Cross-expression analysis reveals patterns of coordinated gene expression in spatial transcriptomics

Abstract

 Spatial transcriptomics promises to transform our understanding of tissue biology by molecularly profiling individual cells *in situ*. A fundamental question they allow us to ask is how nearby cells orchestrate their gene expression. To investigate this, we introduce cross-expression, a novel framework for discovering gene pairs that coordinate their expression across neighboring cells. Just as co-expression quantifies synchronized gene expression within the same cells, cross- expression measures coordinated gene expression between spatially adjacent cells, allowing us to understand tissue gene expression programs with single cell resolution. Using this framework, we recover ligand-receptor partners and discover gene combinations marking anatomical regions. More generally, we create cross-expression networks to find gene modules with orchestrated expression patterns. Finally, we provide an efficient R package to facilitate cross-expression analysis, quantify effect sizes, and generate novel visualizations to better understand spatial gene expression programs.

 Spatial transcriptomics records cells' gene expression alongside their physical locations, enabling 27 us to understand how they influence one another within the broader tissue context¹. Focusing on select genes, imaging-based platforms profile expression at the single cell level, giving a high-29 resolution snapshot of spatial gene expression^{2–7}. They have facilitated numerous studies on 30 defining local spatial patterns^{8–11}, finding gene covariation in spatial niches^{12–16}, elucidating cell-31 cell interactions using ligand-receptor expression^{17–27}, and determining spatial cell type 32 heterogeneity and tissue structure^{10,28–30}. These efforts have resulted in a greater understanding 33 of tissue biology, culminating in the generation of reference atlases $31-34$.

 Imaging-based platforms can now profile up to a few thousand genes in millions of cells^{2–} $\frac{7}{1}$, generating large amounts of data ripe for biological discovery. Historically, large-scale global gene networks have been instrumental in uncovering fundamental biological processes by 37 leveraging the power of high-throughput data to compute gene-gene interactions^{35,36}. A promising 38 approach to exploiting large-scale single-cell RNA-seq data is gene co-expression analysis $37-39$. which investigates how genes covary within cells and therefore discovers modules of functionally 40 related genes. Extending this concept to spatial transcriptomics, a recent study¹² characterized gene covariation within well-defined spatial niches, finding continuous gradients during spinal cord 42 development and localizing cortical somatostatin-positive interneuron subtypes. Another study⁴⁰ used co-expression to create hierarchical tissue structures, revealing the multi-scale organization of the hippocampus. While these studies fruitfully apply gene co-expression within cells to characterize tissue structure at multiple spatial scales, the co-expression framework is silent on patterns between cells as, for example, when a gene expression program in one cell gives rise to a complementary pattern in a neighboring cell.

 Here we introduce cross-expression, a novel conceptual and statistical framework to understand coordinated gene expression as a network *between* neighboring cells. Whereas co- expression captures gene covariation within the same cells, cross-expression measures their coordination between neighboring cells, thereby highlighting how gene expression is orchestrated across the tissue. By developing methods to focus on the conjugate network to cell-cell interaction networks, we are able to investigate novel features that characterize individual genes, cells, and shared patterns across both. For example, we create a cross-expression network, finding that *Gpr20*, a G protein-coupled receptor that appears to line the blood vessels, is a central gene with high node degree and defines visible spatial tracts. Within the same network, we discover an interacting subset of genes enriched in astrocyte-mediated regulation of vascular processes, an essential biological function requiring spatially proximal gene expression. Investigating the relationship between cross-expression and cell type composition, we find that cross-expression

 is frequently driven by compositional differences, where 64% of cross-expressing cell pairs have different cell subtype labels. Using cross-expression to discover anatomical marker combinations, we find gene pairs that cross-express specifically in cells that are located in the thalamus even though the individual genes are highly expressed in other regions. To investigate known gene 64 patterns, we use BARseg^{34,41} to collect mouse whole-brain data with a gene panel containing select ligands and receptors, finding that these genes are highly cross-expressed, thus confirming previous reports that typically target known interaction partners. Collectively, our unbiased framework fully leverages the spatial dimension at the cellular resolution to discover novel genes with coordinated expression, helping us better understand how cells influence one another in tissue.

Results

Cross-expression overview

 Just as co-expression between two genes in single cell data can be conceptualized as the degree to which knowing one gene's expression in a given cell predicts the other gene's expression in the same cell, cross-expression is the degree to which knowing the expression of one gene in a given cell predicts the expression of another gene in a spatially related cell, typically the neighbor. One trivial case where this can occur is when two cells exhibit the same expression pattern; here, the prediction of the neighboring cell effectively captures co-expression and cell type composition. To exclude this, we define cross-expression as the predictions of neighbor expression where co- expression alone provides no performance. Specifically, cross-expression occurs where there is a consistent pattern in which gene A is expressed in a cell without gene B, and gene B is 82 expressed in its neighbor without gene A (Fig. 1a).

 To quantify cross-expression (Fig. 1b), we first consider cell-neighbor pairs where the cells express gene A. We next test if the neighbors express gene B. If many do, given gene B's incidence in the population at large, then these genes are said to cross-express. Additionally, we quantify the effect size by comparing the number of neighbors expressing gene B to the number of cells co-expressing genes A and B (Fig. 1c). Using this procedure on *n* nearest neighbors, we filter for a bullseye-like distribution with low co-expression (center) and high cross-expression (rings). In subsequent text, "gene A" and "gene B" refer to their expression in the central (or reference) and neighboring (or spatially adjacent) cells, respectively, unless indicated otherwise.

91 To explore cross-expression, we use imaging-based spatial transcriptomics²⁻⁷ for several reasons. First, these platforms profile gene expression at the single cell level, allowing us to ask how individual cells influence each other. Second, they share common steps, such as transcript identification and cell segmentation, that allow consistent downstream analysis and interpretation.

95 Third, these platforms have been used to generate large amounts of data^{2–5}, making them suitable for developing and validating the computational framework underlying cross-expression.

 Although we focus on individual cells, groups of cells may form spatial niches and gene expression may be coordinated between niches. To assay cross-expression at this coarser resolution, we average a gene's expression in a cell with its expression in the neighbors (Fig. 1d), thus smoothing it within a spatial niche, with the number of neighbors forming the niche size. Accordingly, cross-expression can be compared across niches by, for example, finding associations between smoothed niche-specific gene expression profiles.

 To enable these analyses, we provide an efficient software package in R that requires the gene expression and cell location matrices as inputs, and outputs a gene-gene p-value matrix that facilitates downstream analyses, such as cross-expression network construction (Fig. 1e). The package also contains functions for computing effect sizes, making bullseye plots, smoothing gene expression, viewing cross-expressing cells *in situ*, and assessing if cross-expression is spatially enriched. Collectively, the cross-expression framework uses spatial information to discover how genes coordinate their expression across neighboring cells, thereby providing a novel analytical framework for deeply exploring spatial transcriptomic data.

 Fig. 1 | Cross-expression analysis. a, Cross-expression is the mutually exclusive expression of 113 genes between neighboring cells. If either cell expresses both genes, the cell pair is not 114 considered to cross-express. **b**, The probability that two genes cross-express is modeled by the considered to cross-express. **b**, The probability that two genes cross-express is modeled by the hypergeometric distribution, where all the cells expressing gene A are sampled and their neighbors expressing gene B are deemed as 'successful trials'. **c**, Cross-expression is compared to co-expression to quantify the effect size, where the number of neighbors with gene B is compared to the number of cells co-expressing genes A and B. 'Sampled cells' (center) are those expressing gene A and neighbors are concentric rings, with the order indicating the *n*-th neighbor. **d**, Averaging gene expression between cells and their neighbors smooths it, extending cross- expression analysis from cell pairs to regions. Number of neighbors is the kernel size. **e**, Software 122 inputs are the gene expression and cell location matrices, and the output is a p-value matrix,
123 vulich enables downstream analyses, such as cross-expression network construction. Created which enables downstream analyses, such as cross-expression network construction. Created with BioRender.com.

Cross-expression recovers ligand-receptor pairs and reveals coordinated gene expression

profiles across the tissue

 To study cross-expression, we used BARseq (barcoded anatomy resolved by sequencing) to collect data from a whole mouse brain. This dataset profiled expression in 1 million cells across 15 sagittal slices, using a gene panel of 104 cortical cell type markers and 25 ligands and receptors, including neuropeptides, their receptors, and monoamine neuromodulatory receptors.

