Discovery and generalization of tissue structures from spatial omics data

Graphical abstract

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In brief

Wu et al. proposed a universal, efficient annotation tool, SCGP, for spatial omics data, which allows unsupervised recognition of multi-cellular tissue structures. They further proposed the SCGP-Extension pipeline to generalize these annotations to unseen samples, achieving robust, consistent tissue structure identification.

Highlights

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- SCGP is a highly flexible and efficient tool for spatial omics annotation
- SCGP shows outstanding performance across 8 distinct spatial omics datasets
- **.** SCGP-Extension generalizes existing tissue structures to unseen samples
- SCGP-Extension effectively addresses common data integration challenges

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Discovery and generalization of tissue structures from spatial omics data

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generalizability of annotations from a reference dataset to new or unseen data also remains a major challenge for methods in this arena. The current work introduces spatial cellular graph partitioning (SCGP) and its reference-query extension pipeline, SCGP-Extension, as unsupervised annotation tools that streamline and simplify this process, enhancing the consistency, reliability, and generalization of structure annotations across large datasets.

SUMMARY

Tissues are organized into anatomical and functional units at different scales. New technologies for highdimensional molecular profiling in situ have enabled the characterization of structure-function relationships in increasing molecular detail. However, it remains a challenge to consistently identify key functional units across experiments, tissues, and disease contexts, a task that demands extensive manual annotation. Here, we present spatial cellular graph partitioning (SCGP), a flexible method for the unsupervised annotation of tissue structures. We further present a reference-query extension pipeline, SCGP-Extension, that generalizes reference tissue structure labels to previously unseen samples, performing data integration and tissue structure discovery. Our experiments demonstrate reliable, robust partitioning of spatial data in a wide variety of contexts and best-in-class accuracy in identifying expertly annotated structures. Downstream analysis on SCGP-identified tissue structures reveals disease-relevant insights regarding diabetic kidney disease, skin disorder, and neoplastic diseases, underscoring its potential to drive biological insight and discovery from spatial datasets.

INTRODUCTION

All human organs exhibit characteristic cellular structures that are required for homeostasis and function. These structures are diverse in form, scale, and function and are typically composed of multiple cell types organized into spatial patterns.

Disruptions to these structures usually indicate a disease pro-cess.^{[1](#page-12-0)[,2](#page-12-1)} Recent advances in *in situ* molecular profiling techniques, including spatial transcriptomics $3\overline{2}$ and proteomics, $8\overline{1}$ have allowed us to observe molecular phenotypes and cell states in their native contexts, where interactions between en-tities at different spatial scales—ranging from cells^{[12](#page-13-1)} and cellular

neighborhoods (CN) $13,14$ $13,14$ to tissue organization 15 and patientlevel characteristics $16,17$ $16,17$ - can be explored. Discovering, annotating, and analyzing these functional units in consistent ways is a major goal for the field of spatial biology.

However, in practice, annotating tissue structures in a way that both incorporates complex molecular information and aligns with our histological understanding has been challenging without manual supervision. Consistency across diverse samples, experiments, or disease conditions is difficult to achieve. More-over, existing analysis pipelines^{[18,](#page-13-7)[19](#page-13-8)} are predominantly cell centric $20-22$ or geared toward small sample numbers. Together, these issues severely limit efforts to analyze and interpret atlasscale spatial biology datasets.

In recent studies, computational methods integrating molecular profiling with spatial information have been proposed. Some of these methods aim to improve the analysis of cell-level characteristics, such as better cell-type prediction 23,24 23,24 23,24 23,24 and intercellular communication modeling.^{[12](#page-13-1)[,25](#page-13-12)} Another line of research focuses on annotating larger structures or spatial domains, exploring their interactions and disease relevance. Such annotations are performed based on clustering of cell-type composition^{13[,26](#page-13-13)} or locally smoothed cell features,^{15[,27](#page-13-14)} topic modeling,²⁸ Bayesian modeling,²⁹ optimal transport,³⁰ graph Fourier transform,^{[31](#page-13-18)} and graph neural networks.^{[16](#page-13-5)[,32,](#page-13-19)[33](#page-13-20)} Many of these methods are unsupervised and lack the ability to generalize—that is, the resulting annotations or model cannot be extended to previously unseen samples. When new data are introduced, model retraining or refitting is necessary to annotate unseen data. Consequently, downstream analysis on structures of interest are restricted to only the training/fitting data, as consistent annotations on out-of-sample data cannot be reliably acquired. Similarly, recognizing tissue structures from unseen hematoxylin and eosin (H&E)-stained pathology images is a widely studied task often resolved in a supervised manner.^{34–37} However, these tools tend to be more tissuetype specific, require substantial annotated training datasets, and do not incorporate molecular information.

In this work, we present a universal, robust, and generalizable tissue structure segmentation tool called spatial cellular graph partitioning (SCGP). SCGP is a fast and flexible method designed to identify the anatomical and functional units in any spatial transcriptomic or proteomic dataset. We further introduce a reference-query extension pipeline, SCGP-Extension, which allows a small set of reference annotations to be generalized to previously unseen query samples. This powerful data alignment and integration method can address challenges ranging from experimental artifacts and batch effects to disease condition differences and more, greatly enhancing SCGP's robustness and versatility. To the best of our knowledge, SCGP is the first data-type-agnostic semantic segmentation tool that is purpose built for generalization. We demonstrate its applications to eight datasets collected on different tissue types using diverse profiling techniques including CO-detection by indexing (CODEX), Visium, imaging mass cytometry (IMC), and multiplexed error-robust fluorescence *in situ* hybridization (MERFISH), totaling more than 2.5 million cells. The tissue structures identified by SCGP are evaluated against expert annotations, benchmarked extensively against related software tools, and applied in downstream analysis that reveals disease-relevant biological insights.

