular mechanisms associated with psychiatric illness.

Study Overview

Here we applied single cell and spatial transcriptomic approaches to create a large-scale, data-driven spatial map of gene expression at single cell resolution in the adult human DLPFC in order to identify spatial domains, define cell-cell communication (CCC) patterns, and perform spatial registration of cell types across brain disorders (Fig. 1A-B). Using the Visium spatial transcriptomics platform (24), we measured spatial gene expression in fresh frozen postmortem human tissue blocks from 10 neurotypical adult donors (Table S1) in three positions spanning the rostral-caudal axis of the DLPFC (anterior [Ant], middle [Mid] and posterior [Post]) for a total of 30 tissue sections [n=10 per position] (Fig. 1A). In parallel, we performed snRNA-seq (10x Genomics 3' gene expression) on a subset of the same DLPFC samples (n=1-2 blocks per donor) to generate matched snRNA-seq and spatial transcriptomic data for 19 tissue blocks (Fig. 1A-C). To preserve Layer (L)1, blocks were microdissected across sulci in the plane perpendicular to the pia that extended to the graywhite matter junction. The morphology of each tissue block was assessed with RNAscope multiplex single molecule fluorescent in situ hybridization (smFISH) using regional and laminar marker genes to ensure dissection consistency (Fig. 1D). Sample orientation was confirmed by expression of genes enriched in the gray matter (SNAP25), white matter (WM; [MBP]), and L5 (PCP4; Fig. 1C, Fig S2, Fig S3, Fig S4). For Visium, 4,866 (4.1%) spots with low library size were excluded (Fig S5), resulting in a total of 113,927 spots across 30 tissue blocks and 10 donors. Downstream analyses at the gene-level (Fig S6) and

spot-level (Fig S7) were not impacted by tissue artifacts, including wrinkles, shears, and folds (see Methods: Evaluating the impact of histology artifacts on Visium H&E data). For snRNA-seq, 54,394 nuclei across 19 tissue blocks from 10 donors passed quality control and were included in the study (Fig. 1C). Using these integrated datasets, we performed several analyses including unsupervised clustering, spot deconvolution, CCC analyses, and spatial registration of PsychENCODE single cell datasets (Fig. 1B).

Identification of data-driven spatial domains at different resolutions across DLPFC

We selected an unsupervised clustering method for robust identification of laminar spatial domains (SpDs) in Visium data by benchmarking three algorithms, graph-based clustering, SpaGGN and BayesSpace (13, 15, 25, 26), using our previously published DLPFC Visium data (12 sections from 3 donors; (12)). We compared data-driven clustering accuracy against manual layer annotations (Fig S9, Fig S10). Among the algorithms tested, BayesSpace most accurately identified spatial domains (SpDs) consistent with the histological cortical layers. Therefore, we used this clustering algorithm to identify 7 unsupervised SpDs approximating the 6 cortical layers and WM. To relate unsupervised SpDs to the classic histological layers, we pseudo-bulked spots (by gene, sum the expression for all spots in the same SpD in a given sample) within each SpD across individual tissue sections to generate SpDspecific expression profiles and performed differential expression (DE) analysis to identify genes enriched in each SpD. Next, we performed "spatial registration" by correlating the enrichment statistics computed on BayesSpace SpDs with those from manually annotated cortical layers (12) to approximate the most strongly associated histological layer for a given BayesSpace SpD (Fig. 1E). We annotated the association of a specific spatial domain (SpD) at cluster resolution k to a classic histological layer using the term $\operatorname{Sp}_k D_d \sim L$, where L refers to the histological layer most strongly associated to domain d following cluster registration at resolution k. For example, spatial domain 7 at cluster resolution k=7 (Sp₇D₇) was mostly strongly associated with white matter (Fig. 1E) and is annotated as Sp₇D₇~WM. We found that k=7 was not sufficient to fully separate histological layers, especially superficial L2-L4, suggesting the presence of higher resolution data-driven SpDs and highlighting the challenges of manually annotating spatial domains (12).

We next evaluated how increasing cluster resolution (*k*) influenced the identification and cluster registration of unsupervised SpDs (Fig. 2). As expected, clustering at *k*=2 reliably separated white and gray matter (Fig S11, Fig S12). Next, we evaluated three clustering resolutions: a broad resolution *k*=9, a data-driven fine resolution *k*=16 (Fig S13, Fig S14), and a super-fine resolution *k*=28. Hereafter, we refer to these SpDs as Sp₉D, Sp₁₆D, and Sp₂₈D, respectively. Broad clustering most accurately recapitulated the classic histological layers with clear separation of Sp₉Ds enriched in genes expressed in L1–6 and WM (Fig. 2A–B, Fig S15). At fine clustering resolution, SpDs were largely laminar with two or more Sp₁₆Ds registering to a given histological layer, suggesting the presence of molecularly-defined sublayers (Fig. 2A–B, Fig S16). At super-fine resolution, many SpDs lacked a laminar structure, but spots belonging to Sp₂₈Ds frequently mapped back to a single broad or fine resolution spatial domain (Fig S17, Fig S18). To evaluate the unsupervised

architecture of cortical layers in the human DLPFC in more depth, we focused on broad and fine SpDs based on the robust laminar features of these domains.

Enrichment of differentially expressed genes in unsupervised spatial domains

To identify the molecular signatures of SpDs within each clustering resolution, we next performed differential expression (DE) analysis with linear mixed-effects modeling using the Sp₉D- or Sp₁₆D-specific expression profiles. As previously described, we employed three different statistical models: ANOVA model, enrichment model, and pairwise model (12). As expected, all three models confirmed that unsupervised clustering at broad or fine resolution identified biologically meaningful SpDs with significantly DE of genes across the laminar architecture of the cortex (5,931 FDR<5% unique enriched genes in at least one Sp₉D, see Methods: Layer-level data processing and differential expression modeling). Although we did identify 512 unique genes that were differentially enriched across the anterior-posterior axis of the DLPFC, SpD had a much stronger effect on gene expression compared to anatomical position (anterior, middle, posterior, Fig S19).