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- 132 Because receptors and their corresponding ligands are often expressed in nearby cells^{17–27}, we

 reasoned that these genes should show cross-expression. As an example, we find that across the cortical somatosensory nose region and visceral areas (Fig. 2a), the neuropeptide somatostatin *Sst* and its cognate receptor *Sstr2* are cross-expressed (Fig. 2b, left, p-values ≤ 0.01 and 0.05, respectively). Indeed, these genes are consistently expressed across neighboring cells (Fig. 2b, right), a pattern that is otherwise difficult to discover without prior knowledge.

 Next, we explore the bullseye plots, which allow us to quantify the effect size by comparing cross-expression with co-expression. For *Sst* and *Sstr2* in the somatosensory nose (2,015 cells) and visceral regions (1,603 cells), we see a bullseye pattern with low co-expression and high cross-expression that decreases for distant neighbors (Fig. 2c). Specifically, for these regions the bullseye score ratio between the first neighbor and the central cell is 1.8 and 1.6, respectively, whereas the ratio between the averaged second-to-tenth neighbor and the central cell is 1.3 and 144 1.2. These findings suggest that for central cells expressing one gene in a pair, a higher proportion of adjacent neighbors, but not the more distant ones, express the other gene within the local spatial niche, underscoring the specificity and resolution with which patterns of coordinated gene expression can be recovered. We next compare the bullseye plots for gene pairs with and without cross-expression (Fig. 2d), finding that the former match the patterns just described. To quantify 149 this, we compare the bullseve scores of the nearest neighbors with those of cells expressing gene A, discovering that this ratio is much greater for genes that cross-express than for those that do not (Fig. 2d, inset, Mann-Whitney U test, p-value ≤ 0.001, median ratios: 1.5 and 0.9, respectively). Notably, this ratio is approximately 1 for genes that do not cross-express, suggesting that here gene B is expressed in neighbors and cells alike. Hence, the bullseye approach intuitively visualizes and quantifies the effect size, making it suitable for downstream analysis, such as comparing cross-expression between different regions.

 We next conducted brain-wide analysis and found that 20% of possible ligand-receptor gene pairs and 4% of non-signaling gene pairs are cross-expressed, thus generating novel candidates that potentially encode functionally relevant interactions. In fact, these patterns are spatially enriched, where most gene pairs cross-express in a few slices and some cross-express in multiple slices (Extended Data Fig. 1a). We now highlight some notable examples of cross-161 expression for both signaling and non-signaling genes. The dopamine receptor D_1 (*Drd1*) and proenkephalin (*Penk*) are strongly cross-expressed (Extended Data Fig. 1b), with discernible 163 spatial enrichment in the striatal regions. *Drd1* is involved in the reward system^{42,43} while *Penk* 164 generates opioids that modulate fear response⁴⁴ and nociception^{45,46}, suggesting that these genes may be involved in avoidance behavior. Indeed, Penk is strongly co-expressed with the dopamine receptor D² (*Drd2*) (Pearson's *R* = 0.72 in scRNA-seq striatal data; *Drd2* is not in our gene panel),

 indicating that the D1 and D2 neurons are spatially intermingled, allowing them to play interrelated 168 roles in motor control⁴⁷. We also find that the somatostatin receptor *Sstr2* cross-expresses with vasoactive intestinal polypeptide receptor 1 (*Vipr1*/*VPAC1*) in the cortex (Extended Data Fig. 1c), suggesting a potential complementary interaction in modulating local neuronal circuits and 171 influencing neuroendocrine signaling pathways^{48,49}. Beyond the signaling genes, we note that the fibril-associated *Col19a1* (collagen type XIX alpha 1 chain), a gene involved in maintaining the 173 extracellular matrix (ECM) integrity^{50,51}, cross-expresses with *C1ql3* (complement C1q like 3) (Extended Data Fig. 1d), whose secretion in the ECM facilitates synapse homeostasis and the formation of cell-cell adhesion complexes 52,53 . Finally, our analysis reveals that *Marcksl1* (myristoylated alanine-rich C-kinase substrate), which is involved in adherens junctions and 177 cytoskeletal processes^{54,55}, cross-expresses with actin beta (Actb) (Extended Data Fig. 1e), 178 hinting at their possible involvement in local tissue architecture⁵⁶. Taken together, the cross- expression analysis not only reveals expected relationships between signaling molecules, but it also discovers genes implicated in the tissue microenvironment. Accordingly, cross-expression is an unbiased framework for finding genes with orchestrated spatial expression profiles, with potential for novel discovery increasing as the gene panel gets larger.

 We have thus far investigated cross-expression between cells and their neighbors. Yet, gene expression may be coordinated between more distant neighbors or between large spatial niches. The former is facilitated by changing the rank of the nearest neighbor tested. The latter is enabled by smoothing a gene's expression in a cell by averaging it with its expression in nearby cells, as shown for cortical layer 4 marker *Rorb* (Fig. 2e) and layer 6 marker *Foxp2* (Extended 188 Data Fig. 1f) in the auditory cortex³⁴.

 Although cross-expression may appear at varying length scales, we focus our analyses at the single cell level to investigate its signature at the finest resolution.

 Fig. 2 | Cross-expression analysis reveals coordinated gene expression between neighboring cells. a, Sagittal brain slices showing cortical somatosensory nose region (top) and visceral area (bottom) as randomly selected regions of interest. **b**, Neuropeptide somatostatin *Sst* and its cognate receptor *Sstr2* cross-express in regions shown in (a). Points indicate cells and colors indicate gene expression (left), with cross-expressing cell pairs highlighted (right). **c**, Bullseye scores for *Sst* and *Sstr2* in the regions shown in (a, b). The scores are reported as ratio of cross- to co-expression. **d**, Bullseye scores for cross-expressing (significant) and non-cross- expressing (not significant) gene pairs in the somatosensory nose region. 'Cell' corresponds to the central ring in (c), and the red rectangle highlights the first neighbor/ ring. Inset, ratio of bullseye scores for the first neighbor to the central cells for cross-expressing and non-cross- expressing genes. Central line, median; box limits, first and third quartiles; whiskers, ±1.5x interquartile range; points, outliers. **e**, Smoothed gene expression for different numbers of neighbors for the auditory cortical layer 4 marker gene *Rorb*. Created with BioRender.com.

Cross-expression is driven by subtle and consistent cell subtype compositional differences

 Having seen that cross-expression recovers coordinated spatial gene expression, we now explore 209 its relationship with cell type heterogeneity. For this purpose, we use another BARseq dataset 210 that was recently used to create a mouse cortical cell type atlas using the same 104 excitatory marker genes as before. Here, we observe that genes cross-express between cells of the same and of different types. For example, *Gfra1* and *Foxp2* are cross-expressed within the same cell type L4/5 IT (intratelencephalic) and between different cell types Car3 or CT (corticothalamic) and L4/5 IT (Fig. 3a). In general, genes vary greatly in terms of the cell type labels of cross-expressing cell pairs (Fig. 3b). For instance, for some gene pairs 40% of the cell pairs have the same cell type label while in others as many as 90% of the cell pairs belong to different cell types (Extended Data Fig. 2a). Moreover, some genes involve many while others involve few cross-expressing cells. For example, in the analyzed data the median number of cross-expressing cell pairs is 2,378, and 27% of genes involve over 4,000 while only 5% involve 400 or fewer pairs (Extended Data Fig. 2b), indicating that the density of gene cross-expression is highly variable. Interestingly, cell type purity – the proportion of cell pairs with the same type – decreases as more cell pairs 222 cross-express (Fig. 3c, Spearman's $\rho = -0.46$), highlighting a potential role for spatially intermingled cell types in patterns of cross-expression.

 To assess the influence of spatial cell type composition more broadly, we use our 225 hierarchical cell type atlas³⁰, where types at a higher-level divide into subtypes at a lower level. Using cross-expressing glutamatergic cells, we find that 64% of the pairs consist of different cell subtypes (Fig. 3d, right-tailed Wilcoxon signed-rank test, different labels ≥ same labels, p-value ≤ 0.0001, Extended Data Fig. 2c), suggesting that subtle cell type differences drive cross- expression. However, for cross-expressing GABAergic cells, we find that only 44% of the pairs have different cell subtype labels (Extended Data Fig. 2d-e right-tailed Wilcoxon signed-rank test, 231 different labels \geq same labels, p-value = 1), reflecting the fact that our gene panel is optimized to detect cell subtype differences between excitatory, but not inhibitory, neurons. Crucially, we observe that cells of one type consistently cross-express with cells of another type (Fig. 3e, Extended Data Fig. 2f), indicating that cross-expression recapitulates patterns of cell type composition. Since cell type labels are assigned based on the expression of many genes, 236 repeated spatial proximity of cell types is one mechanism that generates cross-expression.