RESULTS

Unsupervised partitioning of spatial cellular graphs

To address the task of tissue structure identification, we developed SCGP to perform community detection on specialized graph representations of tissue samples. Nodes in the graphs are small spatial units characterized by spatial coordinates and gene or protein expression at the location [\(STAR Methods](#page-16-0)). In the representative case of multiplexed immunofluorescence (mIF) images, 38 nodes are defined on cells identified through the segmentation pipeline^{[20](#page-13-9)} ([Figure 1A](#page-3-0)). However, this concept of nodes can be further extended to accommodate broader spatial transcriptomics and proteomics data, such as spots in spatial transcriptomics sequencing measurements^{[7](#page-12-3)} or small square patches in single-molecule fluorescence images.^{[3](#page-12-2)} In this study, we primarily present and discuss cell- and spotbased SCGP analysis. An alternative patch-based SCGP experiment can be found in [Figure S10](#page-12-4).

Two types of edges are constructed between nodes [\(Fig](#page-3-0)[ure 1](#page-3-0)B; [STAR Methods\)](#page-16-0). Spatial edges are constructed between nodes based on Delaunay triangulation of node coordinates. These edges aim to capture the adjacency relationships between cells. Feature edges are constructed between nodes that share similar expression profiles.

The Leiden graph community detection algorithm 39 is then applied to the graphs, yielding partitions that represent the different tissue structures ([Figure 1C](#page-3-0)). The central aspect of this method is the joint contribution of two types of edges. Spatial edges guarantee the spatial continuity of the identified tissue structures, differentiating the method from cell-type clustering in that multi-cell tissue structures will appear as cohesive entities. Feature edges interrelate tissue structures of the same type even if they are spatially separated (e.g., two glomeruli from different kidney samples), ensuring the consistency of tissue structure interpretation across samples. This design resonates with a customized tool developed for identifying spatial modules in spatial enhanced resolution omics sequencing (stereo-seq), 27 though SCGP used much sparser feature edges to avoid fragmentation of tissue structures (1–4 nearest neighbors; [STAR Methods;](#page-16-0) [Figures S3E](#page-12-4) and S3F). Ablation experiments [\(Figures S3A](#page-12-4) and S3B) show that both types of edges are necessary for the operation of SCGP. We first verified the effectiveness of SCGP using various types of simulated data representing diverse tissue structures ([Figure S1](#page-12-4)). Results showed consistent and accurate recognition of all ground-truth structures by SCGP across all conditions.

SCGP identifies structures in kidney tissues

To examine the ability of SCGP to recognize known tissue structures, we assessed its performance on a cohort of 17 tissue sections from 12 individuals with diabetes and various stages of dia-betic kidney disease (DKD).^{[40](#page-14-1)} Tissue samples were imaged using the mIF platform CODEX^{[9](#page-13-23)} and further annotated for four major kidney compartments: glomeruli, blood vessels, distal tubules, and proximal tubules. This cohort will be referred to as the DKD Kidney dataset [\(STAR Methods\)](#page-16-0) in the subsequent text.

Together with SCGP, we applied a diverse set of unsupervised annotation tools^{[13](#page-13-2)[,15](#page-13-4)[,24](#page-13-11),[28,](#page-13-15)[32](#page-13-19)} to the DKD Kidney dataset. All

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Figure 1. Workflow for SCGP and SCGP-Extension

(A) Raw mIF images of example kidney samples show multiple tissue structures.

(B) Graph representations of mIF images are constructed on nodes (white circles) representing cells or other spatial units. Spatial edges (solid lines) and feature edges (dashed arrows) are constructed to reflect spatial closeness and feature similarity.

(C) Leiden graph community detection identifies partitions representing tissue structures.

(D) The query sample shares similar structures to the reference partitions.

- (E) Graph representation of the query sample is constructed with additional pseudo-nodes (colored circles in the white box) extracted from the reference partitions. Reference-query edges (brown dashed arrows) are constructed between query nodes and pseudo-nodes.
- (F) Leiden graph community detection yields both existing partitions that align with reference and novel partitions that were previously unseen.

methods were applied to the combination of all 17 samples containing 137,654 cells, i.e., in a joint partitioning manner. Clustering/partitioning outputs on representative samples are visual-ized in [Figures 2A](#page-4-0) and [S2A](#page-12-4), with the leftmost images illustrating the raw mIF images with key biomarkers. Due to the unsupervised nature of the output, we reordered the output clusters of each method in accordance with manually annotated compartments. In [Figure 2](#page-4-0)A, the top image for each column shows the output, and the bottom image highlights the mismatches. Across samples, partitions annotated by SCGP frequently demonstrated the highest fidelity to manual annotations, and they were robust to cell segmentation noise [\(Figures S3C](#page-12-4) and S3D).