Analysis of DEGs identified using the enrichment model allowed for characterization of previously unrecognized data-driven SpDs, such as Sp₉D₁ and Sp₁₆D₁ (Fig. 2B). At both k=9 and k=16, domain 1 is adjacent to histological L1 and enriched for genes associated with blood vessels and brain vasculature, such as *CLDN5* (p=2.04e-75; Fig. 2C-D). Due to its thinness, this vascular-rich meninges layer was not manually annotated in our previous study (12), demonstrating the utility of unsupervised approaches to robustly identify biologically meaningful SpDs. Data-driven clustering at fine resolution also revealed molecularly-defined sublayers, including two adjacent laminar spatial subdomains (Sp₁₆D₁₄~L1 and Sp₁₆D₂~L1; Fig. 2E) enriched in L1 marker genes, including *RELN* (p=6.98e-17, p=3.19e-19 respectively) and AQP4 (p=9.37e-21, p=9.43e-12 respectively). Pairwise tests across these L1-related subdomains highlighted differential expression of SPARC (enriched in $Sp_{16}D_{14}\sim L1$, p=7.61e-14) and HTRA1 (enriched in $Sp_{16}D_{2}\sim L1$, p=1.26e-07), and principal component analysis (PCA) further confirmed the unique nature of these molecularly-defined sublayers (Fig. 2E-G). We also identified multiple SpDs associated with histological L4 (Sp₁₆D₅-L4 and Sp₁₆D₉-L4), L5 (Sp₁₆D₄-L5 and Sp₁₆D₁₆-L5), and L6 (Sp₁₆D₇-L6 and Sp₁₆D₁₂-L6; Fig. 2B). Taken together, this analysis validated the biological relevance of data-driven SpDs and increased the understanding of the molecular neuroanatomy across the laminar architecture of the adult human DLPFC.

Identification of molecularly and spatially distinct neuronal populations across cortical layers

To add single cell resolution to our molecular maps, we performed snRNA-seq on a subset of the same tissue blocks used for Visium (Fig. 1). Following assessment of quality control metrics (Fig S20, Fig S21), we performed batch correction (Fig S22, Fig S23, Fig S24) and data-driven clustering to generate 29 fine-resolution clusters across 7 broad cell types represented throughout the anterior-posterior DLPFC axis (Fig. 3A, Fig S25). Further, we

spatially registered all clusters to the histological layers using manually annotated Visium data from previous work (12) (Fig. 3B). Given that molecularly-defined excitatory neuron populations in the cortex have distinct laminar identities (27), we systematically assigned a histological layer to excitatory neuron clusters (Table S2), resulting in the identification of 13 layer-resolution clusters with distinct marker genes (Fig. 3B–D, Fig S26). At both fine-resolution and layer-resolution, our clusters strongly correlated with those derived from the reference-based mapping tool, *Azimuth* (https://azimuth.hubmapconsortium.org) (28, 29) (Fig S27).

To gain further insight into the relationship between our snRNA-seq clusters and Visium unsupervised SpDs, we performed spatial registration of fine resolution snRNA-seq clusters with Visium SpDs at k=9 and k=16 (Fig. 3B). We refined the spatial positioning of our 29 fine resolution snRNA-seq clusters and validated the laminar associations of broad and fine unsupervised SpDs. For example, we showed that snRNA-seq excitatory neuron clusters Excit_06 and Excit_08, which spatially registered to histological L6, were also highly correlated with Visium Sp₉D₇, a domain enriched for L6-associated genes (Fig. 2B). These snRNA-seq clusters were also highly correlated with a single spatial domain (Sp₁₆D₇), thereby further refining their anatomical position to upper L6. Inhibitory GABAergic populations were also assigned to specific spatial locations. For example, Inhib_05 uniquely registered to histological L2, which was confirmed with strong correlations to Sp₉D₃ and Sp₁₆D₈ showing enrichment for L2-associated genes (Fig. 2B). We also showed registration of snRNA-seq endothelial cell populations to vascular spatial domains (Sp₉D₁ and Sp₁₆D₁) enriched in HBA1 and CLDN5. Whereas endothelial cells showed the strongest correlation with vascular-associated SpDs and glial cells showed the strongest correlation with L1 and WM-associated SpDs (Fig. 2B), these cell types are indeed distributed across the cortical layers necessitating higher resolution spatial mapping approaches, such as spot deconvolution.

Defining cell type composition of unsupervised spatial domains using spot deconvolution

Given that individual Visium spots in the human DLPFC contain an average of 3 cells per spot (12), we used our paired snRNA-seq data to perform cellular deconvolution of Visium spots to better understand the cell type composition of unsupervised SpDs. First, we benchmarked 3 spot-level deconvolution algorithms, *SPOTlight, Tangram*, and *Cell2location* (14, 30, 31), using a gold standard reference dataset acquired with the Visium Spatial Proteogenomics (Visium-SPG) assay. Visium-SPG replaces H&E histology with immunofluorescence staining, enabling us to label and quantify 4 broad cell types across the DLPFC, including NeuN (neurons), OLIG2 (oligodendrocytes), GFAP (astrocytes), and TMEM119 (microglia) (Fig. 4A, Fig S28). After verifying marker genes for each snRNA-seq cluster (Fig S29) and confirming the utility of these genes for spot deconvolution (Fig. 4B, Fig S30, Fig S31, Fig S32), we applied *SPOTlight, Tangram*, and *Cell2location* to our Visium-SPG data and calculated the predicted cell type counts per spot at broad and fine resolution (Fig. 4B, Fig S33).

To quantify algorithm performance, we took 2 complementary approaches: 1) evaluating the localization of laminar cell types to their expected cortical layer (Fig. 4C, Fig S34, Fig S35) and 2) comparing predicted cell type counts to those obtained from immunofluorescent images using a classification and regression tree (CART) strategy to categorize nuclei into the 4 immunolabeled cell types (Fig. 4D, Fig S36, Fig S37, Fig S38, Fig S39). Using the first approach, we found that Tangram and Cell2location performed best across all cell types, but SPOTlight failed to accurately map excitatory neuron subtypes to the correct layer (Fig. 4B). Using the second approach, we found that the predicted counts from Tangram and Cell2location also had the highest correlation to CART-calculated counts (Fig. 4D) and Tangram showed the most consistent performance at both broad and layer level resolution across all cell types and samples (Fig. 4E). Finally, we applied *Tangram* and *Cell2location* to our H&E Visium dataset to predict the cellular composition of SpDs across the anteriorposterior axis of the DLPFC (Fig. 4F-G, Fig S40, Fig S41) and found that *Tangram* and Cell2location showed differences in the predicted cell counts per spot (Fig. 4F) whereas, for both tools, the predicted cellular composition of SpDs was consistent across samples at both broad (k=9) and fine (k=16) resolution regardless of DLPFC position (anterior, middle, posterior).

Spatial mapping of ligand-receptor (LR) interactions associated with schizophrenia (SCZ)

To add clinical relevance to this integrated DLPFC dataset, we next sought to identify interacting cell types and spatially map ligand receptor (LR) interactions associated with neuropsychiatric disorders. We focused on genetic risk for schizophrenia (SCZ) because receptors occur more frequently in SCZ risk genes than would be expected of a brain-expressed gene list of this size (p<0.0001, Fig S42A). First, we identified interacting cell types using cell-cell communication (CCC) analysis (18), which uses a data-driven approach to predict crosstalk between sender and receiver cells based on known LR interactions (Fig. 5A, Table S3). In parallel, using the OpenTargets and Omnipath databases (32, 33), we identified 834 LR pairs (Table S4) associated with genetic risk for SCZ. Of these, 90 interactions overlapped with those identified in CCC analysis where at least one of the interactors was associated with SCZ risk (Table S5).