 Fig. 3 | Cross-expression patterns are discovered independently of cell type labels but are driven by cell type heterogeneity. a, Cells of the same (yellow) and different (green) types cross-express genes *Gfra1* and *Foxp2* in the auditory cortex. Discovering cross-expression relations between this or any other gene pair does not require cell type labels. **b**, Numerous cells cross-express for each gene pair, with the dot size indicating the number of cell-neighbor pairs and the color showing the proportion of pairs with the same label (cell type purity). **c**, Cell type purity against the number of cross-expressing cell-neighbor pairs. Each point is a gene pair from (b), and shaded area is 95% confidence interval. **d**, Number of cell-neighbor pairs with the same or different cell subtype labels given that they were both labeled 'glutamatergic' at the higher level of the cell type hierarchy. Each point is a cross-expressing gene pair. **e**, Heatmap showing the normalized frequencies of cell type label combinations between cross-expressing cells. Created with BioRender.com.

 Cross-expression discovers anatomical marker combinations that delineate the thalamus and cortical layer VI

 Having found patterns of cross-expression within regions, we next tested for spatial organization that reflects anatomical structure. While some anatomical structures, such as cortical layers, have well-defined markers, others are difficult to characterize due to lack of marker genes. We asked whether cross-expressing genes can delineate anatomical regions. An important difference between cross-expression and co-expression is that the former will generally increase independence/dimensionality within the dataset while the latter will decrease it, providing a much larger scope for useful combinatorial markers. Since a gene panel of size *N* contains (259 Iarger scope for useful combinatorial markers. Since a gene panel of size *N* contains $\binom{N}{2}$ pairs, we reasoned that the quadratic space likely contains suitable marker combinations. To assess this, we used Vizgen's MERFISH (multiplexed error-robust fluorescent in situ hybridization) data (data availability section) obtained from coronal mouse brain slices, with a panel of 483 genes, yielding 116,403 gene pairs. We registered the brain slices to Allen Common Coordinate 264 Framework version 3 (CCFv3) atlas⁵⁷ to obtain region annotations for each cell, giving us a reference against which the marker-delineated regions could be compared.

 Surprisingly, we found that cross-expression between *Lgr6* and *Adra2b* delineates the thalamus even though these genes are widely expressed in the brain (Fig. 4a). Specifically, while 48% of *Lgr6*- and 57% of *Adra2b*-expressing cells are thalamic, 91% of their cross-expressing cell pairs are in the thalamus (Extended Data Fig. 3a), underscoring the spatial enrichment of their cross-expression signature (Extended Data Fig. 3b). We find that *Lgr6* also cross-expresses with *Ret* in the thalamus despite brain-wide expression of both genes (Fig. 4b, Extended Data Fig. 3c). Next, we examined whether *Adra2b* and *Ret*, both of which cross-express with *Lgr6*, show enriched co-expression in the thalamus. We find that they are indeed co-expressed within the thalamus but not in rest of the brain (Fig. 4c), e.g., 89% of their co-expressing cells are in the thalamus, thus serving as robust combinatorial markers.

 To evaluate whether the combinatorial marker-based approach is reliable, we asked whether single gene markers, when assessed for cross-expression, rediscover the anatomical 278 Iocations. Using the BARseq cortical cell type atlas data, we assessed cross-expression between cortical layer 6 marker *Foxp2* and ubiquitously expressed gene *Cdh13*. We discover that cross-expression between these genes delineates layer 6 boundary (Fig. 4d), further supporting the view that combinatorial anatomical markers can be discovered using cross-expression.

 Indeed, the layer 6 boundary recovered by cross-expression captures additional L6 IT neurons whereas *Foxp2*-based boundary overlooks these cells, indicating that combinatorial markers can refine extant anatomical regions. More generally, this process leverages the spatial enrichment of cross-expression, where the distance between cross-expressing cells is smaller than the distance between cross-expressing and randomly selected cells. Once spatial enrichment is discovered, our framework can help refine anatomical regions and link them to patterns of coordinated expression across cells that are independent of co-expression.

 Fig. 4 | Cross-expression can discover combinatorial anatomical markers. a, Comparing the thalamus (left) to the rest of the brain, genes *Lgr6* and *Adra2b* are widely expressed across multiple brain regions (middle) but are preferentially cross-expressed in the thalamus (right). **b**, Same as in (a) but for genes *Lgr6* and *Ret*. **c**, Genes cross-expressing with *Lgr6* in (a) and (b) co-express in the thalamus. **d**, Cross-expression of *Cdh13* with cortical layer 6 marker *Foxp2* (middle) recapitulates layer 6 boundaries (right, cf. left). Created with BioRender.com.

Cross-expression network reveals *Gpr20* **as a central gene and discovers possible interaction partners between astrocytes and the brain microvasculature**

 So far, we have assessed cross-expression between gene pairs to discover ligand-receptor interactions, cell type differences, and anatomical markers. However, each gene may cross- express with many others and thus form clusters of genes with coordinated expression. These relationships can be analyzed using networks, where nodes represent genes and edges indicate cross-expression (Fig. 5a, left). Within a network, if nodes A and B connect to node C but not to each other, they form a second-order edge (Fig. 5a, right). Both types of relationships are important, as in genetic interaction networks, because genes are joined not only by similarity but also by a form of complementarity. Representing cross-expression as a network is therefore a potentially powerful formalism, especially because it allows for the application of a substantial body of existing gene network methods.

 Using the MERFISH data, we created a cross-expression network (Fig. 5b), which contains 200 genes with 382 first-order, 217 second-order, and 107 dual-order edges. We observe that *Gpr20*, a G protein-coupled receptor, is a central gene with a high node degree of 40 while the other genes form a median of 4.8 edges (Extended Data Fig. 4a). We performed gene ontology (GO) enrichment for genes cross-expressed with *Gpr20*, finding functional groups like 'regulation of macromolecule biosynthetic process', 'regulation of gene expression', and 'regulation of metabolic process' (Extended Data Fig. 4b, all p-values ≤ 0.05). While some of these genes are co-expressed with astrocytic and microglial cell type markers (Extended Data Fig. 4c), their global co-expression with the endothelial marker is higher, where the co-expression profiles were computed using neighbors cross-expressed with *Gpr20* rather than the entire dataset (Extended Data Fig. 4d, Mann-Whitney U test, endothelial vs others, all p-values ≤ 0.01; remaining pairwise comparisons, all p-values > 0.05).

 Noting that the neighbors of *Gpr20*-positive cells are involved in the microvasculature, we next viewed the spatial distribution of cells expressing *Gpr20*, finding that they form contiguous linear streaks resembling blood vessels (Fig. 5c; anterior slice from mouse brain 1 shown). To test this observation, we looked at whether the neighbors of *Gpr20*-positive cells also express this gene and compared it to randomly selected cells, which constitute the expectation that *Gpr20* is uniformly expressed across space. Consistent with the visualization, we find that cells with *Gpr20* are surrounded by neighbors that also express this gene, a pattern that disappears for neighbor order of 50 or more cells (Fig. 5d-e, area under curve (AUC), neighbors vs random cells, 0.69 vs 0.49; right-tailed Wilcoxon signed-rank test, neighbors vs random cells, p-value ≤ 0.0001). Having seen that cells with *Gpr20* possibly reflect blood vessels, we asked whether these cells are

 themselves vascular or whether they line the vasculature, especially since the cells that cross- express with *Gpr20* are endothelial. We find *Gpr20* is poorly co-expressed with *Igfr1* (Pearson's $R = 0.0024$), the vascular/endothelial marker^{58–60} in our gene panel, suggesting that it lines but does not mark the blood vessels. Moreover, it is lowly co-expressed with other cell type markers (average Pearson's *R*, astrocytes = –0.0027, microglia = 0.0018, oligodendrocytes = –0.022, neurons = –0.0025), eschewing cell type characterization. Taken together, *Gpr20*, a salient topological feature of our cross-expression network, seems to be expressed in diverse cell types that line the blood vessels, reflecting its possible role in the modulation of the microvasculature.