To evaluate the performance of each method quantitatively, we calculated the adjusted Rand index (ARI) between

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Figure 2. Unsupervised annotations of the DKD Kidney samples

(A) Annotations from SCGP and other unsupervised annotation methods recognized tissue structures aligned with manually annotated compartments. Nodes representing cells are colored according to the assigned clusters/partitions in the top images, and colors not listed in the legend (e.g., cyan) refer to clusters/ partitions that cannot be matched to any compartment. Mismatched nodes are highlighted in red in the bottom images.

(B) ARIs were calculated between unsupervised annotations and manual annotations. SCGP performed significantly better than all other methods (Wilcoxon signed-rank test).

(C) For each manually annotated compartment, F1 scores were calculated between manual annotations and the most overlapped cluster/partition.

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manually annotated compartments and unsupervised partitions [\(Figure 2B](#page-4-0)). SCGP achieved the top performance with a median ARI of 0.60, significantly outperforming all other methods (Wilcoxon signed-rank test). Since ARI downweighs smaller/rarer compartments (e.g., glomeruli, blood vessels), we calculated additional alignment and accuracy measurements, including independent F1 scores for each compartment [\(Figures 2C](#page-4-0), [S2B](#page-12-4), and S2C; [STAR Methods](#page-16-0)). These metrics better reflect performances on less common structures, many of which have great functional significance. We found that unsupervised discovery of tissue architecture with graphs (UTAG)^{[15](#page-13-4)} and SCGP performed the best at recognizing glomeruli (F1 = \sim 0.8), while SpiceMix 24 24 24 and SpaGCN 32 32 32 excelled at recognizing tubule structures. Overall, SCGP achieved the best average F1 score in discerning all manually annotated compartments.

To further validate SCGP-identified partitions, we highlighted their relative protein expression ([Figures 2](#page-4-0)D and [S2D](#page-12-4)). The heat-map corresponds well to our expectation, with CCR6^{[42](#page-14-2)} and Nes- \sin^{43} among the top biomarkers for glomeruli and CXCR3^{[44](#page-14-4)} and $MUC1⁴⁵$ $MUC1⁴⁵$ $MUC1⁴⁵$ for proximal and distal tubules. Notably, by grouping performance metrics by sample disease progression, 41 we found that the quality of unsupervised partitions degraded substantially with disease progression (severe DKD, class IIB/III). This indicates how normal tissue structures and functions are dysregulated in DKD and highlights the challenge of performing consistent annotations across disease states ([Figure 2E](#page-4-0)).

SCGP identifies major structures of the brain

Next, we assessed SCGP's performance on spatial transcriptomics measurements of human and mouse brains. We first em-ployed a Visium^{[7](#page-12-3)} dataset of human dorsolateral prefrontal cortex $(DLPFC).$ ^{[46](#page-14-7)} In contrast to the mIF approach, the Visium platform features a grid of spatially barcoded oligonucleotide arrays that can be used for mRNA capture and library preparation. Each array (i.e., spot) may contain multiple cells. We adapted our method by treating each spot as a node and defining edges based on the grid and gene count information [\(STAR Methods\)](#page-16-0).

DLPFC contains 12 samples annotated with 7 compartments: 6 cortical layers (L1–L6) and white matter. We directly compared our method against existing tools developed for this modality, including BayesSpace,^{[29](#page-13-16)} SpaGCN, and SpiceMix, in recognizing manually annotated layers. Clustering/partitioning was performed independently on each sample. Results on a representative sample are demonstrated in [Figure 3](#page-6-0)A, revealing a clear layer-wise pattern in the tissue structures. Among the benchmarked methods, SCGP and SpiceMix achieved the top ARI scores, exhibiting superior alignment with the ground-truth compartments marked by dashed lines. Quantitative metrics ([Figures 3](#page-6-0)B and [S4E](#page-12-4)) demonstrated that SCGP achieved comparable, if not superior, performance(median ARI = 0.56 , median F1 = 0.65) to unsupervised annotation tools designed specifically for Visium spatial transcriptomics.

We subsequently tested SCGP on a MERFISH-profiled section of a whole mouse brain, 47 where SCGP readily recognized

major brain parcels, exhibiting an overall ARI score of 0.75 [\(Fig](#page-6-0)[ure 3C](#page-6-0)). Furthermore, SCGP with higher granularity accurately identified finer structures aligned with detailed parcellation [\(Figure S5\)](#page-12-4).

Next, we further evaluated if jointly partitioning multiple sections would improve performance. Following the experiment design by Chidester et al., 24 24 24 we applied unsupervised annotation tools to the combination of four samples in DLPFC collected from the same donor ([Figures S4A](#page-12-4)-S4C). While BayesSpace, SpaGCN, and SpiceMix performed better when jointly partitioning samples, UTAG and SCGP yielded worse results. Motivated by the aforementioned challenge of partitioning different disease states as well as how joint partitioning might suffer from intersample variance, we explored means of generalizing consistent annotations across conditions.

SCGP extends existing tissue structures to unseen samples

In practical applications, generalizing a set of curated annotations across different experiments, conditions, or disease states is required to conduct analyses at larger scales of data and a long-standing goal of data integration. Generalization is useful for performing inference on prospective unseen data, suppressing unwanted noise or batch effects, comparing samples across disease conditions, and detecting previously unseen and unannotated disease states. Existing unsupervised annotation tools have limited or non-existent support for this functionality. Most methods (e.g., UTAG, SpaGCN) explicitly require either retraining or refitting the clustering model or the addition of separate prediction models to extend existing partitions. Therefore, to address this need, here, we present a specialized reference-query extension pipeline, SCGP-Extension.