We chose to prioritize 18 inter- and intra-cellular interactions where both counterparts showed disease association, including 9 interactions involving the protein tyrosine kinase, FYN (Fig. 5A, Fig S42). A consensus LR pair was identified between these complementary data-driven and clinical risk-driven approaches: the membrane-bound ligand ephrin A5 (*EFNA5*) and its receptor ephrin type-A receptor 5 (*EPHA5*). As part of this signaling cascade, we also evaluated the intracellular interaction between *EFNA5* and *FYN*, which was one of the 18 SCZ-associated LR pairs (Fig. 5A).

The Ephrin/Eph signaling system is critical for neuronal wiring during brain development and neural plasticity and synaptic homeostasis in adulthood (34). To better understand the role of this SCZ-associated signaling pathway in the adult DLPFC, we next characterized the cell types mediating this interaction using our snRNA-seq data. We identified enrichment

of *EFNA5*, *EPHA5*, and *FYN* in excitatory neuron populations (Fig. 5C), which are the dominant sender and receiver cells for this LR interaction (Fig. 5B). Particularly, Excit_L5/6 neurons most specifically expressed the ligand *EFNA5* (*Specificity Measure*, *SPM* = 0.6847) and its intracellular partner *FYN* (*SPM* = 0.5345), whereas Excit_L6 neurons most specifically expressed the receptor *EPHA5* (*SPM* = 0.6508, Fig S42B). Furthermore, Excit_L5 and L6 neurons showed the highest co-expression of *FYN* and *EFNA5*, co-expressing in 87.25% of this population (Fig. 5D).

Since EFNA5-EPHA5 is a contact-dependent interaction (34), we used the Visium data to spatially map sites of likely *EFNA5* and *EPHA5* crosstalk. Across data-driven SpDs, the highest proportion of spots co-expressing EFNA5 and EPHA5 localized to Sp₀D₇~L6 (median (interquartile range) = 0.0196 (0.0137), p=4.0e-09, Fig. 5E–F) compared to other Sp_9Ds ($Sp_9D_1 = 0$ (0), $Sp_9D_2 = 0$ (0), $Sp_9D_3 = 0.0052$ (0.0078), $Sp_9D_4 = 0.014$ (0.0142), $Sp_9D_5 = 0.0091$ (0.0102), $Sp_9D_6 = 0$ (0.0016), $Sp_9D_8 = 0.0077$ (0.0123), Sp_9D_8 = 0.0016 (0.0104)). Consistent with snRNA-seq specificity analysis (Fig S42B–C), spots co-expressing EFNA5 and EPHA5 showed a higher predicted proportion of Excit_L5/6 neurons and Excit L6 neurons compared to spots lacking co-expression (Fig. 5H, Fig S43). Although Excit L5 is the dominant *cell2location* predicted cell type among all spots (Fig S41, Fig S43B), it is even higher among these co-expression spots (median predicted proportion = 0.211) than among other spots (Fig S43A-B). Spatial network analyses further supported that co-localization of EFNA5 and EPHA5 occurs frequently in spots containing Excit_L6 neurons - with strongest co-localization relationships between Excit_L6/Excit_L5 neurons, Excit L6/Excit L4 neurons and Excit L6/oligodendrocytes (Fig. 5I, Fig S42D). Spatial mapping of EFNA5 and FYN interactions also showed significant co-expression of these genes in Sp₉D₇~L6 (p=0.0046, Fig S42F–G) with frequent co-localization between Excit_L5/L6 and Excit_4 neurons (Fig S42H-I). In summary, we demonstrate the utility of this integrated single cell and spatial transcriptomic data for identifying and mapping disease-associated interactions in spatially localized cell types across the human DLPFC.

Spatial registration of cell populations across neuropsychiatric disorders

To leverage the large amount of snRNA-seq data collected across the PsychENCODE consortium (PEC) (35, 36), we spatially registered eight DLPFC snRNA-seq datasets generated in the context of several brain disorders (Autism Spectrum Disorder [ASD], SCZ, bipolar disorder, and Williams Syndrome) to both the histological layers and unsupervised SpDs annotated in Visium data (Fig. 6A, Fig S44). Across the consortium, in neurotypical controls, we found that excitatory neuron subtypes with a laminar annotation spatially register to the relevant histological layers and converge on the same unsupervised SpDs. As expected, most inhibitory populations registered to multiple histological layers and unsupervised SpDs, with the exception of Pvalb and VLMC subtypes, which mapped to Sp₉D₈~L4 and Sp₉D₁~L1, respectively. Finally, glial populations also showed expected spatial registration with astrocytes strongly mapping to L1-associated SpDs, oligodendrocytes and OPCs strongly mapping to the WM, and endothelial, pericyte (PC), and smooth muscle cells (SMC) mapping to the newly characterized vascular domain Sp₉D₁.

Next, we spatially registered snRNA-seq data from 30,147 nuclei derived from the human prefrontal cortex (Brodmann areas unspecified) from control donors in a study of ASD (Fig. 6B) (20). As expected, glial populations and laminar excitatory cell types mapped to the relevant SpDs (L2/3 neurons mapped to Sp₉D₃~L2 and Sp₉D₅~L3). We were also able to provide laminar assignments to some cell populations, such as mapping NRGN neuronal subtypes to Sp₉D₅~L3 and Sp₉D₈~L4. For each cell type, we next used clinical gene set enrichment analyses to assess the SpD enrichment of cell type-specific DEGs between individuals with ASD compared to neurotypical controls (Fig. 6C). Across many cell types, we observed multiple Sp₀Ds enriched for ASD DEGs. Sp₀D₃~L2 showed significant enrichment of genes differentially expressed in L2/3 nuclei between individuals with ASD and neurotypical controls (p=2.60e-11), highlighting that these L2/3 DEGs are core Sp₀D₃~L2 marker genes. We also identified spatial enrichments for DEGs expressed in inhibitory neuron and neurogranin populations, including Sp₉D₄~L5 for SV2C inhibitory neurons and Sp₉D₇~L6 for Neu_NRGN_I neurons. Finally, to demonstrate how this largescale dataset can be used to provide spatial information about genes associated with neuropsychiatric disease, we performed gene set enrichment analysis of bulk RNA-seq DEGs identified in a companion PEC study of PTSD and MDD (PEC study 6) (23). For both DLPFC and ventral medial prefrontal cortex (mPFC), we demonstrated that vasculature domain Sp₉D₁~L1 and L1-associated domain Sp₉D₂~L1 are enriched in DEGs associated with both PTSD and MDD. This is consistent with previous studies implicating neuroimmune signaling in PTSD (37), and current PEC single cell analyses that implicate glial and vascular- cells in both MDD and PTSD (PEC study 6) (23). Together, these spatial registration and clinical gene set enrichment analyses add anatomical context to cell type identities and provide valuable biological insights into molecular changes associated with brain disorders, including ASD, MDD, and PTSD.

