Charting Tissue Expression Anatomy
by Spatial Transcriptome Deconvolution
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We create data-driven maps of transcriptomic anatomy with a probabilistic framework for unsupervised pattern discovery in spatial gene expression data. Convolved negative binomial regression is used to find patterns which correspond to cell types, microenvironments, or tissue components, and that consist of gene expression profiles and spatial activity maps. Expression profiles quantify how strongly each gene is expressed in a given pattern, and spatial activity maps reflect where in space each pattern is active. Arbitrary covariates and prior hierarchies can be specified to support and leverage complex experimental designs.

Transcriptomic patterns in mouse brain and olfactory bulb correspond to neuroanatomically distinct cell layers. Moreover, batch effects are successfully addressed, leading to consistent pattern inference for multi-sample analyses. On this basis, we identify known and uncharacterized genes that are spatially differentially expressed in the hippocampal field between Ammon’s horn and the dentate gyrus.

The analysis of spatially stratified, transcriptome-wide data [1–3] poses additional challenges compared to classical analysis of bulk RNA sequencing (RNA-Seq) samples. In the classical setting, annotated covariates dictate how samples are to be grouped and compared. These covariates are typically known, and often controlled for, as is the case when performing differential gene expression (DGE) analysis for bulk sequencing count data in DESeq2 [4]. In contrast, the covariates determining gene expression in space are often unknown and can change both gradually or abruptly. For example, the number of infiltrating immune cells per unit area often follows smooth gradients, yet tissue boundaries dramatically impact gene expression over small distances.

If the cell types underlying a sample are well characterized, it becomes analytically possible to determine mixing proportions of the cell types’ known expression profiles in spatial gene expression data. But in the general case, there exists the dual discovery problems of expression profiles and spatial distributions.

Two recently introduced statistical tests, Trendsceek and SpatialDE, quantify the extent of spatial variation of individual genes’ expression [5, 6]. Transcending individual genes, the “automatic expression histology” feature of SpatialDE implements a gene clustering approach to hidden pattern discovery. Such clustering formulations, however, appear challenged by genes that participate in multiple, spatially overlapping expression programs, and grade-of-membership formulations seem more appropriate for hidden pattern discovery.

Joint analysis of multiple data sets substantially increases statistical power and sensitivity but is challenging due to the presence of batch effects. Both non-parametric [7, 8] and model-based approaches are used to address such effects in the analysis of single cell RNA sequencing (scRNA-Seq) data. The model-based methods perform regression for the count data based on known sample-level covariates and allow for the discovery of unknown ones.

Log-normal expression models are used for bulk [9, 10] and single cell [11, 12] RNA-seq data. However, they do not faithfully reflect the discrete count nature of RNA-Seq data. Alternatives are discrete count expression models, such as models based on the Poisson [13] or the negative binomial [14, 15] distributions, with the latter being better suited for modeling over-dispersed gene expression data.

ZINB-WaVE [14] offers a zero-inflated negative binomial (ZINB) regression framework for unknown covariate discovery, including gene-level covariates. Embedded in a hierarchical probabilistic ZINB model, scVI [15] utilizes deep neural networks to model gene-level responses based on both a latent space and known sample-level covariates.

Interpolating properties of bulk RNA-Seq and
scRNA-Seq. Spatial Transcriptomics [3] (ST) count data reflects the gene expression of multiple, but comparatively few cells. Since not all of these cells need to be of the same type, probabilistic models of ST data should admit a mixture interpretation on the level of the counts in each spot. Among the above-mentioned methods, only our previously described Poisson regression based method [13] admits such an interpretation but the rigid mean-variance coupling of the Poisson distribution limits its usefulness.

Results

Here we describe spatial transcriptome deconvolution (STD), a hidden pattern discovery method, which uses convolved hierarchical negative binomial regression to identify transcriptomic patterns in space. We first give an overview of the method and study performance using synthetic data, before we present several applications with real biological data.

Spatial Transcriptome Deconvolution

Figure 1 illustrates an application of our method, STD. It performs inference with a probabilistic model for the counts observed for each gene in each spot. Then it computes expected values of marginal relative frequencies: gene expression profiles and spatial activity maps for a set of transcriptomic factors. Like for other model-based methods, these serve as an effective, lower-dimensional representation of the data, and are used for downstream analysis.

In order to leverage increased statistical power, the method is designed for the joint analysis of multiple Spatial Transcriptomics (ST) [3] data sets. On a high level, the inputs to the method are one or more ST count matrices, a specification of the experimental design, as well as an adaptation of the probabilistic model to the specific application.

Core model

We assume that the observed count $x_{gt}$ for gene $g$ in spot $s$ is the sum of hidden counts $x_{gts}$ due to $T$ transcriptomic factors,

$$x_{gt} = \sum_{t=1}^{T} x_{gts}, \quad (1)$$

and that these in turn are negative binomially distributed,

$$x_{gts} \sim \text{NB} \left( r_{gts}, \rho_{gts} \right), \quad (2)$$

according to eq. (5), with rate and odds parameters $r_{gts}$ and $\rho_{gts}$. Notably, the odds parameters $\rho_{gts}$ are restricted to not depend on the factor (see Online Methods).

The choice of the number of factors is a critical parameter of our approach; it determines whether sufficient factors are available or whether factors are overallocated. This choice is currently left to the user, and it may necessitate some experimenting. In the future, we envisage to employ non-parametric process priors to perform inference across numbers of factors.

Regression and experimental design

When jointly analyzing a set of count matrices resulting from multiple ST experiments, both technical and biological variation between the samples needs to be accounted for. To this end, our method performs regression for the negative binomial distribution's log rate and log odds parameters, with a probabilistic model adapted to the specific application by the analyst. Thus, in addition to the data, the analyst needs to provide the experimental design and to adapt the model to it, by specifying the regression equations as well as the probabilistic prior structure. The framework offers flexible modeling choices for the prior structure: hierarchical probabilistic structures composed from common exponential-family distributions. The graphical probabilistic structure of the model is specified using conventional mathematical notation in model specification files, as described in the Online Methods.

In this communication, we will consider only non-hierarchical, standard normal priors on the regression coefficients.

Visualization

Designed in a grid arrangement, the geometries of the ST microarrays exhibit slight irregularities due to technical variability in the printing process. To both accurately represent individual spots' positions and to address missing data in a visually unobtrusive manner, we utilize Voronoi-tessellation for visualization of inference results (fig. S1).