SCGP-Extension begins with the annotation of a small group of high-quality reference samples, for example, by using SCGP. Resulting partitions are assumed to represent the ground-truth structures and referred to as reference partitions [\(Figure 1](#page-3-0)D). Next, we define pseudo-nodes as representative data nodes for each reference partition ([STAR Methods](#page-16-0)). These pseudo-nodes are explicitly added into the SCGP graphs of unseen query samples and connected to other query nodes based on feature similarity [\(Figure 1](#page-3-0)E). Finally, the graph community detection step is conducted as before, with partitions of the pseudo-nodes pre-assigned. Consequently, query nodes resembling pseudo-nodes are assigned the corresponding partition, while query nodes that do not resemble any reference partitions form their own groups and assigned as newly discovered or ''novel'' partitions ([Figure 1](#page-3-0)F). On simulated data, SCGP-Extension robustly identified both known and novel partitions [\(Figures S1D](#page-12-4) and S1E).

To demonstrate how SCGP-Extension improves partitioning performance, we examined its application in the DLPFC joint partitioning experiment. Here, we established reference

⁽D) Signature protein biomarkers for SCGP-identified partitions match expectations of kidney tissue structures.

⁽E) SCGP annotations on samples with different classes of DKD show varying levels of alignment and accuracy. Diabetes mellitus (DM, with healthy kidney) and DKD classes represent the different progression stages assigned by a nephrologist following the Tervaert classification.^{[41](#page-14-6)} See also [Figure S2.](#page-12-4)

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Figure 3. Unsupervised annotations of human and mouse brain samples

(A) A representative DLPFC sample was annotated using SCGP and other unsupervised annotation methods. Note that layer 2 and layer 4 were not fully recognized. Boundaries between ground-truth layers are annotated as dashed lines.

(B) ARIs were calculated between unsupervised annotations and ground-truth layers on 12 samples, with each sample annotated independently.

(C) A mouse brain section profiled by MERFISH was annotated using SCGP, from which major brain parcels were identified. Boundaries between parcels are annotated as dashed lines.

See also [Figures S4](#page-12-4) and [S5.](#page-12-4)

annotations on one sample [\(Figure 3A](#page-6-0)) and extended them to the rest, simulating the application scenario where a small set of gold-standard annotations are acquired and generalized to the entire dataset. SCGP-Extension considerably outperformed joint clustering approaches in alignment and accuracy ([Figures S4](#page-12-4)A and S4D). In addition, we evaluated approaches that directly utilized partial ground-truth annotations, consisting of both predictive modeling and SCGP-Extension approaches, where SCGP-Extension using partial labels demonstrated the best performances ([Figure S4D](#page-12-4)).

SCGP-Extension can also generalize across disease conditions and help identify unique disease states. To demonstrate this, we applied SCGP-Extension to generalize primary partitions defined on healthy DKD samples to severe DKD cases. In this scenario, glomeruli exhibit elevated collagen four and decreased native marker expression (e.g., CCR6). Unsupervised SCGP, joint partitioning methods, and supervised classifiers (XGB Prediction) all struggled to correctly identify tissue characteristics of severe DKD [\(Figures 4](#page-7-0)A, [2](#page-4-0)E, and [S2](#page-12-4)A), often misrecognizing fibrotic glomeruli [\(Figure 4A](#page-7-0), arrows). However, SCGP-Extension preserved most of the original partitions while uncovering two additional structures: the purple partition outlined fibrotic glomeruli, characterized by the depleted native biomarkers and enriched collagen expression, and the red partition, characterized by elevated CD45 and CD68 expression ([Figures 4](#page-7-0)A and 4B), suggesting the infiltration of immune cells

(e.g., macrophages). Compared to joint partitioning and predictive modeling, SCGP-Extension delivered more accurate results both visually and quantitatively [\(Figure 4C](#page-7-0)). Furthermore, it highlighted unique disease-specific partitions with ready biological interpretations, offering valuable insights into disease progression.

We next assessed the performance of SCGP in another healthy versus disease comparison. Here, kidney tissue samples from patients who experienced transplant rejection were parti-tioned (TR Kidney dataset, [STAR Methods](#page-16-0)), characterized by heavily deformed native kidney structures, substantial inflammation, and immune cell infiltration. Although the measured protein markers and disease context were different, our analysis derived the same set of tissue structures as before [\(Figure 4D](#page-7-0)). Extending partitions from reference samples with heavy inflammation to query samples with minimal immune activities produced consistent structures, which enabled direct comparisons be-tween conditions ([Figures 4](#page-7-0)D, [S6](#page-12-4)A, S6B, and S6G). Notably, SCGP-Extension effectively partitioned a region containing background signal artifacts, demonstrating effective handling of data issues that would typically require manual intervention to find and fix [\(Figures S7A](#page-12-4)–S7D).