Discussion

Here we generated a large-scale, transcriptome-wide, data-driven molecular map across the anterior-posterior axis of adult human DLPFC from ten neurotypical control donors. This highly integrated single cell and spatial gene expression reference dataset enabled identification of unsupervised SpDs, which were characterized in terms of both cellular composition and domain-enriched genes, at different resolutions across the DLPFC. We provide a landmark molecular neuroanatomical atlas that complements our understanding of classic cortical cytoarchitecture through identification and characterization of discrete molecularly-defined layers and sublayers. In particular, we annotated a vasculature-rich meninges layer and several molecularly distinct subdomains in histological L1, 4, 5 and 6. An advantage of Visium over snRNA-seq approaches is the ability to capture transcripts in the cell cytoplasm and neuropil, which we speculate may influence identification of higher resolution spatial domains, particularly for demarcating laminar transitions and at the gray/white matter junction (38–40).

In this study, we provide a roadmap for the implementation and biological validation of unsupervised spatial clustering approaches in human brain tissue. Whereas manual annotation of spatial domains is feasible for a limited number of samples in brain regions where neuroanatomical boundaries are well-characterized (12, 22), the application of

data-driven clustering methods is critical for future studies that aim to analyze spatial gene expression changes across diagnostic cohorts groups to identify changes in spatially-resolved cell types. Furthermore, unsupervised approaches will be essential for spatial profiling in brain regions that lack clear molecular or histological boundaries since they allow for identification of unknown or unexpected SpDs as well as SpDs that may be technically difficult to manually annotate (for example the meninges). Although here we evaluated several of the spatial clustering algorithms, including *SpaGCN* and *BayesSpace* (13, 15), there are more tools coming online for both spatially variable gene detection, including *nnSVG* (41), and also spatial domain identification, including *GraphST* and *PRECAST* (16, 17). Large-scale, integrated datasets, including the present study, continue to offer developers of computational tools opportunities to develop methods scalable to atlas-level data, while also extracting meaningful biological information.

Visium offers transcriptome-wide information at spatial resolution; however, a limitation of the platform is that spots often contain multiple cells and cell types. We believe that this might be overcome using spot-level deconvolution tools, as discussed here. We rigorously benchmarked the utility of these tools against Visium-SPG data in the DLPFC, where we manually assigned spots to the histological layers while also immunolabeling 4 broad cell populations in the same tissue sections. From these data we predicted the proportion of cell types in each spot, allowing us to achieve cellular resolution for our spatio-molecular map. We note that although spatial registration correlation approaches can be informative for broadly mapping cell types to SpDs, spot deconvolution approaches provide higher resolution mapping across the cortical layers. This is especially evident for glial cell populations, which show the strongest spatial registration correlations to L1 and WM, but are indeed localized to all other SpDs when performing spot deconvolution. This is also true for endothelial cells, which are strongly associated with vasculature-enriched Sp₉D₁~L1, but are similarly distributed across the cortex when utilizing spot deconvolution approaches. As spot deconvolution tools continue to improve with the emergence of more large-scale single cell and spatial transcriptomic datasets, we expect that algorithms will be able to map finer resolution and rarer cell types.

While other imaging-based, spatially-resolved transcriptomics platforms, such as Xenium and MERFISH (42, 43), directly measure transcripts in individual cells, only a limited number of genes can currently be probed. In contrast, the discovery-based approach afforded by Visium, as well as its scalability to a large number of samples, allowed for robust identification of previously unrecognized spatial marker genes across many donors. These Visium-identified genes can be followed up at single cell resolution in smaller cohorts using probe-based approaches, such as MERFISH, which was recently applied to the middle and superior temporal gyri in the human brain (44). As spatial transcriptomics technologies continue to evolve, it will be important to functionally validate these spatial domains and consider the biological significance of these molecularly-defined compartments. Indeed, previous spatial transcriptomic studies in the mouse brain identified unique astrocyte layers that diverge from classic excitatory neuron layers (45), suggesting that the cortex likely has a more complex laminar architecture than previously appreciated. Future studies should evaluate the conservation of laminar spatial domains between rodent and human brain and explore their functional and clinical relevance.

In terms of clinical relevance, alterations in neural activity patterns within the DLPFC are noted in several neurodevelopmental and neuropsychiatric disorders (46-49), and it is hypothesized that changes in molecular signaling cascades may contribute to these alterations in activity. To gain insight into molecular dysfunction in DLPFC in the context of disease, we used our integrated molecular atlas to spatially map cell type-specific LR interactions that are associated with genetic risk for SCZ. For example, we highlight the interaction between EFNA5 and EPHA5 in excitatory L5/6 and L6 neuron subtypes in deep cortical layers, which is consistent with results from the most recently released SCZ genome-wide association study (GWAS) that identified enrichment of SCZ risk genes in glutamatergic neurons (50). Not only is EFNA5 the locus of a GWAS-identified common SCZ risk variant, but it is also differentially expressed between individuals with SCZ and neurotypical controls in specific cell types (21). Spatially mapping disease-relevant LR pairs, which are often highly specific and druggable targets, can provide valuable insights into pathophysiology and can help prioritize spatially restricted targets for therapeutic development. In combination with our interactive web resources, this highly integrated single cell and spatial transcriptomic data from neurotypical control DLPFC can be used to accelerate research across a variety of brain disorders by allowing researchers to search for relevant genes of interest, spatially register clinical gene sets, and explore disease-associated cell types for complementary assays, such as in vitro disease models.

Finally, spatial registration of eight DLPFC snRNA-seq datasets collected across the PsychENCODE consortium in the context of different neuropsychiatric disorders (35) revealed a convergence of excitatory, inhibitory, and non-neuronal cell types in relevant spatial domains. We observed increased confidence of inhibitory neuron mapping in our current expanded study compared to (12), likely due to the larger donor/sample number and data-driven clustering approach, which allowed for identification of finer resolution SpDs. Furthermore using ASD as an example (20), we demonstrated the utility of our data-driven molecular atlas for localizing cell-type specific DEGs to specific SpDs. For example, ASD DEGs in VIP inhibitory neurons were enriched in L3, L5, and L6-associated spatial domains, while those in SV2C inhibitory neurons were enriched only in the L5-associated domain. Together, this analysis provides anatomical context for cell type-specific gene expression changes and molecular mechanisms associated with neurodevelopmental disorders and psychiatric illness.

In summary, we provide a large-scale, highly integrated single cell and spatial transcriptomics resource for understanding the molecular neuroanatomy of the human DLPFC. We share web-based tools for the scientific community to interact with these datasets for further interrogation of molecular pathways associated with brain disorders.