Maps of transcriptomic anatomy

When dealing with spatial maps for numerous features—whether genes or factor activities—it is frequently helpful to condense the information present in them. This can be done with dimensionality reduction techniques from machine learning, such as t-distributed stochastic neighbor embedding (t-SNE) [16], principal component analysis (PCA), or uniform manifold approximation and projection (UMAP) [17]. By applying such a technique, we compress the information across all features into three components. After rescaling into the unit cube, we use these as coordinates in color space to colorize the spots in spatial plots. Feature similarity is thus encoded by colors, and when the features reflect transcriptome-wide gene expression data we refer to such plots as maps of transcriptomic anatomy (fig. S2).

Synthetic data experiments

To study the sensitivity of the model to properties of the input data, we simulate synthetic data based on different sets of ground truth parameters. For each such set, the simulated data is deconvolved and the result of the deconvolution is compared to the ground truth (fig. 2a).
Figure 1: Overview of Spatial Transcriptome Deconvolution (STD). Spatial Transcriptomics (ST) is performed for multiple sections from multiple individuals, yielding a set of gene maps, one for each gene and each sample, quantifying the number of reads, \( x_{gs} \), observed for gene \( g \) in spot \( s \). STD performs inference to determine point estimates for all parameters, and subsequently computes expectations of marginal relative frequencies. Thus, the gene maps are decomposed by STD into a set of transcriptomic factors, each comprising one gene expression profile and for each sample a spatial activity map. Expression profiles quantify how strongly each gene is expressed in a given factor, and spatial activity maps reflect where in space each factor is active. Only four factors are indicated, and only ten genes per expression profile. Figure S1 illustrates how Voronoi tesselation is here used to visualize spatial gene expression. In addition to the data, the inference of STD depends on the experimental design and the probabilistic model is adapted to the specific application. The model is composed of three parts: 1) the core model specifying that the observed counts \( x_{gs} \) are the sum of the hidden counts \( x_{gts} \) of several transcriptomic factors \( t \), which in turn are negative binomially distributed; 2) the regression equations for the rate and odds parameters of the negative binomial distribution, \( r_{gts} \) and \( \rho_{gs} \); 3) the probabilistic structure specifying the prior distributions of the regression coefficients. The regression equations and prior structure are adapted to the specific application. To illustrate application adaptation, the rate regression equation contains terms \( r_g \), \( r_{g indiv} \), and \( r_{g treat} \) for gene-dependent terms respectively specific to the sample, individual, and treatment. Finally, the spatial activity maps can be summarized across the factors to yield maps of transcriptomic anatomy, either quantitatively for visual inspection or qualitatively—using hierarchical clustering—for down-stream analyses. Figure S2 illustrates how quantitative maps of transcriptomic anatomy are created.
We evaluate the success of the deconvolution by computing the Pearson correlation between the expected number of reads generated by the true and inferred models in the gene-factor and spot-factor marginals. Details about the data generation and performance evaluation are described in the Online Methods.

Overall, the performance of the model is positively associated with and highly sensitive to the number of reads (fig. 2b) and the heterogeneity of the transcriptomic factors (figs. 2d and 2e), both in terms of their spatial activities and gene profiles. In contrast, the performance is less sensitive to the number of spots in the input data (fig. 2c). Crucially, for parameter values inferred from real data (annotated in figs. 2b to 2e), the data is deconvolved with high accuracy.

Our method outperforms the Poisson regression framework of Berglund et al. [13] (figs. 2b and 2c), even when using a Poisson source model similar to their inference model (Online Methods, fig. S3).

**Mouse olfactory bulb**

We jointly analyze sixteen mouse olfactory bulb ST libraries from seven individuals, including four previously-unpublished libraries (table T1 and Online Methods).

The factor analysis was performed for twenty factors utilizing staging, 20% dropout frequency, and adaptive down-sampling to equate sequencing depth (Online Methods). Results for a subset of samples and factors are shown in fig. 3; fig. S6 displays results for all samples and factors.

Sequencing depth is comparable across the samples (fig. 3a), and the relative proportions of reads and spots attributed to the different factors are approximately constant across the samples (figs. S4c and S4d). Factors are ordered by decreasing number of attributed reads, and top-ranking factors have markedly more reads attributed than lower ranking ones (fig. 3b).

Microscopy images of the H&E-stained cryo-sections and inferred spatial factor activity maps are shown in figs. 3c and 3d. The spatial activities of most inferred factors reflect the olfactory bulb anatomy across individuals and replicates, while for four factors the correspondence to anatomy is less clear (factors 6, 8, 12, 20 in figs. 3d and S6b). Summarizing the spatial factor activities by t-SNE or UMAP (figs. 3e and 3f) yields colorizations of the spots consistent with anatomical position across individuals and replicates. But when the same summarization techniques are applied directly to the read counts (figs. S7 to S9), rather than to the spatial factor activities, then uncorrected effects are visible between samples that impede identification of corresponding regions across individuals and replicates.

We partitioned the spots into five sets based on hierarchical clustering of the spatial factor activities (figs. 3g and S6e). From outside inwards, these clusters correspond to the olfactory nerve layer, the glomerular layer,
the plexiform and mitral cell layers, as well as two for the granular cell layer: a peripheral and a central one. The outer plexiform, mitral, and inner plexiform layers cannot clearly be resolved by partitioning into more clusters (results not shown), presumably due to insufficient spatial resolution of the array spots. Applying hierarchical clustering directly to the read counts (fig. S10), rather than to the spatial factor activities, again exhibits uncorrected between-sample effects.

Subsequently, we performed DGE analysis between all pairs of clusters (Online Methods). We intersected the sets of genes that are significantly up-regulated in a given cluster in all pairwise analyses (supplementary dataset D1) to define sets of genes that are specific to that cluster. We then retrieved images of in situ hybridizations (ISH) for the cluster-specific genes from the Allen brain atlas [18]. Inspection of representative ISH images (fig. S11) reveals that the cluster-specific genes share common, cluster-specific spatial expression patterns in the Allen brain atlas.

**Coronal brain sections**

We analyze four ST libraries prepared from mouse coronal brain sections, neighboring sections from two individuals. These sections contain parts of the hippocampus, cortex, cerebral nuclei, thalamus, hypothalamus, as well as several cross-cutting fiber tracts.