To show that SCGP-Extension can identify novel anatomical structures in query samples, we evaluated unsupervised annotation methods on an IMC dataset collected on healthy lung spec-imens (Lung IMC dataset, [STAR Methods](#page-16-0)), which were manually

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annotated for anatomical structures. SCGP achieved remarkable alignment with ground truth, but tissue structures across samples were disintegrated and assigned to different partitions [\(Figure S8A](#page-12-4)). To address this issue, we conducted primary SCGP on a small subset of samples and extended partitions to the remaining samples. Notably, SCGP-Extension identified two anatomical structures (submucosal glands and cartilage) that were absent in the reference while preserving all known structures [\(Figure 4](#page-7-0)E). SCGP-Extension also achieved significantly better alignment scores (median ARI = 0.564 ; [Figures 4](#page-7-0)F, [S8B](#page-12-4), and S8C) than all other benchmarked methods (Wilcoxon signed-rank test).

SCGP-Extension can also help mitigate batch effects between experiments. On a cohort of skin samples collected from four separate experiments (UCSF Derm dataset, [STAR Methods](#page-16-0)), we employed unsupervised annotation methods to define tissue structures. The UCSF Derm dataset comprised samples of different skin conditions sharing similar anatomical structures ([Table S1](#page-12-4)). Both UTAG and SCGP failed to link tissue structures from different samples [\(Figures S7](#page-12-4)E and S7F). We then attempted SCGP-Extension by defining reference partitions on samples from one experiment ([Figure S6](#page-12-4)C) and extending them to the rest ([Figures S6D](#page-12-4)–S6F). Regardless of biases in the experiments and differences in disease conditions, SCGP-Extension successfully recognized consistent partitions across samples reflecting anatomical structures [\(Figures 4](#page-7-0)G and [S7H](#page-12-4)).

SCGP partitions assist downstream analysis of disease states

Partitions acquired by unsupervised annotation using SCGP reflect the anatomical and functional structures of the subject tissues. In this section, we demonstrate how these tissue structures can facilitate the analysis of disease states and enable the discovery of key insights regarding disease-relevant partitions.

In the DKD Kidney experiments, samples were collected from individuals with different DKD classes. Based on the partitioning of these samples, we were interested in exploring correlations between tissue structures and disease progression. [Figure 5A](#page-9-0) illustrates three representative samples of different DKD classes along with their tissue structures annotated by SCGP and SCGP-Extension. Clear visual differences between samples can be observed: tubules and glomeruli were denser in the healthy sample, while these structures gradually deformed over the course of DKD, accompanied by fibrosis and infiltration

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of immune cells. These changes were also reflected in the tissue structures: a significant increase (*p* < 0.001, Jonckheere-Terpstra test⁴⁸) in the area proportion of the basement membrane partition was observed across samples ([Figure 5](#page-9-0)B), reflecting the degradation of normal kidney structures. Immune aggregate (red) and fibrotic glomerular (purple) partitions were identified in the severe DKD sample with SCGP-Extension, which were not present in healthy and mild DKD samples.

Furthermore, to characterize how DKD affects glomerular functions, we assessed the protein biomarker expression of individual glomerulus (dashed circles in [Figure 5](#page-9-0)A), annotated by deriving connected components of the SCGP glomeruli partitions. The results suggested that glomeruli undergo significant loss of native proteins (CCR6) throughout the disease progression (*p* < 0.001, Jonckheere-Terpstra test; [Figure 5C](#page-9-0)), with high intra-sample and inter-sample heterogeneity.

We next applied the partition-based analysis to the UCSF Derm dataset comprising normal samples and atopic dermatitis samples. Representative samples visualized in [Figure 5D](#page-9-0) show notable differences in terms of epidermal thicknesses and immune cell densities. We verified these visual signatures using SCGP partitions: the area proportion of the immune aggregate partition demonstrated significant increases in atopic dermatitis samples $(p < 0.001$, two-sided two-sample t test; [Figure 5](#page-9-0)E). Thicknesses of the epidermal layers were characterized using the contours of the corresponding partition, which showed significant increases as well (*p* < 0.001, two-sided two-sample t test; [Figure 5](#page-9-0)F). Patients with atopic dermatitis also exhibit a much more heterogeneous distribution of epidermal thickness.⁴⁹ Thus, SCGP empowers rapid quantitative insights from spatial molecular data.

Atlas-scale structural annotation of TMEs

Lastly, we assessed the applicability of SCGP in identifying structures in neoplastic diseases, which are typically characterized by the complete disruption of normal cellular architectures and uncontrolled growth of abnormal cells. We focused our analysis on tumor microenvironments $(TMEs)$,^{[50](#page-14-11)} highly structured and complex systems containing cancer cells and a variety of non-malignant cell types, including immune cells and stromal cells.

Identifications of major structures in TMEs, such as tumor and stromal regions, are crucial tasks that facilitate further investigation of critical topics including tumoral heterogeneity, dynamics

Figure 4. Versatile applications of SCGP-Extension

(G) In the UCSF Derm dataset, SCGP-Extension consistently partitioned samples from multiple experiments with different skin conditions. See also [Figures S6–S8.](#page-12-4)

⁽A) Compared to joint partitioning or predictive models, SCGP-Extension better recognized the fibrotic glomeruli (purple dashed circles) in severe DKD samples and identified a previously unseen immune aggregates partition (red boxes).

⁽B) Extension to severe DKD samples improved partition alignment and accuracy.

⁽C) Heatmap shows signature protein biomarkers for extended partitions in the severe DKD samples; note the additional fibrotic glomeruli and immune aggregates partitions.