 Cross-expression driven by cell types might be particularly common when two genes which cross-express with a third gene are co-expressed together, reflecting some common transcriptional program jointly cross-expressing with neighboring cells. To investigate this, we reduced co-expression further by specifying that cross-expressing genes must show lack of significant co-expression, a procedure that yielded a subnetwork, which we further curated by removing genes with fewer than two edges. Indeed, we find that two genes that independently cross-express with another gene tend to be co-expressed (Fig. 5f, Extended Data Fig. 5a) and, as expected, belong to the same cell types, as revealed by their co-expression with cell type marker genes (Extended Data Fig. 5b). Confirming these results, the subnetwork genes are enriched in GO groups like 'endothelial cell proliferation', 'positive regulation of vascular endothelial growth factor production', and 'regulation of endothelial cell migration' (Fig. 5g, all p- values ≤ 0.05). These results indicate that while cross-expressing genes are present in specific cell types, the relations between them are functionally suggestive as opposed to simply reflecting cell type compositional differences, especially since the cell type markers are not cross- expressed. For example, the astrocytic *EGFR* (epidermal growth factor receptor) cross-expresses with the vascular *Flt4*/*VEGFR-3* (FMS-like tyrosine kinase 4), *Tek*/*Tie2* (TEK tyrosine kinase/ angiopoietin-1), and *Tie1* (tyrosine kinase with immunoglobulin-like and EGF-like domain 1). These three vascular receptors promote angiogenesis via the *VEGF* (vascular epidermal growth 356 factor) ligand^{61,62}, prevent endothelial cell apoptosis^{63,64}, and negatively regulate angiogenesis⁶⁵, respectively, thus reflecting their potential role in the brain microvasculature in coordination with the astrocytes, whose endfeet ensheathe the blood microvessels to constitute the blood-brain barrier (BBB).

 Within the same subnetwork, the astrocytic gene *Ppp1r3g* (protein phosphatase 1 361 regulatory subunit 3G), which helps convert glucose to glycogen⁶⁶, cross-expresses with *Epha2* 362 (ephrin type-A receptor 2), whose activity makes the BBB more permeable⁶⁷, likely enabling glucose's transport and eventual conversion into glycogen, thereby making this cross-expression relation relevant for energy metabolism. Indeed, this observation can be used to generate hypotheses about the (directional) relationship between energy needs within a local microenvironment and the remodeling of the microvasculature, making cross-expression a powerful approach with which to form testable hypotheses. More broadly, the cross-expression framework can be combined with well-established approaches, such as network analysis, to generate biological insights from high-throughput spatial transcriptomics data.

 Next, we asked whether cross-expression networks change across the brain. Because gene expression is regional, slices from various areas should show cross-expression between distinct genes. We assessed this by forming networks for each slice in our sagittal BARseq data. As expected, we find that adjacent slices have similar networks than distant slices (Fig. 5h, 374 Spearman's $\rho = -0.9$), a trend also seen in our BARseq coronal data (Extended Data Fig. 6a, 375 Spearman's $\rho = -0.87$) but not when the two datasets are mixed and the "distance" reflects the 376 difference in the order of slices (Extended Data Fig. 6b, Spearman's ρ = 0.094). Hence, cross-expression is sensitive to broad spatial variation in gene expression.

 Fig. 5 | Networks of cross-expression. a, Cross-expression (edges) between genes (nodes) forms a network (left), where second-order edges (right) between genes share a node (first-order). **b**, Example cross-expression network, with first-order node degree represented by size and edge color showing first-, second-, or dual-order (first-order and second-order) connections. Threshold for the second-order edges is 4. Node color shows community membership assigned by Louvain clustering the second-order network. **c**, Cells are colored based on *Gpr20* expression. Numbered rectangles in the central figure correspond to zoomed-in versions. **d**, Number of neighbors with *Gpr20* given that the source cells also express this gene. **e**, Cumulative sums (after L1 normalization) from (d) for true and randomly selected neighbors. Identity line is dashed. **f**, Subnetwork created from (b) by pruning edges with significant co-expression and then removing

 nodes with degree 1. Nodes are colored by cell types based on their co-expression with marker genes. **g**, GO functional groups for genes in the subnetwork in (f). **h**, Similarity in the network structures of nearby and distant brain slices. Shaded area is 95% confidence interval. Created with BioRender.com.

Cross-expression signal is replicable across datasets, and global co-expression between spatial and single cell datasets indicates reliable cell segmentation

396 Two sources of non-biological variation in spatial transcriptomics^{2–7} are batch effects, which result from technical differences between experimental runs, and cell segmentation, which draws boundaries around and assigns transcripts to cells, a process that can alter gene expression profiles and affect downstream analysis, including cross-expression.

 We assess batch effects by comparing cross-expression between corresponding slices across biological replicates. The MERFISH data contains three replicates with three slices each, where the slices are sampled from roughly the same position. We find that the cross-expression signature is highly similar across replicates. For example, the average correlation for the anterior slices between the three replicates is 0.83 (Fig. 6a), with similar findings for the middle and 405 posterior slices (Extended Data Fig. 7a-b, Spearman's $\rho = 0.81$ and 0.8, respectively).

 We next assessed the degree to which cross-expression within the BARseq sagittal or coronal experiments³⁴ is similar to that between experiments. To this end, we compared cross- expression patterns between brain slices. As expected, the cross-expression profiles are more similar within brains than between brains (Fig. 6b, Mann-Whitney U tests, FDR-corrected, coronal 410 vs sagittal, p-value = 0.2, coronal vs mixed, p-value \leq 0.001, and sagittal vs mixed, p-value \leq 0.001), suggesting that the sectioning procedure samples different brain regions and therefore reveals distinct underlying gene expression profiles. Supporting this result, we find that the same 413 anatomical regions (per Allen CCFv3 brain atlas⁵⁷) across brains have more similar cross- expression profiles than do different regions within or between brains (Fig. 6c, Mann-Whitney U 415 test, different regions vs same regions, p-value \leq 0.001). Noting that the sagittal and coronal brains contain the same regions in the dorsal to ventral directions, we asked whether the cross- expression is similar in this shared dimension. Here, we computed the density of cross-expressing cells in the dorsal to ventral direction and compared these distributions across the two brains, finding that 99% (without FDR correction) of the genes did not have significantly different density profiles (Fig. 6d), suggesting that the cross-expression patterns are highly similar across batches at the whole-brain level.

 Having found that the cross-expression profiles are generally robust, we assessed cell segmentation at a global level by comparing gene co-expression between the single cell RNA-

424 sequencing (scRNA-seq)³¹ and spatial transcriptomic data. We reasoned that scRNA-seq does not require segmentation and therefore captures genes co-expressed within the cell's boundaries (Fig. 6e). Because cell segmentation alters transcript assignment, it could change co-expression in spatial transcriptomic data. Reassuringly, we find a strong association between gene co- expression in the scRNA-seq and spatial transcriptomic data (Fig. 6f, Pearson's *R* = 0.83). We further examine whether this correlation is sufficiently strong by comparing co-expression 430 between scRNA-seq and single-nucleus RNA-sequencing (snRNA-seq)⁶⁸ (Fig. 6g, Pearson's $R =$ 0.86), finding agreement between the two comparisons (*R* = 0.83 vs. *R* = 0.86). These results imply similar levels of technical variability between platforms while suggesting that gene co-expression is congruent between scRNA-seq and spatial transcriptomic data.

434 The data in our work was processed using CellPose⁶⁹, a deep learning-based cell 435 segmentation algorithm. A recent benchmarking study⁷⁰ showed that it outperforms other methods on a variety of metrics. In fact, it uses the nuclear stain DAPI as a cell landmark and forms boundaries using cytoplasmic signal, such as the transcript distributions, making it the state-of- the-art segmentation algorithm on a variety of assessments. Further, the cell segmentation 439 algorithms are continuously being improved⁷¹, allowing users to re-segment and reanalyze their data. Most importantly, the analysis conducted using the cross-expression framework may suffer if segmentation is performed poorly, but the validity of the concept and the soundness of its statistics do not rely on this potential artefact and, with rapid improvements in data quality, the 443 inferences drawn from it will become increasingly more reliable.

 Moreover, we assessed the relationship between cross-expression and noise in gene expression measurement. Since the algorithm requires binarizing the expression matrix, an appropriate threshold needs to be applied prior to analysis. To count a gene as expressed in a cell, we applied thresholds of 1 to 10 molecules, finding that the cross-expression patterns are generally concordant across these noise levels (Extended Data Fig. 8a-b, median Pearson's *R* = 0.88). Importantly, our framework is agnostic to and compatible with multiple models of gene 450 expression noise⁷², and once an appropriate threshold has been applied, the resultant expression matrix can be used for cross-expression analysis.

 Finally, we explored the patterns of cell-neighbor relations and found that over 60% of cells are the nearest neighbors of exactly one cell but the remaining cells are the nearest neighbors of two or more cells (Extended Data Fig. 9a). Patterns such as these may be biologically important if the 'neighbor' cell plays a central role in the local microenvironment, so deviations from one-to- one mappings should be captured by statistical analyses. To investigate that our results are consistent across these patterns, we compared cross-expression in one-to-one against many-to one mappings and with the full dataset, finding an average Pearson's correlation of 0.96 (Extended Data Fig. 9b). Importantly, our procedure is consistent with the assumption of independent sampling because while a cell may be the nearest neighbor of multiple cells, each cell-neighbor pair is statistically independently.