Like for the olfactory bulb data, the factor analysis was again performed for twenty factors utilizing staging, 20% dropout frequency, and adaptive down-sampling to equate sequencing depth (Online Methods). Results for all samples and factors are shown in fig. 4, and read and spot statistics across samples and transcriptomic factors are displayed in fig. S5.

Inspection of the spatial factor activity maps reveals that at least 17 of the 20 factors correspond to neuroanatomical brain structures (fig. 4b). In particular, several factors correspond to cortical areas; one each for layer 1, for layers 2 and 3, for layers 4 and 5, and for layer 6, for the striatum-like amygdalar nuclei and for the cortical olfactory areas. Four structures are identified within the hippocampal field: the pyramidal cell layer in Ammon’s horn (one factor for CA1 and CA2, another for CA3), the granule cell layer in the dentate gyrus, and the molecular layer. Thalamic structures are represented by three factors, for the dorsal thalamus, the ventral thalamus, and for the reticular nucleus. Further factors correspond to the hypothalamus, the caudoputamen, the lateral ventricle, the third ventricle, and white matter.

The remaining three factors appear to capture residual signal, as they exhibit more diffuse spatial activities and do not capture the same anatomical structures in both individuals. Factors 11 and 18 capture specific but different anatomical structures within the individuals, and factor 10 competes with the white matter factor in one sample.

**Figure 3:** Transcriptomic patterns in mouse olfactory bulb sections. (a) Number of reads across samples, colors indicate factor. (b) Number of reads across factors, colors indicate sample. (c) H&E-stained microscopy images for 4 of 16 sections. All sections are shown in fig. S6. (d) Spatial factor activity maps for 13 of 20 factors. All factors are shown in fig. S6. (e) t-SNE summarization of factor activities. (f) UMAP summarization of factor activities. (g) Hierarchical clustering of factor activities into five clusters.
Figure 4: Transcriptomic patterns in mouse coronal brain sections. (a) H&E-stained microscopy images. (b) Spatial factor activity maps, and neuroanatomical regions co-incident with factor activity. (c) t-SNE summarization of factor activities. (d) UMAP summarization of factor activities. (e) Hierarchical clustering of factor activities into twelve clusters. Abbreviations: HPF hippocampal field, TH thalamus, C1–C6 cortical layers 1–6, CA1–CA3 Ammon's horn regions 1–3.
Figure 5: Differential gene expression analysis between the clusters corresponding to dentate gyrus and Ammon's horn. (a) Expression log fold change of dentate gyrus over Ammon's horn as a function of mean expression. Top 5 most significantly up- and down-regulated genes are labeled. Circle area is proportional to negative log p-value, p-value < 0.01 marked red. (b–e) In situ hybridization images from the Allen mouse brain atlas of genes significantly differentially expressed between Ammon's horn and dentate gyrus.

Summarizing the spatial activity maps by t-SNE or UMAP consistently colorizes spots in corresponding anatomical positions within and across individuals and replicates (figs. 4c and 4d). Similarly, hierarchical clustering of the spatial activities into twelve clusters consistently partitions the spots into distinct, neuroanatomically-defined regions (figs. 4e and S10d).

We performed pairwise DGE analyses for all pairs of clusters (supplementary dataset D2). Functional enrichment analysis for these DGE results yields almost exclusively meaningful categories from the various ontologies (supplementary dataset D3), such as “calcium ion-binding” (GO-MF), “dendrite” and “axon” (GO-CC), “chemical synapse transmission” (GO-BP), “dopaminergic synapse” and “glutamatergic synapse” (KEGG 2016), “neuronal system” (Reactome 2016), among others.

Considering one such pairwise comparison closer, we find 350 genes differentially expressed (adjusted p-value < 0.01) between the dentate gyrus and Ammon’s horn clusters, of which 233 are expressed higher in dentate gyrus (fig. 5a). “Dentate gyrus” and “Field CA1, stratum oriens” are the categories from the “Allen_brain_up” ontology that are respectively most enriched for genes significantly differentially regulated in this comparison (adjusted p-values $5.7 \times 10^{-21}$ and $3.8 \times 10^{-18}$). In situ hybridizations from the Allen brain atlas constitute further orthogonal validation of the spatial specificity for the most differentially regulated genes from this comparison (figs. 5b to 5e).

Transcriptomic maps mirror anatomy

Juxtaposition of summarized spatial factor activities with the corresponding region in the Allen mouse brain reference atlas (figs. 6a, 6b, 6d, and 6e) reveals that the resulting spot colorization highlights the same gross anatomical features. This is noteworthy because the reference atlas is based on manually-curated expert knowledge while our results are based on automatic analysis of transcriptome-wide data and do not incorporate any prior knowledge.

Further reducing the dimensionality of the spatial factor activities to two dimensions reveals that the olfactory bulb data is modeled by a topologically one-dimensional manifold (fig. 6c), while the brain data exhibit discrete islands corresponding to disjoint clusters of spots (fig. 6f).

Discussion

We presented a probabilistic method to model spatial gene expression count data by convolved negative binomial regression and thereby simultaneously infer unknown gene expression profiles and their unobserved mixing proportions. It supersedes our related, recently introduced deconvolution method based on Poisson regression [13].
The present method is related to and generalizes the negative binomial regression models of DESeq2 and ZINB-WaVE, although it does not currently implement a zero-inflation model. While it is quite possible to augment our model accordingly, it is conceivable that our mixture approach may constitute a modeling alternative for zero-inflation. Further, instead of relying on DESeq2 for DGE analysis, it appears worthwhile to investigate the benefits of directly performing DGE analysis based on the parameters inferred by our model. Another relevant future addition to our method might be to allow for Gaussian processes to be used as prior distributions for the spatial coefficients specified in the regression equations. This could help to discover spatially smoother distributions. Another avenue for future improvement is to improve scalability to large data sets. To date, we have successfully applied our method to datasets approaching $10^8$ spots, and speed-ups could be achieved for even bigger data sets by not considering every spot in each iteration.

The applications to the olfactory bulb and brain sections demonstrated that our method identifies anatomical regions from the spatial, transcriptome-wide data alone, without requiring additional prior knowledge. Furthermore, the identified patterns are consistent across neighboring sections and individuals, and this consistency indicates that our method successfully corrects technical batch effects. Importantly, when dimensionality reduction or hierarchical clustering are applied directly to gene expression data—rather than our spatial factor activities—these un-corrected batch effects are evident and may confound down-stream analyses.

The Allen mouse brain atlas project provided orthogonal validation for the spatial patterns revealed by our analyses, both in terms of functional enrichment results and in terms of in situ hybridization imaging data for individual genes found differential in our hippocampal DGE analysis.