⁽D) In the TR Kidney dataset, SCGP partitioned a cohort of kidney samples with heavy inflammation (top row). SCGP-Extension further extended the partitions to control samples with minimal immune responses (bottom row).

⁽E) In the Lung IMC dataset, SCGP-Extension recognized an anatomical structure, submucosal glands (purple), in the query samples that was not seen in reference samples.

⁽F) ARIs calculated between unsupervised annotations and manual annotations show that SCGP-Extension achieved significantly better alignments (Wilcoxon signed-rank test).

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of tumor-immune cell interactions, and therapeutic targeting, among others. Extensive studies have been proposed to perform tumor classification and segmentation on histopathology images using various deep learning methods, $36,51-53$ $36,51-53$ most of which adopt supervised learning strategies. Spatial omics, on the contrary, facilitate the detection of these structures in TMEs in a fully unsupervised manner.

We benchmarked SCGP against other annotation tools to a cohort of head and neck tumor samples (University of Pittsburgh Medical Center-Head and Neck Cancer [UPMC-HNC]; [STAR](#page-16-0) [Methods\)](#page-16-0) containing 36 tumor cores and 175,000 cells. As illus-trated in [Figures 6](#page-11-0)A and [S9](#page-12-4)A, a distinct separation between tumor and stromal regions was observed after partitioning at a coarse resolution. Quantitative metrics showed a median ARI exceeding 0.8 and an F1 score of 0.94 from SCGP [\(Figure 6B](#page-11-0)), which not only significantly outperformed competing unsupervised methods but also matched the reported performance of H&E-based supervised segmentation tools. These outcomes are further verified in another cancer study [\(Figures S9](#page-12-4)B and S9C).

We also performed more granular partitioning of TMEs. By deploying our SCGP-Extension pipeline, we acquired consistent an-notations of TMEs at a higher granularity ([Figure 6](#page-11-0)C). In addition to the tumor/stroma separation, several novel partitions with distinct biomarker profiles and cell-type compositions were identified [\(Fig](#page-11-0)[ure 6](#page-11-0)D). For example, an immune-rich partition was isolated from stroma regions, which exhibited much denser immune cell populations including B cells, T cells, and macrophages. Similarly, a unique CD15/granulocyte-rich partition was detected in the tumor regions of multiple samples, which showed distinct morphology and biomarker enrichment. Such results highlight the power of tissue structure annotations, from which distinct spatial organization of different cell types can be discovered and examined.

DISCUSSION

In this work, we present SCGP, an unsupervised annotation tool for spatial transcriptomics and proteomics measurements. SCGP embeds spatial information and molecular features into graph representations and performs Leiden graph community detection to identify partitions corresponding to anatomical and functional structures. The reference-query extension pipeline, SCGP-Extension, robustly generalizes existing tissue structures to unseen samples and recognizes unique disease states and anatomical structures. Our experiments demonstrate the power of SCGP and SCGP-Extension in identifying tissue structures in various data cohorts and show how tissue structures assist downstream biomedical research and discoveries.

We compared SCGP against other unsupervised annotation tools that similarly utilize both spatial information and molecular profiling output. Methods such as CN, UTAG, and Spatial-LDA define different concepts of neighborhoods, usually based on distance thresholding, and annotate them through unsupervised clustering. These methods tend to be inflexible to structures of different spatial scales. Another class of methods also uses graph representations of tissue samples to model cellular organization. Computational tools including latent variable modeling and graph neural networks are applied on these graphs to annotate nodes (cells) according to their tissue context. However, spatially disconnected samples are difficult to annotate with these tools. SCGP and SCGP-Extension follow a graph representation approach and enhance it with nearest-neighbor feature edges to weave multiple samples into one cohesive graph.

Moreover, SCGP-Extension is the first method that addresses the long-standing need of generalizing structures to previously unseen samples. SCGP-Extension resembles supervised learning tools in that models apply knowledge learned from the reference samples (i.e., training data) to unseen query samples (i.e., test data). Simultaneously, SCGP-Extension can isolate previously unseen tissue structures during inference. In practice, we have demonstrated that SCGP-Extension can help overcome common challenges including experimental artifacts, batch effects, and different disease conditions. More importantly, it can help uncover unique disease states and anatomical structures.

SCGP and SCGP-Extension offer outstanding running time and memory consumption performances compared to many unsupervised annotation tools, some of which require extensive parameter estimations or optimizations [\(STAR](#page-16-0) [Methods](#page-16-0); [Table S2\)](#page-12-4). In practice, this advantage allows for applications to much larger datasets and facilitates model parameter tuning to obtain optimal partitions.

SCGP and SCGP-Extension are validated on both simulated and real datasets to model various spatial proteomics and transcriptomics measurements effectively, capturing tissue structures of diverse shapes and sizes in systems with varying levels of complexities. Minimal parameter adjustments [\(STAR](#page-16-0) [Methods](#page-16-0)) are required. The two major parameters for SCGP are the density of feature edges and the granularity of partitioning [\(Figures S3E](#page-12-4)-S3G). One additional parameter, the extent of extension, is introduced in SCGP-Extension to control the balance between generalizing existing partitions and exploring new partitions. In examples involving different disease conditions, certain tissue structures may experience changes in their expression profiles (e.g., fibrosis, immune infiltration), and the

Figure 5. Downstream analysis of disease states with SCGP-identified partitions

(F) Epidermal layers show significant thickening in atopic dermatitis samples (*p* < 0.001, two-sided two-sample t test).