 We enable these analyses by providing a highly efficient R package. A laptop with 16 GB RAM can test for cross-expression in large datasets containing hundreds of thousands of cells and thousands of genes within minutes (Fig. 6h). At present, most (commercial) imaging-based 465 platforms cannot profile gene panels of this magnitude $2-7$, though such capabilities are anticipated 466 and are being developed⁷³. Our software's performance makes it well-suited for analyzing current and future spatial transcriptomic datasets.

 Fig. 6 | Assessing batch effects, cell segmentation, and software runtime. a, Correlation between cross-expression signatures across three biological replicates. **b**, Correlation between cross-expression signatures within (sagittal or coronal) and between (mixed) brains. Positive signal between brains likely reflects the fact that the sagittal and coronal brains both contain regions in the dorsal to ventral direction. **c**, Correlation between cross-expression signatures between the same anatomical regions across brains or between different anatomical regions

 across or within brains. **d**, Density of cross-expressing cells in the dorsal to ventral directions is compared across the sagittal and coronal brains. Significant p-values (without FDR correction) indicate that a cross-expressing gene pair has different densities across the two brains. Red dotted line is the significance threshold at alpha = 0.05. **e**, Single cell RNA-sequencing (scRNA- seq) profiles cells' gene expression without cell segmentation. Co-expression between scRNA- seq and spatial transcriptomic data helps diagnose segmentation artifacts. **f**, Gene co-expression in spatial transcriptomic and in scRNA-seq data. Each point is a gene pair. **g**, Gene co-expression in single-nucleus RNA-sequencing (snRNA-seq) and in scRNA-seq data. Same gene panel is used in (f) and (g). **h**, Software runtime for varying numbers of genes and cells using a personal laptop with 16 GB RAM. Created with BioRender.com.

Discussion

 Cross-expression allows us to study gene-gene networks that reflect how cells influence each other through coordinated gene expression between neighboring cells. Using this framework, we recapitulated known ligand-receptor interactions at the single cell level, revealing biologically meaningful tissue phenotypes. We further showed that cross-expression can be discovered without cell type labels but often reflects cell subtype compositional differences. Moreover, it helps us discover paired markers for anatomical regions, such as the thalamus, and is amenable to network formulations, finding genes like the *Gpr20* as central nodes and revealing the relationships between astrocytes and brain microvasculature. Together, cross-expression is a powerful way of analyzing spatial transcriptomic data and allows us to study gene-gene relations between adjacent cells, thereby fully harnessing the high-throughput of these technologies.

 The cross-expression framework complements current approaches analyzing spatial 498 transcriptomic data, such as those exploring niche-specific co-expression patterns^{12–16}. Specifically, niche-specific cross-expression networks may be compared with co-expression networks to examine if inter-cellular relations are associated with intra-cellular gene programs and vice versa. This may be approached at different, potentially hierarchical spatial scales to reveal spatial gene expression programs within the tissue. Moreover, the cross-expression patterns can be quantified in multiple ways, such as using mutual information or graphlets, allowing investigations into the best approaches that capture the signal of interest. For example, just as co-expression relations can be measured using the Pearson's correlation coefficient, cross-expression patterns may be investigated from numerous perspectives to discover the most robust formalism. In this sense, the cross-expression framework introduced here is primarily a novel way of conceptualizing gene-gene relations within spatial transcriptomics data, thereby serving as a powerful framework for research in tissue biology. For instance, it can be used to 510 study cancer⁷⁴, where tumor progresses via signaling with the stromal tissue, as well as 511 neurodegenerative diseases like Alzheimer's⁷⁵ or senescence⁷⁶, where the progression of pathology is spatially mediated, making it a broadly useful approach for numerous problems.

 Cross-expression is not restricted to imaging-based spatial transcriptomics. Instead, it can be applied to any biological assay that provides cell-by-features and cell-by-coordinate matrices. 515 For example, it can be extended to spatial proteomics⁷⁷, with potential to discover ligand-receptor 516 interactions. Likewise, it may be applied to spatial translatomics⁷⁸ to focus on translating mRNAs that are more likely to form functional proteins, making conclusions about cell-cell relations more 518 robust. In fact, with the increasing resolution of spatially barcoded RNA capture based methods^{3–} \textdegree , the framework may be extended transcriptome-wide to understand relations between spots at near single cell resolution.

521 A key challenge in imaging-based spatial transcriptomics²⁻⁷, including the datasets used in this work, is the size and constitution of the gene panel, which sets an upper limit on biological discovery. Although our framework will become more powerful as the quality of spatial transcriptomic data, especially the gene panel, increases, care must be taken to not interpret the results in mechanistic terms. Instead, the coordinated gene expression between neighboring cells should be viewed as a target for experimental validation. In this sense, the cross-expression framework radically narrows the space of gene-gene relations by identifying pairs that are potentially biologically meaningful, making the problem experimentally tractable. Overall, cross- expression is a powerful addition to the growing list of analytical techniques and, most importantly, it offers a unique perspective on using spatial transcriptomic data for driving biological discovery.

Extended Data Figures

 Extended Data Fig. 1 | Cross-expression across tissue slices and regions for ligand- receptor and non-signaling genes. a, Distribution of the number of gene pairs cross-expressed in different slices. Dataset has 15 slices sampled sagittally from the left hemisphere of a mouse brain. **b-e**, Cells are colored by gene expression (left) and cross-expressing cells are highlighted (center). Right, distances between cross-expressing cells are compared with those between cross-expressing and randomly selected cells. Smaller distances mean that cross-expressing cells are nearer each other (spatial enrichment) than expected by chance (p-values ≤ 0.01, left- tailed Mann-Whitney U test). Genes include ligands and receptors (b, c) and non-ligands and non- receptors (d, e). **f**, Smoothed gene expression for different numbers of neighbors for the auditory cortical layer 6 marker gene *Foxp2*. Created with BioRender.com.

543

Extended Data Fig. 2 | Relationship between cross-expression and cell type heterogeneity.

 a, Proportion of cross-expressing cell pairs belonging to the same cell type label. **b**, Number of cell-neighbor pairs involved in cross-expression. **c**, Proportion of cell-neighbor pairs with different cell subtype labels given that both were labeled 'glutamatergic' at the higher level in the cell type hierarchy. **d**, Number of cell-neighbor pairs with the same or different cell subtype label given that both were labeled 'GABAergic' at the higher level in the cell type hierarchy. Each point is a cross- expressing gene pair. **e**, Same as in (c) but for 'GABAergic' cells. **f**, Proportion of neighbor cell types against which cell types cross-express. Created with BioRender.com.

 Extended Data Fig. 3 | Combinatorial anatomical markers discovered using spatially enriched cross-expression. a, Proportion of *Adra2b*- and *Ret*-expressing cells in the thalamus (red) and the proportion of cell pairs in the thalamus (blue) when cross-expressing with *Lgr6*. **b-c**, Distances between cross-expressing cells versus those between cross-expressing and 558 randomly chosen cells for genes Lgr6 and Adra2b (c) and for Lgr6 and Ret (d). Smaller distances randomly chosen cells for genes *Lgr6* and *Adra2b* (c) and for *Lgr6* and *Ret* (d). Smaller distances mean that cross-expressing cells are nearer each other (spatial enrichment) than expected by chance (p-values ≤ 0.01, left-tailed Mann-Whitney U test). Created with BioRender.com.

 Extended Data Fig. 4 | Exploration of *Gpr20* **and its cross-expressing genes. a**, Distribution of node degree, with *Gpr20* highlighted. **b**, Gene ontology (GO) functional groups for genes cross- expressed with *Gpr20*. **c**, Co-expression of genes cross-expressed with *Gpr20* (right) against cell 565 type marker genes (bottom). For each gene, co-expression was computed using cells involved in
566 cross-expression and not the entire dataset. **d**, Distribution of cell type marker genes' co- cross-expression and not the entire dataset. **d**, Distribution of cell type marker genes' co-expression across the genes in (c). Created with BioRender.com.

Extended Data Fig. 5 | Exploration of the MERFISH cross-expression (sub)network. a, Co-

expression of genes in the subnetwork. **b**, Co-expression between genes in the subnetwork (right)

and cell type marker genes (bottom). Created with BioRender.com.

 Extended Data Fig. 6 | Cross-expression network similarity between slices. a, Slice-specific cross-expression networks compared and shown as a function of distance between slices. **b**, Same as in (a) but slice-specific networks compared between sagittal and coronal datasets, where the "distance" is the difference in slice ID's. Shaded areas are 95% confidence intervals. Created with BioRender.com.