To conclude, the spatial factor activity maps inferred by our method quantitatively reflect the spatial expression profiles of corresponding cell types and tissue anatomy. As such they constitute an explanatory interpretation for patterns observed across thousands of genes and provide a starting point for down-stream analyses. Finally, condensing the activity maps of all factors offers a data-driven way to create qualitative and quantitative maps of transcriptomic anatomy.
Software availability  The method is implemented in C++ and available as free software from https://github.com/SpatialTranscriptomicsResearch/std-nb under the terms of the GNU GPL v3 license. Data and results are available in the same place.

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Contributions  J.M. designed and implemented the method, designed the experiments, and wrote the manuscript. L.B. contributed to the implementation and designed the experiments, and wrote the manuscript. J.La. and J.Lu. supervised the project.

Competing interests  The authors declare no competing interests.

References

1. Ke, R. et al. In situ sequencing for RNA analysis in preserved tissue and cells. Nature Methods 10, 857–860 (2013).
2. Lee, J. H. et al. Highly multiplexed subcellular RNA sequencing in situ. Science 343, 1360–1363 (2014).
3. Ståhl, P. L. et al. Visualization and analysis of gene expression in tissue sections by spatial transcriptomics. Science 353, 78–82 (2016).
4. Love, M. I., Huber, W. & Anders, S. Moderated estimation of fold change and dispersion for RNA-seq data with DESeq2. Genome Biology 15, 550 (2014).
5. Edsgård, D., Johnsson, P. & Sandberg, R. Identification of spatial expression trends in single-cell gene expression data. Nature Methods. doi:10.1038/nmeth.4634 (2018).
6. Svensson, V., Teichmann, S. A. & Stegle, O. SpatialDE: identification of spatially variable genes. Nature Methods. doi:10.1038/nmeth.4636 (2018).
7. Butler, A., Hoffman, P., Smibert, P., Papalexi, E. & Satija, R. Integrating single-cell transcriptomic data across different conditions, technologies, and species. Nature Biotechnology. doi:10.1038/nbt.4096 (2018).
8. Haghverdi, L., Lun, A. T. L., Morgan, M. D. & Marioni, J. C. Batch effects in single-cell RNA-sequencing data are corrected by matching mutual nearest neighbors. Nature Biotechnology. doi:10.1038/nbt.4091 (2018).
9. Gagnon-Bartsch, J. A. & Speed, T. P. Using control genes to correct for unwanted variation in microarray data. Biostatistics (Oxford, England) 13, 539–552 (2012).
10. Risso, D., Ngai, J., Speed, T. P. & Dudoit, S. Normalization of RNA-seq data using factor analysis of control genes or samples. Nature Biotechnology 32, 896–902 (2014).
11. Pierson, E. & Yau, C. ZIFA: Dimensionality reduction for zero-inflated single-cell gene expression analysis. Genome biology 16, 241 (2015).
12. Prabhakaran, S., Azizi, E., Carr, A. & Pe'er, D. Dirichlet Process Mixture Model for Correcting Technical Variation in Single-Cell Gene Expression Data in Proceedings of The 33rd International Conference on Machine Learning (eds Balcan, M. F. & Weinberger, K. Q.) 48 (PMLR, New York, New York, USA, 2016), 1070–1079.
13. Berglund, E. et al. Spatial maps of prostate cancer transcriptomes reveal an unexplored landscape of heterogeneity. Nature communications 9, 2419 (2018).
14. Risso, D., Perraudeau, F., Gribkova, S., Dudoit, S. & Vert, J.-P. ZINB-WaVE: A general and flexible method for signal extraction from single-cell RNA-seq data. bioRxiv. doi:10.1101/292037 (2018).
15. Lopez, R., Regier, J., Cole, M. B., Jordan, M. & Yosef, N. Bayesian Inference for a Generative Model of Transcriptome Profiles from Single-cell RNA Sequencing. bioRxiv. doi:10.1101/292037 (2018).
16. Van der Maaten, L. & Hinton, G. Visualizing High-Dimensional Data Using t-SNE. Journal of Machine Learning Research 9, 2579–2605 (2008).
17. McInnes, L. & Healy, J. UMAP: Uniform Manifold Approximation and Projection for Dimension Reduction. arXiv: 1802.03426v1 [stat.ML] (2018).
18. Lein, E. S. et al. Genome-wide atlas of gene expression in the adult mouse brain. Nature 445, 168–176 (2007).
Online Methods

Materials

Slides with spatially barcoded arrays

To generate the Spatial Transcriptomics data, Codelink Activated Slides (Surmodics) with 1007 distinct capturing oligonucleotides attached were used [1]. Briefly, the oligonucleotides comprising T7 RNA polymerase promoter sequence, 18-mer unique barcode, 9-mer semi-randomized or 7-mer randomized UMI and a poly-20TVN capture region were immobilized in 100 µm spots with center-to-center distance of 200 µm. Six 6200 µm × 6600 µm subarrays were printed onto one glass slide.

Tissue collection and sectioning

Adult C57BL/6 mice were sacrificed and the brains were removed from the cranial cavity. Olfactory bulbs were dissected out on ice, snap-frozen in dry ice/isopentane slurry and then embedded in OCT (Sakura). The left hemisphere was put into a mold filled with cold OCT and snap-frozen in isopentane pre-cooled with liquid nitrogen. Olfactory bulbs and the left hemisphere were sectioned on the cryostat at 10 µm thickness. Sections were placed on the spatially barcoded arrays with 1 sec tion per well.

Fixation, staining and imaging

Sections were fixed for 10 min in 3.6 % to 3.8 % formaldehyde (Sigma) in PBS, washed, then treated for 1 min with isopropanol and air-dried. To stain the tissue, sections were incubated in Mayer’s Hematoxylin (Dako) for 7 min and then Bluing buffer (Dako) for 2 min, followed by Eosin (Sigma) for 20 s. After drying, slides were mounted with 85 % glycerol and imaged using Metafer Slide Scanning Platform (Metasystems). Raw images were stitched together using VSlide software (Metasystems).