⁽A) Three representative samples of different DKD classes are illustrated; note the fibrosis of glomeruli and increase in the area of the basement membrane partition (light blue-colored nodes). Glomeruli are segmented by dashed circles.

⁽B) Boxplot shows that the area proportion of the basement membrane partition significantly increases (*p* < 0.001, Jonckheere-Terpstra test) in DKD samples. Each dot represents a tissue sample.

⁽C) Expression of native proteins (CCR6) significantly decreases (*p* < 0.001, Jonckheere-Terpstra test) in glomeruli of patients with DKD. Each dot represents an individual glomerulus, and each box summarizes glomeruli from one patient. Note the heterogeneity observed within a single tissue sample (A, bottom row) and between patients from the same DKD class.

⁽D) Two samples from the UCSF Derm dataset with different skin conditions are illustrated.

⁽E) Area proportion of the immune aggregate partition significantly increases in atopic dermatitis samples (*p* < 0.001, two-sided two-sample t test).

Figure 6. Annotation of major structures in TMEs by SCGP

(A) Coarse partitioning identified tumor/stroma regions in TMEs.

(B) ARIs and F1 scores were calculated between unsupervised annotations and manual tumor/stroma annotations. SCGP achieved significantly better performances than other methods (Wilcoxon signed-rank test).

(C) Fine-grained partitions detected additional tissue structures in TMEs.

(D) Heatmaps show distinct cell-type enrichment and protein expression profiles of different tissue structures identified by SCGP. See also [Figure S9.](#page-12-4)

decision of whether to integrate or separate these structures will depend on the specific downstream applications.

Looking ahead, SCGP opens new opportunities for analyzing and understanding spatially resolved molecular profiling data. By inserting a middle layer between cell-level annotations and sample-level characteristics, it facilitates analysis tailored to specific structures of interest at scale.

Limitations of the study

We acknowledge the presence of certain caveats in SCGP and SCGP-Extension, which hinder their applications in specific scenarios. One major shortcoming is that SCGP appears to be less suitable for identifying thinly layered structures. Due to the design of the hybrid graph, spatial edges are isotropic for the purpose of community detection. Thin-layer structures exhibiting much denser spatial connections in their normal directions than tangential directions are harder to detect [\(Figures S1A](#page-12-4) and [S4](#page-12-4)A). In the DLPFC study, alternative methods (e.g., BayesSpace, SpiceMix) can identify thinner structures under more granular settings, while SCGP recognizes structures in the orthogonal direction. Refining the spatial edges to reflect the anisotropy of tissues would be a direction to improve the performance of SCGP.

Another limitation is that the outputs of SCGP and SCGP-Extension are hard class assignments. This lack of flexibility may limit applications in scenarios where probabilistic or mixture

outputs are preferred. One way to accommodate this limitation is to collect ensembled outputs from running the algorithm using different random seeds.

Case studies presented in this work used solely biomarker expression to define tissue structures. This can be further improved by integrating multi-modal data. For instance, inclusion of morphology information (e.g., through embeddings of H&E staining) would greatly enrich the feature space and allow SCGP to account for cellular- and tissue-level morphological differences.

In addition, the concept of reference-based annotation can be further extrapolated to prior-based annotation. Specifically, instead of fully unsupervised partitioning, incorporating prior knowledge of the expected tissue structures (e.g., signature biomarkers, sizes, and shapes) could potentially yield better-aligned results and will be a direction for future improvements. Lastly, we acknowledge that evaluations in this study were not exhaustive of all possible methods and datasets. We selected representative methods spanning diverse approaches to establish the benchmark for SCGP. Many recent methods^{54–57} were not included due to time constraints despite their competitive performances. Moreover, quantitative performances (i.e., ARI, F1 scores) were calculated based on manual annotations, which are subjective and may contain intrinsic biases. To mitigate dataset-specific biases, we performed comparisons across diverse datasets from different institutions, though realistic performances will still vary based on the specific tissue/disease environments.

STAR+METHODS

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

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SUPPLEMENTAL INFORMATION

Supplemental information can be found online at [https://doi.org/10.1016/j.](https://doi.org/10.1016/j.crmeth.2024.100838) [crmeth.2024.100838](https://doi.org/10.1016/j.crmeth.2024.100838).

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AUTHOR CONTRIBUTIONS

Conceptualization, Z.W. and A.E.T.; methodology, Z.W.; software, Z.W., E.A.G.B., and B.C.; validation, A.K., E.W., and M.K.R.; investigation, Z.W.; resources, N.A.B., V.C., R.J.C., J.B.C., and M.A.; data curation, A.K. and M.M.; writing – original draft, Z.W. and A.E.T.; writing – review & editing, Z.W. and A.E.T.; supervision, J.Z., A.T.M., and A.E.T.

DECLARATION OF INTERESTS

Z.W., A.K., M.M., E.A.G.B., B.C., M.K.R., A.T.M., and A.E.T. are affiliated with Enable Medicine as employees and/or shareholders. J.Z. is a member of Enable Medicine's scientific advisory board.

DECLARATION OF GENERATIVE AI AND AI-ASSISTED TECHNOLOGIES IN THE WRITING PROCESS

During the preparation of this work, the authors used GPT-4 in order to polish the writing of the manuscript. After using this tool/service, the authors reviewed and edited the content as needed and took full responsibility for the content of the publication.