579 **Extended Data Fig. 7 | Cross-expression network similarity between replicates. a-b**, Cross-580 expression networks compared between three replicates for the middle (a) and posterior (b) 581 slices. Created with BioRender.com. slices. Created with BioRender.com.

582

 Extended Data Fig. 8 | Cross-expression network similarity at different levels of gene expression noise thresholds. a, Cross-expression networks compared after applying different noise thresholds, which are the minimum number of molecules a gene must express within a cell to be considered as detected. **b**, Distribution of the network similarities across noise levels, with the median indicated using the dotted line. Created with BioRender.com.

588

 Extended Data Fig. 9 | Patterns of cell-neighbor mappings and their relationship with cross-expression. a, Cells considered as 'nearest neighbor' by other cells reported as a proportion of total cell-neighbor relations. '1' is one-to-one mapping and '2-4' is many-to-one mapping. **b**, Cross-expression networks computed using one-to-one mappings, many-to-one mappings, and the full dataset (both mappings). Created with BioRender.com.

594

595 **Online Methods**

596 We first explain the theoretical underpinnings of our approach and outline the features of the 597 associated R package. We then specify how these are used in various analyses.

598

599 **Statistics of cross-expression between a gene pair**

 Cross-expression is the mutually exclusive expression of a gene pair across neighboring cells. To assess whether gene A's expression in cells and gene B's expression in their spatial neighbors is significant, we use a simple sampling procedure and model the probabilities using the hypergeometric distribution

$$
P(X = k) = \frac{\binom{k}{k} \binom{N-K}{n-k}}{\binom{N}{n}}
$$
\n(1)

 where *N* is the population size, *K* is the number of successes (or success states), *n* is the number of samples or draws, and *k* is the number of observed successes. The form $\binom{a}{b}$ $\;$ of samples or draws, and k is the number of observed successes. The form $\binom{a}{b}$ is the binomial coefficient giving us the number of distinct *b*-sized groups from *a*-entries.

Equation 1 outlines all the ways in which success can be observed—($\frac{K}{\nu}$ 607)—*and* (product rule) all the ways in which failure can be obtained— $\binom{N-K}{n-k}$ 608 $-$ rule) all the ways in which failure can be obtained— $\binom{N-K}{n-k}$ —normalized by all possible ways of generating our sample $\binom{N}{n}$ 609 is generating our sample $\binom{N}{n}$, making the outcome probabilistic by bounding it between [0,1].

 Traditionally, the *n* samples are assessed for the presence of some property *k*. Here, we sample cell-neighbor *pairs* conditioned on the cell expressing gene A and ask whether the neighbor expresses gene B. Thus, the sample size *n* is the number of cells with gene A, the number of observed successes *k* is the number of neighbors with gene B whose corresponding cells express gene A, and the number of success states *K* is the total number of neighbors with gene B. The population size *N* is the total number of cells, including those that co-express A and B and those that express neither gene. We are interested in the probability of observing *k* or more neighbors with B when *n* cells with A are sampled. To this end, we modify the hypergeometric cumulative distribution function (CDF)

$$
P(X \ge k) = 1 - P(X < k) = 1 - \sum_{i=0}^{k-1} \frac{\binom{K}{i} \binom{N-K}{n-i}}{\binom{N}{n}}
$$
\n(2)

619 to calculate the probability of *k* or more successes. A value lower than alpha α indicates an 620 unusually large number of neighbors expressing gene B when cells expressing gene A are 621 sampled, implying statistically significant cross-expression between this pair.

622

623 **Statistics of cross-expression between all gene pairs**

We need to assess cross-expression across all gene pairs, which rise quadratically by $\binom{N}{2}$ 624 We need to assess cross-expression across all gene pairs, which rise quadratically by $\binom{N}{2}$ or $N(N-1)$ 625 $\frac{N(N-1)}{2}$ for *N* genes. For example, a panel with 500 genes contains around 125,000 pairs whereas 626 a panel with 1,000 genes has approximately 500,000 pairs. To efficiently explore this space, we 627 implement the procedure above using matrix operations and specialized packages in R.

 We begin with a cells-by-genes expression matrix **E** and a cells-by-coordinates location matrix **L**, where the coordinates in our data are cell centroids on two-dimensional slices. We input 630 L into RANN package's function^{79,80} nn2 with search type as standard, which implements a kd- tree (or optionally a bd-tree) search algorithm to explores data subspaces and efficiently find the *n*-th neighbors. Using the neighbor indices, we re-order the expression matrix **E** to generate the neighbors-by-genes matrix **E′**. The value of *n* can be changed to generate paired gene expression matrices, where the corresponding rows represent cells and their *n*-th nearest neighbors.

 Our aim is to use **E** and **E′** to compute *N* (population), *K* (neighbors with B), *n* (cells with A), and *k* (neighbors with B when their corresponding cells express gene A) for each gene pair. 637 These four values can be inputted into R's $_{\rm phyper}$ function for all gene pairs, facilitating efficient computation. The population size *N* is the total number of cells and is the same across all pairs. To compute *n*, we binarize **E** based on expression or lack thereof, and compute co-occurrences using the dot product

$$
\mathbf{C} = \mathbf{E}^T \cdot \mathbf{E} \tag{3.1}
$$

641 where \mathbf{c}_{ii} is the number of cells expressing gene *i* and \mathbf{c}_{ii} (for $i \neq j$) is the number of cells co-642 expressing genes i and j . We perform

$$
\mathbf{U}_{ij} = \mathbf{C}_{ii} - \mathbf{C}_{ij} \tag{3.2}
$$

643 where U_{ij} is the number of cells uniquely expressing gene i. We implement this by extracting the 644 diagonal of C, and "broadcast" it against its off-diagonal entries, thus aligning the corresponding 645 values before subtraction. For each pair, this gives us the number of cells *n* uniquely expressing 646 each gene. We perform an analogous calculation for *K* using **E′** instead of **E**, giving us the number 647 of neighbors uniquely expressing each gene within a gene pair.

 We now turn to *k*, the number of neighbors observed with gene B given that their corresponding cells express gene A. Using binarized matrices **E** and **E′**, we compute the number of cell-neighbor pairs such that the cells express gene A without gene B and the neighbors express gene B without gene A

$$
\mathbf{X} = (\mathbf{E} \odot (1 - \mathbf{E}')) \tag{4.1}
$$

$$
\mathbf{Y} = ((1 - \mathbf{E}) \odot \mathbf{E}')
$$
 (4.2)

$$
\mathbf{Q} = \mathbf{X}^T \cdot \mathbf{Y} \tag{5}
$$

652 where \odot is the Hadamard (elementwise) product and ${\bf Q}_{ij}$ is the number of cell-neighbor pairs with 653 mutually exclusive expression. In X, E contains '1' in cells where a gene is expressed and $1 - E'$ 654 contains '1' in neighbors where a gene is *not* expressed. Their elementwise product has '1' to 655 indicate genes' presence in cells and their absence in neighbors. **Y** shows the analogous 656 procedure for genes' presence in the neighbor and their absence in cells. Hence, the dot product 657 of **X** and **Y** gives Q, a genes-by-genes asymmetric matrix, whose entries show the number of cell-658 neighbor pairs with mutually exclusive expression. (Q is asymmetric because the number of cell-659 neighbor pairs in the A-to-B and B-to-A directions are not always identical.) This is *k* or observed 660 successes. These steps generate four number – *N*, *K*, *n*, and *k* – per gene pair. We input these 661 into R's phyper function in accordance with equation (2), giving us corresponding p-values.

662 Since Q is asymmetric, we obtain two p-values per gene pair, one in the A-to-B and the 663 other in the B-to-A direction. We perform Benjamini-Hochberg⁸¹ false discovery rate (FDR) multiple test correction on the entire p-value distribution. For each gene pair, we then assess whether or not cross-expression is observed in either direction and use the lower FDR-corrected p-value as the final output, which is provided both as an edge list and as a gene-by-gene p-value matrix **P**.