Tissue permeabilisation

To separate six subarrays from each other and to create reaction chambers for individual sections, the slide was placed in an ArrayIT hybridization cassette. To pre-permeabilize the tissue, sections of olfactory bulbs were incubated for 30 min at 37 °C with Exonuclease I Reaction Buffer (NEB) mixed with 0.2 µg/µL BSA (NEB). Sections from the hippocampal region were incubated for 20 min at 37 °C with 0.2 U/µL collagenase (Thermo Fisher Scientific) in HBSS buffer (Thermo Fisher Scientific) supplemented with 0.2 µg/µL BSA. After washing in 0.1x SSC buffer (Sigma), sections of olfactory bulbs and the hippocampal region were permeabilized with 0.1 % pepsin/HCl (Sigma) at 37 °C for 10 min and 6 min, respectively. Then, wells were carefully washed with 0.1x SSC buffer.

Reverse transcription and library preparation

Following permeabilisation, reverse transcription mix was added to each well and incubated overnight at 42 °C as described previously [1]. Next, tissue was removed and the surface probes with bound mRNA/cDNA were cleaved from the slide [1]. 65 µL of the reaction mixture containing the released probes were collected from each well and 2nd strand synthesis, cDNA purification, in vitro transcription, aRNA purification, adapter ligation, post-ligation purification, a second 2nd strand synthesis and purification were carried out using an automated MBS 8000 system as described previously [2]. Purified cDNA was then PCR amplified using Illumina Indexing primer [1]. The indexed libraries were purified using carboxylic acid beads on an automated MBS robot system [3] and eluted in 20 µL Elution buffer (Qiagen). The length distribution of the libraries was determined by using the DNA HS Kit (Agilent) with Bioanalyzer 2100 according to the manufacturer’s protocol. The concentration of the libraries was measured with Qubit dsDNA HS (Thermo Fisher Scientific) according to the manufacturer’s protocol. The finished libraries were diluted to 4 nM with Elution buffer and sequenced on the Illumina Nextseq platform using paired-end sequencing, according to the manufacturer’s protocol.

Staining of the slide spots and image alignment

After cleaving of the probes from the glass surface, slide was incubated with hybridisation mixture containing Cyanine-3 labelled probes, as described previously [1]. Fluorescent images were acquired using the same scanning platform as for the bright field images. Bright field images and corresponding fluorescent images were aligned manually using Adobe Photoshop CS6 and the spots located under the tissue were selected.

Bioinformatics

Sequenced reads were processed with the ST Pipeline [4, version 1.4.5] in order to obtain matrices of counts where each cell represents the number of unique molecules for a given spot and a given gene. Homopolymer stretches of at least 10 bp and low quality bases (phred-33 score ≤ 20) were removed from R2. Reads were discarded if R2 was shorter than 20 bp. A contaminant filter was applied to the remaining reads using the Ensembl GRCm38 (v.86) non-coding RNA reference. Filtered reads were then mapped to the genome Ensembl GRCm38 (v.86), demultiplexed and annotated using the reference Mouse GenCode vM11 (Comprehensive gene annotation). Unique counts (UMIs) for
each spot/gene combination were computed with default settings of the ST Pipeline. The obtained matrices of counts were processed to replace Ensembl ids by gene names where only protein-coding, long intergenic non-coding and antisense genes were kept. Finally, the matrices of counts were filtered to keep only the spots under the tissue of the corresponding image datasets.

Regression formula notation

Regression formulæ use the following special arithmetic rules. The 1 denotes an intercept term. $a:b$ denotes an interaction term for covariates $a$ and $b$, and $a*b$ expands to $a + a*b + b$. Addition and interaction are idempotent, $a + a = a$, and $a:a = a$.

Spatial Transcriptome Deconvolution

Negative binomial deconvolution

Core model We assume that the observed count $x_{gt}$ for gene $g$ in spot $s$ is the sum of hidden counts $x_{gts}$ due to $T$ transcriptomic factors,

$$x_{gts} = \sum_{t=1}^{T} x_{gts}$$

and that these in turn are negative binomially distributed,

$$x_{gts} \sim \text{NB} \left( r_{gts}, \rho_{gts} \right),$$

with rate and odds parameters $r_{gts}$ and $\rho_{gts}$ according to eq. (5),

$$\Pr \left( x_{gts} = k \mid r, \rho \right) = \frac{\Gamma \left( k + r \right)}{\Gamma \left( k + 1 \right) \Gamma \left( r \right)} \frac{\rho^{k}}{\left( \rho + 1 \right)^{k+r}}.$$

Notably, the odds parameters $\rho_{gts}$ are restricted to not depend on the factor because from this follows a simplified likelihood, $x_{gts} \sim \text{NB} \left( \sum_{t=1}^{T} r_{gts}, \rho_{gts} \right)$; this allows us to marginalize over $x_{gts}$ during inference. Furthermore, in practice, the odds parameters of our models are typically chosen to depend only on the gene, $\rho_{gts} = \rho_{g}$. This restriction is also present in other models, such as DESeq2, ZINB-WaVE, and scVI for which the dispersion parameters only depend on the gene [5, equation 1] [6, eq. 6][7, eqs. 4 and 5].

Rate and odds regression In the framework, the logarithms of the rate and odds parameters are specified in terms of regression formulæ. The default regression formulæ are

$$\log r \sim 1 + \text{gene} * \text{factor} + \text{factor} * \text{spot}$$

$$\log \rho \sim 1 + \text{gene},$$

see the preceding section for an explanation of the formula notation. These regression formulæ correspond to the following regression equations:

$$\log r_{gts} = r + r_{g} + r_{s} + r_{ts} + r_{s}$$

$$\log \rho_{gts} = \rho + \rho_{g},$$

where the indices $g, t, s$ denote covariate dependence on gene, factor, and spot, respectively.

Covariates Aside from the above-mentioned covariates (intercept, genes, spots, and factors), it may often be necessary to include additional terms in the rate and odds regression formulæ. For example, when data of multiple sections are analyzed then section + gene:section terms can capture technical noise. Furthermore, by writing design files, the samples may be annotated with additional covariates according to the experimental design, and terms depending on these covariates may be used in the formulæ.

In this way, when sections from multiple different biological conditions are analyzed, for example different cancer types, then biological variation can be captured by gene:cancer terms. Thus, when data are available that control for different cancer types and that comprise multiple sections as repeat experiments, then the following rate regression formulæ may be appropriate:

$$\log r \sim 1 + \text{gene} * \left( \text{factor} + \text{cancer} + \text{section} \right) + \text{factor} * \text{spot}$$

Probability distributions and hierarchies In order to adapt the model to the experimental design of the specific application, the framework offers flexible modeling choices for the coefficient prior structure. Available prior distributions include the normal, beta, and gamma distributions; and arbitrary directed graphical probabilistic hierarchies can be built out of these.