Materials and methods summary

Detailed materials and methods can be found in the supplementary materials. Three tissue blocks from the anterior, middle, and posterior positions along the rostral-caudal axis of human DLPFC were microdissected from the post-mortem brains of 10 donors for a total of 30 tissue blocks. The 10x Genomics Visium Spatial Gene Expression protocol was performed on all blocks (n = 30). The 10x Genomics Visium SPG and 3' Single

Cell Gene Expression protocols were performed on a subset of blocks (n = 4 and 19, respectively). Visium and Visium-SPG data was processed with 10x Genomics SpaceRanger (51). Dimensionality reduction was performed by using scater (52), and batch correction was performed with Harmony (53). Unsupervised clustering was performed on the Visium data with BayesSpace (13) defining data-driven SpDs. An optimal number of SpDs was determined with fasthplus (54). DEGs between SpDs were found with ANOVA, enrichment, and pairwise models with tools from spatialLIBD (55), powered by limma (56). snRNAseq data were processed with 10x Genomics CellRanger (57). Reduced dimensions were calculated with GLM-PCA (58), and batch correction was performed with *Harmony* (53). Graph-based and hierarchical clustering described in (59) identified cell type populations, which were annotated by using established marker genes (59, 60). Spatial registration was applied to snRNA-seq and SpDs with tools from spatialLIBD (55). Images from Visium and Visium-SPG were processed with *VistoSeg* (61). Nuclei were segmented with *Cellpose* (62). A decision-tree cell type classifier was built with scikit-learn (63) on the basis of a training set classified by an expert. Cell segmentation and classification from Visium-SPG was used to benchmark spot deconvolution algorithms Tangram, Cell2Location, and SPOTlight (14, 30, 31). Spot deconvolution on the 30 Visium samples was performed with both Tangram and Cell2Location. CCC was performed with LIANA (18). The SCZ risk gene list was obtained from *OpenTargets* (32). Spatial registration was performed on the uniformly processed PsychENCODE snRNA-seq datasets (35). Enrichment statistics were calculated with spatialLIBD for the DEGs in an ASD snRNA-seq dataset (20) and bulk RNAseq PTSD and MDD dataset (23).

Supplementary Material

Refer to Web version on PubMed Central for supplementary material.

Acknowledgements

Data were generated as part of the PsychENCODE Consortium, supported by: U01DA048279, U01MH103339, U01MH103340, U01MH103346, U01MH103365, U01MH103392, U01MH116438, U01MH116441, U01MH116442, U01MH116488, U01MH116489, U01MH116492, U01MH122590, U01MH122591, U01MH122592, U01MH122849, U01MH122678, U01MH122681, U01MH116487, U01MH122509,R01MH094714, R01MH105472, R01MH105898, R01MH109677, R01MH109715, R01MH110905, R01MH110920, R01MH110921, R01MH110926, R01MH110927, R01MH110928, R01MH111721, R01MH117291, R01MH117292, R01MH117293, R21MH102791, R21MH103877, R21MH105853, R21MH105881, R21MH109956, R56MH114899, R56MH114901, R56MH114911, R01MH125516, R01MH126459, R01MH129301, R01MH126393, R01MH121521, R01MH116529, R01MH129817, R01MH117406, and P50MH106934 awarded to: Alexej Abyzov, Nadav Ahituv, Schahram Akbarian, Kristen Brennand, Andrew Chess, Gregory Cooper, Gregory Crawford, Stella Dracheva, Peggy Farnham, Michael Gandal, Mark Gerstein, Daniel Geschwind, Fernando Goes, Joachim F. Hallmayer, Vahram Haroutunian, Thomas M. Hyde, Andrew Jaffe, Peng Jin, Manolis Kellis, Joel Kleinman, James A. Knowles, Arnold Kriegstein, Chunyu Liu, Christopher E. Mason, Keri Martinowich, Eran Mukamel, Richard Myers, Charles Nemeroff, Mette Peters, Dalila Pinto, Katherine Pollard, Kerry Ressler, Panos Roussos, Stephan Sanders, Nenad Sestan, Pamela Sklar, Michael P. Snyder, Matthew State, Jason Stein, Patrick Sullivan, Alexander E. Urban, Flora Vaccarino, Stephen Warren, Daniel Weinberger, Sherman Weissman, Zhiping Weng, Kevin White, A. Jeremy Willsey, Hyejung Won, and Peter Zandi. Additional data were provided to the PsychENCODE Consortium, supported by 2015 and 2018 NARSAD Young Investigator grants from Brain & Behavior Research Foundation awarded to: Nikolaos Daskalakis.

We would like to thank Linda Orzolek and the Johns Hopkins Single Cell and Transcriptomics Core facility for executing all sequencing. We also acknowledge Elizabeth Engle and the Johns Hopkins Tumor Microenvironment Lab core facility for assistance with the Vectra Polaris slide scanner. We would like to thank the Joint High Performance Computing Exchange (JHPCE) for providing computing resources for these analyses. While an Investigator at LIBD, Andrew E. Jaffe helped secure funding for this work, and contributed to the conceptualization

of the spatial registration framework. We thank William S. Ulrich for his assistance in deploying the Samui interactive websites. We thank R. A. Miller for assistance with uploading data to the NIMH Data Archive. We thank Daniel R. Weinberger and members of the PsychENCODE consortium for feedback on the manuscript. We thank Amy Deep-Soboslay and her diagnostic team for curation of brain samples. We thank the neuropathology team, especially James Tooke, for assistance with tissue dissection. We thank the physicians and staff at the brain donation sites and the generosity of donor families for supporting our research efforts. Finally, we thank the families of Connie and Stephen Lieber and Milton and Tamar Maltz for their generous support of this work. Schematic illustrations were generated using Biorender. The opinions expressed in this article are the author's own and do not reflect the view of the National Institute on Aging, the National Institutes of Health, the Department of Health and Human Services, or the United States government. A preprint of this work is available at https://doi.org/10.1101/2023.02.15.528722. For the purpose of open access, the authors have applied a CC BY public copyright license to all Author Accepted Manuscripts arising from this submission.

Funding

Funding for these studies was provided by the Lieber Institute for Brain Development (LIBD) and National Institute of Mental Health (NIH) grants U01MH122849 and R01MH123183. AB was supported by NIH National Institute of General Medical Sciences (NIGMS) grant R35GM139580. MGP and MR were funded by Aligning Science Across Parkinson's [Grant numbers: ASAP-000478 and ASAP-000509] through the Michael J. Fox Foundation for Parkinson's Research (MJFF).

Data and materials availability

The source data described in this manuscript are available from the National Institute of Mental Health (NIMH) Data Archive (https://nda.nih.gov/) (64). The source data are also publicly available from the Globus endpoint 'jhpce#spatialDLPFC' and 'jhpce#DLPFC_snRNAseq' that are also listed at http://research.libd.org/globus. The raw data provided through Globus include all the FASTQ files and raw image files. Processed data are publicly available from the Bioconductor package *spatialLIBD* version 1.11.7 (55) through the fetch_data() function. All source code is publicly available through GitHub and permanently archived through Zenodo at https://github.com/LieberInstitute/spatialDLPFC (65) and https://github.com/LieberInstitute/DLPFC_snRNAseq (66).