668

669 **Cross-expression networks**

- 670 We can threshold and binarize **P** at a pre-selected alpha α to form an adjacency matrix **N**, where
- 671 '1' indicates connections (edges) between genes (nodes)

$$
\mathbf{N}_{ij} = \begin{cases} 1 & \mathbf{P}_{ij} \le \alpha \text{ and } i \ne j \\ 0 & \text{otherwise} \end{cases}
$$
 (6.1)

672 This allows us to perform cross-expression network analysis, where higher-order 673 community structure is discovered using shared connections between genes

$$
\mathbf{S}^R = \mathbf{N}^1 \cdot \mathbf{N}^2 \cdot \dots \cdot \mathbf{N}^{R-1} \cdot \mathbf{N}^R
$$
 (6.2)

- 674 where we restrict *R* = 2 to discover second-order connections between genes.
- 675

676 **Cross-expression at multiple length scales**

 Cross-expression is coordinated gene expression between neighboring cells. Yet, these patterns may be present at larger length scales, requiring us to understand associations between regions. To facilitate this, we smooth the expression of each gene in a cell by averaging it with its expression in *n* nearby cells. Using the RANN package, we find the indices of each cell's *n* nearest neighbors, and make the corresponding values '1' in the cells-by-cells matrix **C**

$$
\mathbf{C}_{ij} = \sum_{k=1}^{s} \mathbb{I}\left\{(i,j) = (i_k, j_k)\right\}
$$
 (7.1)

682 where the indicator function $\mathbb I$ specifies

683
$$
\begin{cases} 1 & \text{if } (i,j) = (i_k, j_k) \\ 0 & \text{otherwise} \end{cases}
$$

684 and

$$
s = c + (c \times n) \tag{7.2}
$$

685 where *c* is the number of cells and *n* is the number of neighbors. Here, *s* is the total number of 686 row-column indices *i*-*j* that *k* iterates over. We perform averaging using the expression matrix **E**

$$
\mathbf{S} = \frac{1}{n} (\mathbf{C} \cdot \mathbf{E}) \tag{8}
$$

687 where S_{ij} is the *j*-th gene's average value in *i*-th cell across *n* neighbors. The smoothed gene 688 expression matrix **S** can be used for downstream analysis.

689

690 **Bullseye scores as effect size**

691 The bullseye scores quantify the effect size by comparing cross-expression with co-expression.

692 Here, the number of neighbors with gene B is compared to the number of cells co-expressing

693 genes A and B. We use binarized cell and neighbor expression matrices **E** and **E′**, respectively

$$
\mathbf{B}_n = \mathbf{E}^T \cdot \mathbf{E}_n' \tag{9.1}
$$

 where *n* is the *n*-th neighbor, giving us *n* gene-by-gene asymmetric matrices **B***n*. The *i*-th and *j*-th entries of **B***ⁿ* indicate the number of *n*-th nearest neighbors expressing gene B when cells in **E** 696 express gene A. B_n is a co-occurrence matrix when $n = 0$. Viewing B_n as a tensor with dimensions *i*, *j*, and *n*, for each gene pair we take the cumulative sum and normalize across the neighbors

$$
\mathbf{B}_{ijn} = \frac{1}{n} \sum_{n'=1}^{n} \mathbf{B}_{ijn'} \quad \text{for } n \ge 1
$$
 (9.2)

698 These matrices can be compared with $B_{n=0}$ to find the ratio of cross-expression to co-699 expression and/or $log₂$ -transformed for further analysis. The output is provided as an array of 700 matrices (tensor) or as an edge list, where columns represent different *n* neighbors.

701

702 **Expression of gene pairs on tissue**

 A powerful way of viewing cross-expression is to plot the cells and color them by their gene expression. For a gene pair, a cell can express genes A, B, both, or neither. We make these plots for user-selected gene pairs using the expression matrix **E** and the cell coordinates matrix **L**. We can also exclusively highlight cross-expressing cell-neighbor pairs. Finally, the tissues are often not upright, partly due to their misorientation with respect to the glass slide, making it difficult to interpret the results. Accordingly, we rotate them using user-defined *n*-degree

$$
\theta = n_{degrees} \times \frac{\pi}{180} \tag{11.1}
$$

$$
x' = \cos(\theta) \cdot x - \sin(\theta) \cdot y \tag{11.2}
$$

$$
y' = \sin(\theta) \cdot x + \cos(\theta) \cdot y \tag{11.3}
$$

- 709 where *x*′ and *y*′ are the cell coordinates after counterclockwise rotation. Rotation does not change 710 the distances between cells, so *x*′ and *y*′ can be used for downstream analysis.
- 711

712 **Spatial enrichment of cross-expression**

 Cross-expressing cells may be distributed across the tissue or show spatial localization. To quantify their enrichment, we first average the distance between cell-neighbor pairs. We next compare the distances between all cross-expressing cells to the distances between cross- expressing and randomly selected cells. If the former distance is significantly smaller than the latter distance, then cross-expression is spatially enriched.

718

719 **Data acquisition and preprocessing**

720 *MERFISH brain receptor map data*

 We obtained Vizgen MERSCOPE's mouse brain receptor map from https://info.vizgen.com/mouse-brain-data. This data contains three coronal slices from three replicates, with the middle slice covering the center of the brain. We analyzed slice 2 from replicate 2, which contains 483 genes and 84,172 cells. We filtered cells with fewer than 50 counts and those lacking brain region annotations (see below), leaving around 82,000 cells. The gene panel consists of cell type markers, G protein coupled receptors (GPCRs), and receptor tyrosine kinases (RTKs). We registered the slice to the Allen CCFv3 (Common Coordinate Framework version 3) 728 brain region atlas⁵⁷. To facilitate this, we annotated the cells using Seurat⁸². Here, we created a 729 Seurat object and used SCTransform with the clip.range between -10 and 10. We then ran Principal Component Analysis (PCA), setting the number of components to 30 and specifying the 731 features as genes. Next, we used FindNeighbors and FindClusters with the resolution set 732 to 0.3. The clusters are cell type labels, which help us identify brain structures during registration. 733 (These labels were not used for any analyses.) For registration, we used QuickNii⁸³ (v3 2017) to linearly align the slice to the Allen CCFv3 atlas using discernible regions like the hippocampus 735 and the ventricles as anchors. We then used VisuAlign⁸³ (v0.9) to non-linearly transform the slice to improve alignment with the atlas. This procedure assigns a brain region annotation to every cell. Finally, we rotated the image 40 degrees counterclockwise to make it upright.

 The entire dataset contains 3 replicates with 3 slices each (anterior, middle, posterior), yielding a total of 734,647 cells that we used for additional analyses.

BARseq data

The BARseq data was collected in an effort to create a mouse brain cortical cell type atlas³⁴. Its 104 genes consist largely of excitatory cell type markers (109 total genes), and its 1,161,387 cells were sampled across 40 slices. The cells were iteratively clustered into H1, H2, and H3 types, providing a hierarchical cell type atlas. The H2 types were used during brain registration, which was performed as described above. We filtered cells expressing fewer than 5 genes or with less than 20 counts.

 We also collected a sagittal mouse hemi-brain data (P56 male) from the left hemisphere (20µm thick sections, 300µm distance between slices) with the same gene panel as the coronal data but with 24 additional ligand-receptor pairs (neuropeptides, neuropeptide receptors, monoamine receptors such as cholinergic, adrenergic, serotonergic, and dopaminergic). This data yielded 133 genes assayed across 1,311,001 cells spanning 16 slices. It was collected for 753 this project and was processed similarly to the coronal data³⁴. All experimental procedures were

 carried out in accordance with the Institutional Animal Care and Use Committee at the Allen Institute for Brain Science.

Single-cell RNA-seq (scRNA-seq) and single-nucleus RNA-seq (snRNA-seq) data

758 We obtained the scRNA-seq³¹ and snRNA-seq⁶⁸ data from the Brain Initiative Cell Census Network (BICAN) cell type atlases. These data were collected from dissected tissue regions, giving us the cells' coarse anatomical origin. We removed cells with a doublet score of 30 or above and randomly selected 10,000 cells from each region for subsequent analysis.

Ligand-receptor cross-expression

 We aimed to find cross-expression between known ligand-receptor pairs. In our sagittal data, we selected two slices and within each slice we chose a cortical region. These choices were made randomly. In practice, we chose the visceral area (VISC) in slice 3 and the somatosensory nose region (SSp-n) in slice 5. Next, we selected the well-known neuropeptide somatostatin *Sst* and its cognate receptor *Sstr2* as the candidate pair. Finding their cross-expression significant, we show their expression on tissue and highlight cross-expressing cells. We also compute their bullseye scores and report them as a ratio of cross- to co-expression across 10 neighbors.

Cross-expression and cell type heterogeneity

 We explore the relationship between cross-expression and cell type heterogeneity using the 774 BARseg coronal data³⁴. First, we use *Gfra1* and *Foxp2* to highlight cross-expressing cells and map different cell types to distinct shapes. Second, we count the number of cross-expressing cell- neighbor pairs for numerous genes. Since each cell has a cell type label, we compute cell type purity as the proportion of pairs with the same label. Third, we use the cell type hierarchy to assess if cell-neighbor pairs with the same H1 label have the same H3 label. We first find cross- expressing gene pairs using the entire dataset. Next, using cell pairs with the 'glutamatergic' H1 label, we compute the number of pairs with the same or different H3 labels. We perform a similar analysis using cells labelled as 'GABAergic' at the H1 level. Finally, we compute the frequencies with which cell type label combinations are associated between neighboring cells and normalize this by the expected frequencies of those cell type pairs in the population.