Model specification The graphical probabilistic structure of the model is specified using conventional mathematical notation in model specification files. The regression for the rate and odds parameters can be specified either directly in terms of equations or in terms of formulæ that get translated into equations. The following is an example model specification file:

```plaintext
# Rate equation
rate = rate()+rate(gene)+rate(spod)+rate(gene, spot)+rate(type)

# Odds equation
odds = odds()+odds(gene)

# Coefficient distributions
rate() - Normal(0,1)
rate(gene) - Normal(0,1)
rate(gene, type) - Normal(0,1)
rate(spod) - Normal(0,1)
rate(spot, type) - Normal(0,1)
```

draft-534-g100f761
rate(type)  - Normal(0,1)
odds()  - Normal(0,1)
odds(gene)  - Normal(0,1)

Note that type corresponds to the transcriptomic factor.

The regression equations may be arbitrary mathematical expressions composed of sums, differences, products, divisions, exponentiations, and logarithms.

Instead of the regression equations given above, the following formulas could be used equivalently:

# Rate formula
rate := 1+type*(gene+spot)

# Odds formula
odds := 1+gene

Syntactically, expressions with an equality sign, =, such as in the first example, are parsed as equations, while expression with := are parsed as formulas. Finally, expressions with a tilde symbol, ~, are parsed as probability distribution specifications. If no probability distribution is specified for a coefficient, then it is assumed to be distributed according to the standard normal distribution.

Engineering model flexibility  Technically, our method’s modeling flexibility is enabled by representing the abstract syntax tree (AST) corresponding to the user-specified mathematical expressions in the regression equations and computing symbolic derivatives for gradient-based optimization. Function and derivative evaluation relies on just-in-time code generation and compilation for the expressions from the AST utilizing the LLVM compiler framework [8].

Optimization  The framework offers several parameter optimization schemes based on likelihood gradients, including RPROP [9], AdaGrad [10], and ADAM [11] optionally with a Nesterov-type momentum term [12, 13].

Stochastic gradient  Stochasticity is injected into the learning process by two means: we compute a stochastic approximation to the gradient by randomly ignoring counts \( x_{gs} \) during gradient calculation. Furthermore, it is often beneficial to dynamically down-sample the counts so that all experiments have the same read-per-spatrope ratio. The stochastic gradient approximation linearly speeds up the computation but more importantly helps avoid over-training. Down-sampling of counts to equate the read-per-spot ratios avoids uneven likelihood contributions across the experiments due to sequencing depth, which otherwise frequently results in factors focused more towards explaining samples with higher sequencing depth.

Staging  Optimization is done in multiple rounds, in which increasing numbers of parameters are included into the optimization. In the first stage only global, scalar coefficients are optimized. In the second stage, we additionally optimize scalar coefficients that depend on further covariates such as section or individual. From the third stage optimization includes gene-dependent and spot-dependent coefficients that do not depend on further covariates. Stage four also includes gene-dependent and spot-dependent coefficients that depend on further covariates. The fifth stage finally optimizes all coefficients, including gene- and factor-dependent ones, as well as spot- and factor-dependent ones.

Stages one to four perform fifty iterations of gradient updates, the fifth stage performs 2000 iterations.

Visual summarization and clustering  Spatial patterns across many features—regardless whether genes or factors—can be visually summarized with dimensionality reduction techniques from machine learning, see fig. S2. In this context, we consider matrices that have rows for every spot and columns for every feature (which could be genes or spatial factor activities). Utilizing t-distributed stochastic neighbor embedding (t-SNE) [14] or similar methods, such as principal component analysis (PCA) or uniform manifold approximation and projection (UMAP) [15], the number of columns of the matrices is reduced to three. The data are then rescaled into the unit cube and the rows used as coordinates in color space to colorize the spots in a spatial plot. When spots are colored in this way, similar colors indicate similar gene expression or similar factor activities, indicating similar cell type composition. We refer to such plots as maps of transcriptomic anatomy if the features reflect transcriptome-wide gene expression data.

Hierarchical clustering  Aside from visualizing cell type composition by dimensionality reduction of spatial activity maps, it is also possible to apply hierarchical clustering to the spatial activity maps. Unlike the quantitative visualization that dimensionality reduction based approaches yield, clustering partitions the spots into discrete sets, which can be useful for down-stream analyses.

Differential gene expression analysis  Differential gene expression (DGE) analysis is performed as follows. The analyst decides on a suitable number of clusters by inspecting the hierarchical clustering results. For the chosen number of clusters, the spots are then partitioned into sets for each cluster. For all pairs of clusters, a pair-wise DGE analysis is performed using DESeq2.

The performance of the scaling factor determination of DESeq2 deteriorates with increasing number of spots
because it can only utilize genes that have non-zero counts throughout all spots, and the probability of observing no zero count for a given gene in each of \( n \) samples decreases exponentially in \( n \). Therefore, instead of using all spots of a given cluster (which can number in the hundreds or thousands), we randomly select subsets of about 50–100 spots. This is justified based on the assumption that the spots within a cluster are interchangeable.

**Synthetic data experiments**

**Model**

The synthetic data is generated according to the following model:

\[
x_{gts} \sim \text{NB} \left( \alpha r_g r_s, \rho_g \right)
\]

(11)

For notational brevity, we will use \( \varphi_g \) and \( \theta_s \) in place of \( r_g \) and \( r_s \) in this section and denote them in matrix form as \( \Phi \) and \( \Theta \), respectively.

**Measuring performance**

We evaluate the success of the deconvolution by computing the Pearson correlation between the expected number of reads generated by the true and inferred models in the gene-factor and spot-factor marginals. That is, we use \( r_{gs} \) in this section as a performance measure, where either

\[
X = \sum_s x_{gts} \quad \text{and} \quad \hat{X} = \sum_s \hat{x}_{gts}, \quad \text{or} \quad (12)
\]

\[
X = \sum_g x_{gts} \quad \text{and} \quad \hat{X} = \sum_g \hat{x}_{gts}, \quad (13)
\]

and where \( x_{gts} \) and \( \hat{x}_{gts} \) are the number of reads generated by the true and inferred models, respectively, for a given gene \( g \), transcriptomic factor \( t \), and spot \( s \).