We provide several interactive web applications to explore this highly integrated DLPFC dataset as listed at http://research.libd.org/spatialDLPFC/#interactive-websites. First, we developed two *spatialLIBD* apps (55) that allow users to analyze Visium gene expression, cell segmentation, spot deconvolution, and clinical gene set enrichment results at k=9 and k=16. We added a third *spatialLIBD* app for the position differential expression results and a fourth for the Visium-SPG data. Second, both our Visium H&E (n=30 tissue sections) and Visium-SPG (n=4 tissue sections) data is available through a newly created performant web-based interactive visualization tool, called *Samui Browser* (67), which allows rapid loading, visualization, and custom annotation of high resolution images and corresponding gene expression. Finally, snRNA-seq data at both fine and laminar resolution as well as pseudo-bulked Visium data (k=9, 16, and 28) are available through *iSEE* apps (68) that allows users to visualize expression levels for genes of interest through violin and heatmap plots.

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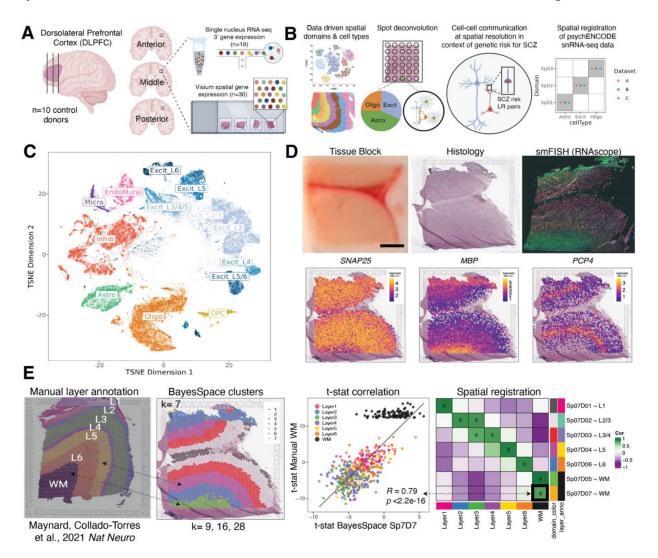


Figure 1. Study design to generate paired single nucleus RNA-sequencing (snRNA-seq) and spatially-resolved transcriptomic data across DLPFC.

(A) DLPFC tissue blocks were dissected across the rostral-caudal axis from 10 adult neurotypical control postmortem human brains, including anterior (Ant), middle (Mid), and posterior (Post) positions (n=3 blocks per donor, n=30 blocks total). The same tissue blocks were used for snRNA-seq (10x Genomics 3' gene expression assay, n=1-2 blocks per donor, n=19 samples) and spatial transcriptomics (10x Genomics Visium spatial gene expression assay, n=3 blocks per donor, n=30 samples). (B) Paired snRNA-seq and Visium data were used to identify data-driven spatial domains (SpDs) and cell types, perform spot deconvolution, conduct cell-cell communication analyses, and spatially register companion PsychENCODE snRNA-seq DLPFC data. (C) t-distributed stochastic neighbor embedding (t-SNE) summarizing layer resolution cell types identified by snRNA-seq. (D) Tissue block orientation and morphology was confirmed by hematoxylin and eosin (H&E) staining and single molecule fluorescent in situ hybridization (smFISH) with RNAscope (SLC17A7 marking excitatory neurons in pink, MBP marking white matter (WM) in green, RELN marking layer (L)1 in yellow, and NR4A2 marking L6 in orange). Scale bar is 2mm.

Spotplots depicting log transformed normalized expression (logcounts) of *SNAP25*, *MBP*, and *PCP4* in the Visium data confirm the presence of gray matter, WM, and cortical layers, respectively (see also Fig S2–Fig S4). (E) Schematic of unsupervised SpD identification and registration using *BayesSpace* SpDs at *k*=7. Enrichment *t*-statistics computed on *BayesSpace* SpDs were correlated with manual histological layer annotations from (12) to map SpDs to known histological layers. The heatmap of correlation values summarizes the relationship between *BayesSpace* SpDs and classic histological layers. Higher confidence annotations (cor > 0.25, merge ratio = 0.1, see Methods: Spatial registration of *Spatial Domains*) are marked with an "X".

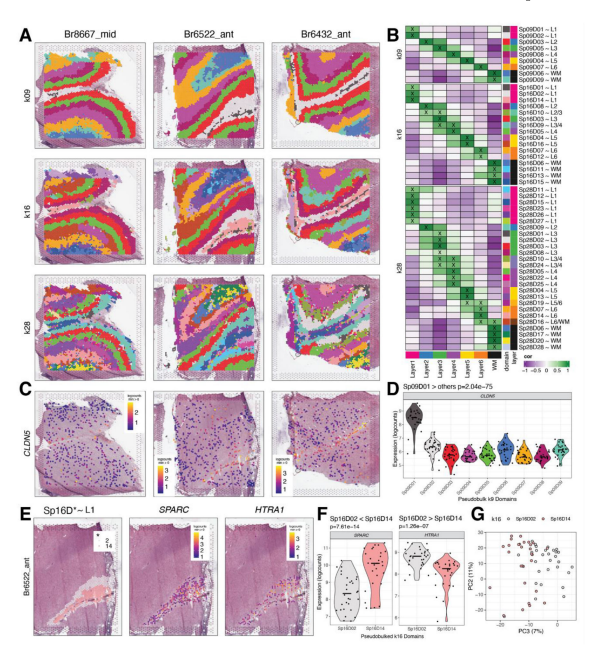


Figure 2. Unsupervised clustering at different resolutions identifying spatial domains (SpDs) and defining molecular anatomy of DLPFC.

(A) BayesSpace clustering at k=9, 16, and 28 (broad, fine, and super-fine resolution, respectively, which we refer to as Sp_kD_d for domain d from SpDs at k resolution) for three representative DLPFC tissue sections (Br8667_mid, Br6522_ant, Br6432_ant). (B) Heatmap of spatial registration with manually annotated histological layers from (12). BayesSpace identifies laminar SpDs at increasing k with the majority of Sp_kDs correlating with one or more histological layer(s). SpDs were assigned layer annotations following spatial registration to histological layers. Annotations with high confidence (cor > 0.25, merge ratio = 0.1, see Methods: Spatial registration of Spatial Domains) are marked with an "X", and this histological layer association is denoted for a given Sp_kD_d by adding

"~L," where L is the most strongly correlated histological layer (or WM). See also Fig S11–Fig S18. (C) Spotplots depicting expression of CLDN5 in vasculature domain 1 at k=9 resolution (Sp₉D₁). (D) Boxplot confirming enrichment of CLDN5 in Sp₉D₁ compared to other Sp₉Ds across 30 tissue sections. (E) Spotplots of representative section Br6522_ant showing identification of molecularly-defined sublayers for histological L1 at k=16 (Sp₁₆D₂ and Sp₁₆D₁₄) and enrichment of HTRA1 and SPARC, respectively. (F) Boxplots quantifying enrichment of SPARC and HTRA1 in Sp₁₆D₁₄ and Sp₁₆D₂, respectively, across 30 tissue sections. (G) PCA plot showing separation of Sp₁₆D₂ and Sp₁₆D₁₄ supporting identification of molecularly distinct SpDs.