Discovering combinatorial anatomical marker genes

 We observed that cross-expression discovers anatomical marker genes that delineate the thalamus. To quantitatively assess this, we made a mask by combining the following regions:

 anterior group of the dorsal thalamus (ATN), intralaminar nuclei of the dorsal thalamus (ILM), lateral group of the dorsal thalamus (LAT), medial group of the dorsal thalamus (MED), midline group of the dorsal thalamus (MTN), ventral group of the dorsal thalamus (VENT), and ventral posterior complex of the thalamus (VP). Importantly, we compared every brain region annotation 792 in our data with Allen CCFv3 atlas⁵⁷ and judged the ones presented here to best mark the thalamic regions. This allowed us to calculate the number of cells expressing each gene within or outside the thalamus. For cross-expressing cells, we considered a pair as thalamic if both cells were part of the regional mask. More generally, potential combinatorial markers can be discovered by assessing if their cross-expression is spatially enriched.

 Our second exploration involved well-known genes *Foxp2* and *Cdh13*, which mark cortical layer 6 and show pan-layer expression in the cortex, respectively. These genes exhibited significant cross-expression, which was spatially enriched in layer 6, whose boundaries we identified using H2 cell type annotation. The spatial enrichment was viewed by comparing tissue 801 plots with and without highlighting cross-expressing cells.

Networks of cross-expression

 Using the MERFISH data, we computed cross-expression p-values between all genes and 805 binarized the matrix at $\alpha \le 0.05$ to create an adjacency matrix. We calculate the node degree as 806 the number of edges formed by each gene and create a network with second-order edges (shared connections) as outlined in equation 6.2. We set the threshold for second-order edges to 4, meaning that two genes are connected if they share at least 4 first-order edges, ensuring that the 809 higher-order network is robust. Next, we use the input package⁸⁴ to perform Louvain clustering (with default parameters) on the second-order network and thus assign genes to communities.

811 We visualize the network using Cytoscape⁸⁵ (v3.10.1), mapping node size to degree, color to node community, and edge color to edge type (first-order, second-order, or both). We use the 813 "organic" layout and apply "remove overlaps" from the y_{Files} app⁸⁶ and tweak the network to 814 further reduce overlaps. Finally, we use the Legend Creator app⁸⁶ to render a legend with node degree size and community assignment.

 Because our network revealed *Gpr20* as topologically salient, we performed gene 817 ontology^{87,88} (GO) enrichment analysis on genes that cross-expressed with it ('test set'). Here, we used the entire gene panel (except *Gpr20*) as the background set and used the hypergeometric test to determine if it significantly overlapped with the test set, giving us p-values for each GO functional group. We report FDR-corrected p-values. Additionally, for each gene cross-expressed with *Gpr20*, we used the cells involved in cross-expression, rather than the entire dataset, to compute co-expression with cell type marker genes and compared these global profiles between marker types.

 Since the cells expressing *Gpr20* visually showed spatial autocorrelation, we assessed their neighbors as well as randomly chosen cells for the expression of *Gpr20*. We L1-normalized 826 the counts for both groups, rendering them into probability distributions, and computed cumulative 827 sums. To calculate the area under curve (AUC), we scaled the neighbor order between 0 and 1 828 and used the $trapz$ function from R's $pracma$ package to calculate the AUC.

 Within the main network, we introduce a further constraint that cross-expressing genes must lack significant co-expression. We curate the subnetwork by removing genes with node degree of 1 and assign cell type labels based on genes' co-expression with marker genes. Like before, we perform GO enrichment using the subnetwork genes as the test set and the gene panel as the background set, and report FDR-corrected p-values.

 To assess whether cross-expression networks are more similar between adjacent slices than between distant slices, we compute slice-specific cross-expression networks and calculate Spearman's correlation between these networks. The correlations are plotted against distances between slices, where the "distance" is the difference in the slice order. As a control, we compute the Spearman's correlations between slice-specific networks obtained from different brains and 839 plot this against the "distance" between the slice ID's.

Cross-expression replicability across batches

 To assess the replicability of the cross-expression signature, we used the MERFISH dataset containing 3 biological replicates (mouse brains) with 3 slices each, where the slices are sampled from approximately the same location across the brains. We compared the slice-specific networks between corresponding slices. Moreover, for slice-specific and brain region-specific networks, we performed comparisons within the sagittal data and within the coronal data as well as between these two datasets. Finally, observing that the dorsal to ventral direction is sampled in both the coronal and the sagittal brains, we compared the densities of cross-expressing cells in the dorsal 849 to ventral directions across these datasets.

Cell segmentation quality control assessment

 We assessed the quality of cell segmentation at a global level by comparing co-expression 853 between the scRNA-seq³¹ and MERFISH spatial transcriptomic data. Since the scRNA-seq was obtained from dissected brain regions, we established correspondence between these and the brain region annotations in the MERFISH data. The regions used in both data are reported in Supplementary Table 1. We included only those genes – and in the same order – as present in the MERFISH data. We calculated gene co-expression using Pearson's correlation and compared 858 these across the two datasets.

 To quantify variability between platforms, we compared gene co-expression between 860 scRNA-seq and snRNA-seq⁶⁸ for the same genes – and in the same order – as above. Because the snRNA-seq was obtained from dissected brain regions, we established correspondence between these and the scRNA-seq data. The regions used in these data are reported in Supplementary Table 2. Like before, we quantified co-expression using Pearson's correlation and compared it across the two datasets.

Gene expression noise thresholds and cell-neighbor relations

 Because gene expression measurement is noisy, we applied thresholds of 1 to 10 molecules, thus specifying the minimum number of counts per cell a gene must have to be considered expressed. We then compared cross-expression networks across these thresholds.

 Additionally, a cell might be the nearest neighbor of one or more cells. To ensure that our framework captures this variability, we compare cross-expression networks for the one-to-one and many-to-one mappings with each other and with that of the full dataset.

Benchmarking the algorithm's speed

 We assessed the speed of the cross-expression algorithm by duplicating our BARseq coronal data, where the gene panel ranged from 2,000 to 8,000 and the number of cells ranged from 20,000 to 200,000. We ran the cross-expression algorithm and calculated the time on a 16 GB Apple M1 Pro macOS Sonoma 14.5 laptop.

Data availability

 The MERFISH/ MERSCOPE data was downloaded from Vizgen's mouse brain receptor map at [https://info.vizgen.com/mouse-brain-data.](https://info.vizgen.com/mouse-brain-data) The BARseq coronal data is deposited at the Brain 883 Image Library (BIL) at [https://api.brainimagelibrary.org/web/view?bildid=ace-cry-zip,](https://api.brainimagelibrary.org/web/view?bildid=ace-cry-zip) with the cell 884 and rolony level data at [https://data.mendeley.com/datasets/8bhhk7c5n9/1.](https://data.mendeley.com/datasets/8bhhk7c5n9/1) The BARseq sagittal data's sequencing images are being deposited to BIL. While it is being approved, we stored the cell-level gene expression and cell metadata at

- [https://drive.google.com/drive/folders/1fk5JDeVJcE71iH1AalCT0il9PN9DTJJm?usp=drive_link.](https://drive.google.com/drive/folders/1fk5JDeVJcE71iH1AalCT0il9PN9DTJJm?usp=drive_link)
- 888 scRNA-seq is at https://alleninstitute.github.io/abc_atlas_access/descriptions/WMB_dataset.html
- and snRNA-seq at [https://docs.braincelldata.org/downloads/index.html.](https://docs.braincelldata.org/downloads/index.html)

Code availability

891 The R package is available at https://github.com/ameersarwar/cross_expression

Acknowledgements

 A.S. received funding from University of Toronto's FAST Fellowship and from NSERC's Canada Graduate Scholarship. X.C. was supported by grants R01MH133181 and DP2MH132940. L.F. received support from R01MH133181 and J.G. from both R01MH133181 and R01MH113005. 897 The content is solely the responsibility of the authors and does not necessarily represent the official views of the National Institutes of Health.

Author Contributions

 J.G. conceived the project. A.S. conducted the analyses, developed the software, and wrote the first draft under supervision from J.G. M.R and H.C. collected the sagittal brain data under

supervision from X.C. L.F. managed, curated, and parsed the datasets. All authors interpreted the

- results and edited the manuscript.
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Competing interests

- L.F. owns shares in Quince Therapeutics and has received consulting fees from PeopleBio Co.,
- GC Therapeutics Inc., Cortexyme Inc., and Keystone Bio. The remaining authors declare no competing interests.
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