**Ground truth parameters**

In each experiment, we generate data for \( |T| = 10 \) factors and let

\[
\log_2 \rho_i = \frac{i}{|G| - 1} \times 8 - 4,
\]

(14)

where \( i \in [0, 1, \ldots, |G| - 1] \) and \( |G| = 1000 \).

We examine model performance along four dimensions: the average number of reads per spot, the number of spots, the heterogeneity of gene profiles, and the heterogeneity of activity maps. In order to measure the latter two quantities, we consider the random vector \( G, T, S \), defined so that \( P(G=g, T=t, S=s) \) is equal to the relative frequency of reads for a given gene \( g \), factor \( t \), and spot \( s \):

\[
p(g,t,s) = \frac{\mathbb{E}[x_{gts}]}{\mathbb{E}\left[\sum_{g,t,s} x_{gts}\right]} = \frac{q_{gt} \varphi_{gt} \rho_g}{\sum_{g,t,s} \varphi_{gt} \theta_s \rho_g},
\]

(15)

We define the heterogeneity of the gene profiles and spatial activity maps, respectively, as

\[
I(G; T) = \sum_{g,t} p(g,t) \log_2 \left( \frac{p(g,t)}{p(g)p(t)} \right), \quad \text{and} \quad (16)
\]

\[
I(S; T) = \sum_{s,t} p(s,t) \log_2 \left( \frac{p(s,t)}{p(s)p(t)} \right). \quad (17)
\]

**Sampling**

We use the following procedure to sample the parameters \( \alpha, \Phi, \Theta \) for an experiment with an average of \( n \) expected reads per spot, \( |S| \) spots, and factor heterogeneities \( i(G; T) \) and \( i(S; T) \):

1. Initialize \( \Phi \) to a \( |G| \times |T| \) matrix and \( \Theta \) to an \( |S| \times |T| \) matrix with each entry set to \( 1/|T| \).

2. Sample \( \Phi \) and \( \Theta \) using the Metropolis-Hastings algorithm on the density \( p(\Phi, \Theta) \propto e^{-E} \), where

\[
E = \delta_{\text{gene}}(\Phi,\Theta)^2 + \delta_{\text{spot}}(\Phi,\Theta)^2,
\]

(18)

and

\[
\delta_{\text{gene}}(\Phi,\Theta) = i(G; T) - I(G; T) \quad \text{and} \quad (19)
\]

\[
\delta_{\text{spot}}(\Phi,\Theta) = i(S; T) - I(S; T). \quad (20)
\]

In order to improve convergence speed, we tune the proposals based on the values of (19) and (20). Formally, we use a time dependent proposal function \( N_t(\Phi, \Theta, S_t) \), where

\[
S_t = \left( k_{t,\text{gene}}, k_{t,\text{spot}} \right) \quad (21)
\]

denotes its state at time \( t \). We let its initial state be \( S_0 = (0, 0) \). \( N_t \) generates new proposals and modifies its state according to the following procedure:

(a) Choose one of

\[
(\Phi, k_{i,\text{gene}}, s_i \mapsto \delta_{\text{gene}}(x, \Theta)) \quad (22)
\]

and

\[
(\Theta, k_{i,\text{spot}}, x \mapsto \delta_{\text{spot}}(\Phi, x)) \quad (23)
\]

at random with equal probability. Call the chosen triplet \( (M, k, f) \).

(b) Select a random set of \( m \) rows in \( M \), where \( m = \lfloor 0.01 \times \text{rows}(M) \rfloor \). Each row is drawn equiprobably from the set of all rows in \( M \) without replacement. Define \( \hat{M} \) to be \( M \) with the selected rows replaced with \( m \) samples from a \( \text{Dir}(\varepsilon^2 u) \) distribution, where \( u \) is a \( |T| \)-dimensional vector of ones.

(c) Let \( \hat{k} = k + 0.001 \times \Delta \), where

\[
\Delta = \begin{cases} 1 & \text{if } \hat{M} > 0 \\ -1 & \text{otherwise.} \end{cases} \quad (24)
\]
The new proposal is
\[ \hat{M}/M \left( \Phi, \Theta \right), \]  
(25)

where the operation \([x/y]z\) denotes replacing variable \(y\) with \(x\) in expression \(z\).

The new state of the proposal function is
\[ S_{t+1} = \left[ k/k \right] S_t. \]  
(26)

3. Given \(\Phi\) and \(\Theta\), we can solve for the scaling parameter \(\alpha\):
\[ \alpha = \frac{n[S]}{\sum_{g,t,s} \varphi_g \theta_{gt} \rho_s}. \]  
(27)

**Inference**

Using the data generated by the model in the preceding section, we infer the parameters of the model
\[ \log r \sim 1 + \text{factor} \times (\text{gene} + \text{spot}) \]  
(28)
\[ \log \rho \sim 1 + \text{gene}. \]  
(29)

We run std-nxt with the following command:

\begin{verbatim}
std-nxt -t10 -i2000 -m model.txt \ --optim=adam --adam_nesterov counts.tsv
\end{verbatim}

**Comparison to Poisson regression** We run the method of Berglund et al. [16] with the following command:

\begin{verbatim}
std -t10 -i2000 counts.tsv --sample=contributions,phi,phi_prior,phi_local,theta,theta_prior,spot,baseline
\end{verbatim}

**Poisson model**

To better understand how our method compares to the one of Berglund et al. [16], we substitute eq. (11) to conform to their inference model:
\[ x_{gsi} \sim \text{Pois} \left( \alpha r_g r_s \right). \]  
(30)

Sampling of \(\alpha\), \(r_g\), and \(r_s\) and inference are performed analogously to what has been described above. The results of the performance evaluation are shown in fig. S3.

**Design and analysis of biological data**

The experimental design for the olfactory bulb and the brain samples is given in table T1 and table T2. For the olfactory bulb samples, we discard the sex covariate since it is confounded with individual B. We thus use the following rate regression formula for both the olfactory bulb and brain analyses:
\[ \log r \sim 1 + \text{gene} \times (\text{factor} + \text{individual} + \text{section}) + \text{factor} \times \text{spot}. \]  
(31)

Analyses were performed with version 0.3-111-g40dcf70 of std-nxt and the following command:

\begin{verbatim}
std-nxt --adjdepth --stage 50 \ --minread_spot 100 --dropout 0.2 \ -v -i 2000 -r 10 -t 20 --optim adam \ --design design.txt --model model.txt
\end{verbatim}

**References**

1. Ståhl, P. L. et al. Visualization and analysis of gene expression in tissue sections by spatial transcriptomics. Science 353, 78–82 (2016).
2. Jemt, A. et al. An automated approach to prepare tissue-derived spatially barcoded RNA-sequencing libraries. Scientific Reports 6, 37137 (2016).
3. Lundin, S., Stranneheim, H., Pettersson, E., Klevebring, D. & Lundeberg, J. Increased throughput by parallelization of library preparation for massive sequencing. *PloS one* 5, e10029 (2010).