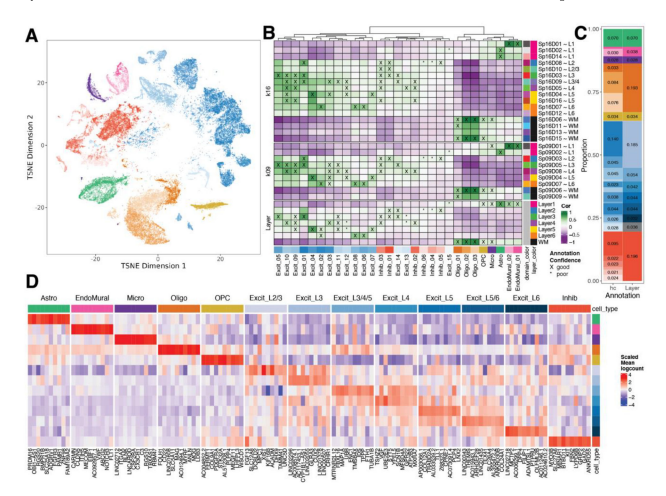


Figure 3. Spatial registration of fine resolution snRNA-seq clusters defining laminar cell types. (**A**) t-distributed stochastic neighbor embedding (t-SNE) plot of 56,447 nuclei across 29 cell type-annotated fine resolution hierarchical clusters (hc; related to Fig S25A). (**B**) Spatial registration heatmap showing correlation between snRNA-seq hierarchical clusters (hc) and manually annotated histological layers from (12) as well as unsupervised *BayesSpace* clusters at k=9 and 16 (Sp₉Ds and Sp₁₆Ds). Hierarchical clusters for excitatory neurons (Excit) were assigned layer-level annotations following spatial registration to histological layers (cor > 0.25, merge ratio = 0.25, see Methods: snRNA-seq spatial registration). For Sp₉Ds and Sp₁₆Ds, annotations with good confidence (cor > 0.25, merge ratio = 0.1) are marked with "X" and poor confidence are marked with "*". (**C**) Summary barplot of cell type composition for hc and layer level resolutions (related to Fig S25B & Fig S26) (**D**) Heatmap of the scaled mean pseudo-bulked logcounts for the top 10 marker genes identified for each cell type at layer-level resolution.

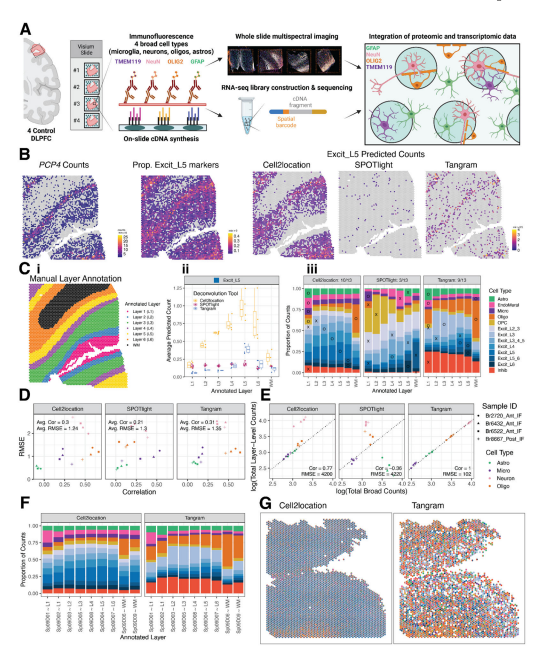


Figure 4. Integration of snRNA-seq and Visium data to benchmark spot deconvolution algorithms and define cellular composition across spatial domains.

(A) Schematic of the Visium-SPG protocol. (B) For Br6522_Ant_IF, counts for L5 marker gene *PCP4* are compared to the proportion of Excit_L5 marker genes with nonzero expression as well as the counts of Excit_L5 cells as predicted by the 3 evaluated deconvolution algorithms. (C) Example of manually annotated layer assignments for Br6522_Ant_IF (i), which are used to benchmark predicted cell type composition across layers. Using Excit_L5 as an example, predicted Excit_L5 counts for each method are averaged across all spots within each annotated layer for each tissue section (ii). These data are summarized across layers and tissue sections for the 13 cell types using a bar plot (iii). An "X" or "O" is placed on the layer with maximal proportion; an "O" is placed for

a "correct" match for the given cell type, and an "X" is placed otherwise. For example, Tangram correctly predicts the maximal proportion of Excit_L5 cells in L5 annotated spots, leading to the placement of an "O" for Excit_L5. The "O"s are tallied for each method to generate a summary score in each facet's title (for example, 9 of 13 cell types were correctly predicted to the expected layer using *Tangram*). (D) Predicted counts for a given method, section, and layer-level cell type are collapsed and compared against the corresponding CART predictions by computing the Pearson correlation and RMSE, forming a single point in the scatterplot (Supplemental Methods: Evaluating performance of spot-deconvolution methods). Each of these values is then averaged to generate a single correlation and RMSE value for each method, indicated in the top left inside each plot facet. (E) Section-wide counts for each cell type are compared between broad and layer-level resolutions, collapsed onto the cell-type resolution used by the CART, where values theoretically should precisely match. (F) The predicted proportion of cells in each Sp₉D, deconvoluted by Cell2location and Tangram, are averaged across all Visium samples (n=30). (G) Cell composition of each Visium spot for Br8667_mid, deconvoluted by Cell2location and Tangram, revealing differences in cell composition prediction.

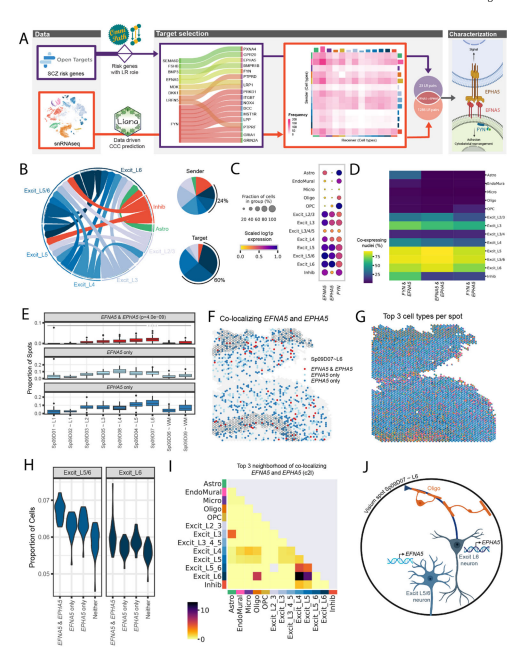


Figure 5. Schizophrenia (SCZ)-associated ligand-receptor (LR) interactions identified by integrative analysis of snRNA-seq and Visium data.

(A) The LR interaction between membrane-bound ligand ephrin A5 (*EFNA5*) and ephrin type-A receptor 5 (*EPHA5*) is a consensus target identified in both data-driven (Table S3) and clinical risk-driven LR (Table S4) analyses. Notably, this interaction also requires an intracellular interaction between *EFNA5* and protein tyrosine kinase (*FYN*), which was also identified among clinical risk targets. (B) Cell-cell communication analysis predicts the sender/receiver cross-talk pattern of *EFNA5-EPHA5* between layer-level cell types visualized in a circular plot. Excit_L5/6 and Excit_L6 neurons account for 24% of the cross-talk as senders and 60% as targets compared to other cell types shown in the pie charts. (C-D) Downstream analysis of snRNA-seq data characterizes *FYN-EFNA5-EPHA5*

signaling pathway, showing these genes are highly enriched (C) and co-expressed (D) in excitatory neuron populations. (E) Across all 30 tissue sections, EFNA5 and EPHA5 are co-expressed in a statistically higher proportion of spots in Sp₀D₇ (median (interquartile range) = 0.0196 (0.0137), p = 4.0e-09) compared to other Sp₉Ds (Sp₉D₁ = 0 (0), Sp₉D₂ $= 0 (0), Sp_9D_3 = 0.0052 (0.0078), Sp_9D_4 = 0.0140 (0.0142), Sp_9D_5 = 0.0091 (0.0102),$ $Sp_9D_6 = 0 (0.0016), Sp_9D_8 = 0.0077 (0.0123), Sp_9D_8 = 0.0016 (0.0104)).$ (**F**) Spotplot of EFNA5 and EPHA5 co-expression in Br8667_mid. (G) Spotplot with spot-level pie charts for Br8667_mid showing the top 3 dominant cell types in each Visium spot predicted by Cell2location (c2l). (H) Visium spots co-expressing EFNA5 and EPHA5 have higher proportions of predicted Excit L5/6 neurons (p=1.8e-12) and Excit L6 (p=3.9e-4) compared to non-coexpressing spots, consistent with snRNA-seq specificity analyses (Fig S42). Few other cell types show this relationship (Fig S43). Complementary analyses of EFNA5 and FYN co-expression are shown in Fig S42. (I) Spatial network analysis of all 30 tissue sections, using top 3 dominant c21 cell types in each spot (exemplified in G with Br8667_mid), confirms EFNA5 and EPHA5 co-expression occurs frequently in spots containing Excit_L6 neurons. Complementary analyses using top 6 dominant c2l cell types as well as Tangram predictions are reported in Fig S42. (J) Schematic of a Visium spot depicting EFNA5-EPHA5 interactions between Excit_L5/6 neurons and Excit_L6. The high colocalization score in the spatial network analysis in (I) suggests oligodendrocytes also likely co-exist with Excit L6 neurons.

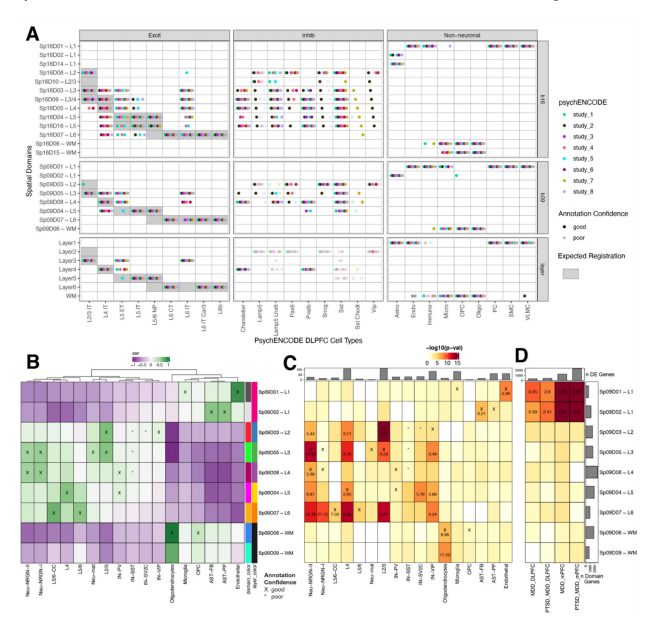


Figure 6. Spatial enrichment of cell types and genes associated with neurodevelopmental and neuropsychiatric disorders.

(A) Dot plot summarizing spatial registration results for eight PsychENCODE (PEC) snRNA-seq datasets from human DLPFC. snRNA-seq data was uniformly processed through the same pipeline and annotated with common nomenclature based on work from Allen Brain Institute (35, 69). Registration was performed for control donors only (see Fig S44 for full dataset) across manually annotated histological layers from (12) as well as unsupervised *BayesSpace* clusters at *k*=9 and *k*=16 (Sp₉Ds and Sp₁₆Ds, respectively). Each dot shows the histological layer(s) or SpD(s) where a dataset's cell type was annotated during spatial registration. Solid dots show good confidence in the spatial annotation, translucent dots show poor confidence in the annotation. IT, intratelencephalon-projecting; ET, extratelencephalon-projecting; CT, corticothalamic-projecting; NP, near-projecting; VLMC, vascular lepotomeningeal cell; OPC, oligodendrocyte precursor cell; PC, pericyte;

SMC, smooth muscle cell. (**B**) Spatial registration of cell type populations from control samples from (20) against unsupervised BayesSpace clusters at k=9 (Sp₉Ds). Higher confidence annotations (cor > 0.25, merge ratio = 0.1, Supplemental Methods: Spatial registration of PsychENCODE and other external snRNA-seq datasets) are marked with an "X". (**C**) Enrichment analysis using Fisher's exact test for Sp₉D- enriched statistics versus differentially expressed genes (DEGs, FDR < 0.05) in Autism spectrum disorder (ASD) for each cell type population. The values are the odds ratios (ORs) for the enrichment in significant (FDR < 0.001) blocks of the heatmap, and the color scale indicates $-\log 10(p$ -value) for the enrichment test. The top bar plot shows the number of DEGs for each cell type. (**D**) Enrichment analysis using Fisher's exact test for Sp₉D- enriched statistics versus differentially expressed genes (DEGs, FDR < 0.05) in Post Traumatic Stress Disorder (PTSD) and/or Major Depressive Disorder (MDD) in bulk RNA-seq of DLPFC and medial prefrontal cortex (mPFC) (23). Top bar plot shows the number of DEGs for each DE test. Left bar plot shows the number of significantly enriched genes for each Sp₉D in both enrichment analyses.