4. Fernández Navarro, J., Sjöstrand, J., Salmén, F., Lundeberg, J. & Ståhl, P. L. ST Pipeline: an automated pipeline for spatial mapping of unique transcripts. *Bioinformatics (Oxford, England)* 33, 2591–2593 (2017).

5. Love, M. I., Huber, W. & Anders, S. Moderated estimation of fold change and dispersion for RNA-seq data with DESeq2. *Genome Biology* 15, 550 (2014).

6. Risso, D., Perraudeau, F., Gribkova, S., Dudoit, S. & Vert, J.-P. ZINB-WaVE: A general and flexible method for signal extraction from single-cell RNA-seq data. *bioRxiv*. doi:10.1101/125112 (2017).

7. Lopez, R., Regier, J., Cole, M. B., Jordan, M. & Yosef, N. Bayesian Inference for a Generative Model of Transcriptome Profiles from Single-cell RNA Sequencing. *bioRxiv*. doi:10.1101/292037 (2018).

8. Lattner, C. & Adve, V. *LLVM: A Compilation Framework for Lifelong Program Analysis & Transformation* in Proceedings of the International Symposium on Code Generation and Optimization: Feedback-directed and Runtime Optimization (IEEE Computer Society, Palo Alto, California, 2004), 75–.

9. Riedmiller, M. & Braun, H. Rprop - A Fast Adaptive Learning Algorithm in Proceedings of the International Symposium on Computer and Information Science VII (1992).

10. Duchi, J., Hazan, E. & Singer, Y. Adaptive subgradient methods for online learning and stochastic optimization. *Journal of Machine Learning Research* 12, 2121–2159 (2011).

11. Kingma, D. & Ba, J. Adam: A method for stochastic optimization. arXiv:1412.6980 [cs.LG] (2014).

12. Nesterov, Y. A method for unconstrained convex minimization problem with the rate of convergence $O(1/k^2)$. *Doklady AN USSR* 269, 543–547 (1983).

13. Dozat, T. Incorporating Nesterov momentum into ADAM (2016).

14. Van der Maaten, L. & Hinton, G. Visualizing High-Dimensional Data Using t-SNE. *Journal of Machine Learning Research* 9, 2579–2605 (2008).

15. McInnes, L. & Healy, J. UMAP: Uniform Manifold Approximation and Projection for Dimension Reduction. arXiv:1802.03426v1 [stat.ML] (2018).

16. Berglund, E. et al. Spatial maps of prostate cancer transcriptomes reveal an unexplored landscape of heterogeneity. *Nature communications* 9, 2419 (2018).
Figure S1: Three ways of visualizing spatial expression patterns. The geometries of the ST microarrays exhibit slight irregularities due to technical variability in the printing process: individual spots may be misprinted, and lack surface probes to capture mRNA; other spots may fuse with neighboring spots due to electrostatic attraction of the printing ink, and may have to be discarded. Points and vertices represent measurement points, with their plot positions corresponding to the spatial coordinates. Single channel information such as the expression of individual genes can be visually encoded in terms of size or color from a gradient palette. Alternatively, the Voronoi tessellation can be colored to convey the desired information.

Figure S2: Visualizing complex spatial information by dimensionality reduction. The input matrices are $n \times k$ matrices corresponding to $n$ spots and $k$ features. The $k$ input features may represent genes or factors, and the matrices typically contain sequencing read counts or factor activities. These $k$ features may be visually represented with $k$ single-channel images. Alternatively, the features may be combined and jointly visualized, making use of multiple color channels. By applying dimensionality reduction methods such as PCA, t-SNE, or UMAP, the $n \times k$ matrices are reduced to $n \times 3$ matrices. The matrices are then rescaled to fit into the unit cube, and the three components are used as coordinates in color space to colorize spots in spatial plots.

Figure S3: Performance evaluation on synthetic data using a Poisson source model. Plots show the correlation between the expected number of reads in the true and inferred models over the spot-factor and gene-factor marginals as a function of the number of reads per spot (a) and the number of spots (b) in the input data. Lines correspond to medians and shaded areas to minima and maxima. Solid lines show results for negative binomial deconvolution, as described in this paper, while dashed lines show results for the Poisson regression framework of Berglund et al. [1].
Figure S4: Reads and spots explained by factors across the mouse olfactory bulb samples. Colors denote factors (a–d) or samples (e–h).

Figure S5: Reads and spots explained by factors across the mouse brain samples. Colors denote factors (a–d) or samples (e–h).
Figure S6: Transcriptomic patterns in mouse olfactory bulb sections. (a) H&E-stained microscopy images. (b) Spatial factor activity maps. (c) t-SNE summarization of factor activities. (d) UMAP summarization of factor activities. (e) Hierarchical clustering of factor activities into five clusters.
Figure S7: Summarization by principal component analysis (PCA) of gene expression (a, c) and factor activities (b, d) in mouse olfactory bulb (a, b) and mouse brain (c, d).
Figure S8: Summarization by t-distributed stochastic neighbor embedding (t-SNE) of gene expression (a, c) and factor activities (b, d) in mouse olfactory bulb (a, b) and mouse brain (c, d).
Figure S9: Summarization by uniform manifold approximation and projection (UMAP) of gene expression (a, c) and factor activities (b, d) in mouse olfactory bulb (a, b) and mouse brain (c, d).
Figure S10: Hierarchical clustering of gene expression (a, c) and factor activities (b, d). Five clusters shown for mouse olfactory bulb (a, b) and twelve clusters for mouse brain (c, d).
Figure S11: Spatial cluster specific genes in the Allen brain atlas [2]. Images from the Allen brain atlas for genes that are expressed significantly higher in one cluster versus all others. Based on clustering results shown in figs. S6e and S10b. From outside inwards, the clusters correspond to the olfactory nerve layer (layer 3), the glomerular layer (layer 5), the plexiform and mitral cell layers (layer 4), as well as two for the granular cell layer: a peripheral (layer 2) and a central one (layer 1).