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STAR★METHODS

KEY RESOURCES TABLE

RESOURCE AVAILABILITY

Lead contact

Further information and requests for resources should be directed to and will be fulfilled by the lead contact, Alexandro E. Trevino ([alex@enablemedicine.com\)](mailto:alex@enablemedicine.com).

Materials availability

This study did not generate new unique materials or reagents.

Data and code availability

- d External spatial proteomics and transcriptomics datasets used in this study (Human DLPFC Visium dataset, adult mouse brain MERFISH dataset, Lung IMC dataset, UPMC head-and-neck cancer CODEX dataset) are publicly available as of the date of publication. Processed CODEX datasets (DKD kidney CODEX dataset, UCSF derm CODEX dataset, TR kidney CODEX dataset, Stanford pancreatic cancer CODEX dataset) have been deposited at zenodo. All accession numbers are listed in the [key](#page-16-1) [resources table.](#page-16-1) Raw data and images will be accessible through links posted at: [https://gitlab.com/enable-medicine-public/](https://gitlab.com/enable-medicine-public/scgp) [scgp](https://gitlab.com/enable-medicine-public/scgp). They will also be available from [lead contact](#page-16-3) (A.E.T.) upon request.
- \bullet SCGP was implemented based on Annotated data^{[61](#page-14-14)} (<https://anndata.readthedocs.io/en/latest/>) and emObject^{[62](#page-14-15)} ([https://docs.](https://docs.enablemedicine.com/emobject/) [enablemedicine.com/emobject/](https://docs.enablemedicine.com/emobject/)). All source codes are publicly available at [https://gitlab.com/enable-medicine-public/scgp.](https://gitlab.com/enable-medicine-public/scgp) An archival DOI is listed in the [key resources table.](#page-16-1)
- d Any additional information required to re-analyze the data reported in this paper will be available from [lead contact](#page-16-3) (A.E.T.) upon request.

EXPERIMENTAL MODEL AND STUDY PARTICIPANT DETAILS

Eight spatial proteomics and transcriptomics datasets collected from diverse tissue types were examined in this work. An overview of the statistics and primary phenotypes of these datasets can be found in [Table S1](#page-12-4). For external datasets, please refer to the source publications for full details. For datasets introduced in this study:

TR Kidney⁶³: Kidney samples were obtained from patients who underwent allograft nephrectomy at Stanford University Medical Center under Institutional Review Board-approved protocols. One TMA was constructed using 2 mm cores of cortical tissue, then stained and acquired on the PhenoCycler Fusion using a 51-plex biomarker panel, from which 5 normal and 17 transplant rejection kidney regions were used in this study. Further clinical or patient demographic data related to these samples was not selected or considered in this study.

Stanford-PC: De-identified tissue samples were obtained from the surgical pathology archives of Stanford Medical Center under Institutional Review Board-approved protocols. Tumor cores (1 mm diameter) were sampled and profiled with CODEX using immune-cell focused biomarker panels. One coverslip containing 148 cores was analyzed in this study. Clinical or patient demographic data related to these samples was not selected or considered in this study. All cores were manually annotated for tumor regions based on epithelial protein markers.

METHOD DETAILS

Datasets

This section briefly describes phenotypes and properties of each dataset analyzed in this study.

DKD Kidney⁴⁰: Kidney samples were obtained from patients with diabetes and healthy kidneys (DM, five individuals), DKD classes IIA, and IIB (two individuals per class), IIA-B intermediate (two individuals), and III (one individual). Twenty-three cores, each 0.5 mm in diameter, were sampled from twelve tissue blocks and assembled into a tissue microarray (TMA). The TMA block was further sectioned into 5 µm slices. One TMA section was imaged and characterized using the CO-Detection by indexing (CODEX) platform. After excluding medulla samples and quality control, a total of 17 cortical section samples across various DKD classes were acquired. Each sample was imaged for 21 protein biomarkers (see columns in [Figure S1B](#page-12-4)).

DLPFC⁴⁶: Spatial gene expression in human postmortem dorsolateral prefrontal cortex tissue sections was profiled using two pairs of ''spatial replicates'' from three independent neurotypical adult donors on the Visium platform, each pair comprising two directly adjacent, 10 μ m serial tissue sections, with the second pair located 300 μ m posterior to the first. In total 12 samples are collected and examined. We downloaded the filtered count matrices for all 12 samples from the spatialLIBD project.^{[64](#page-14-20)} In the independent partitioning experiment ([Figure 3\)](#page-6-0), we filtered the count matrices to exclude spike-in genes, mitochondrial genes, and genes that have nonzero expression in fewer than three spots. Expression matrices were normalized to have the same total counts per spot (median of all pre-normalize spots), log-transformed and reduced to the top 50 principal components. In the joint partitioning exper-iment [\(Figure S2\)](#page-12-4), we followed the preprocessing steps outlined in SpiceMix²⁴: Genes having nonzero expression in less than 10% of spots were removed. Expression matrices were normalized to have total counts of 10,000 per spot and log-transformed. We further reduced the expression to the top principal components.

Adult Mouse Brain (ABCA-1)⁴⁷: This dataset comprises in total 147 coronal sections of a single adult mouse brain, profiled with a 1122 gene panel using MERFISH. Cells were annotated for cell types and mapped to the Allen mouse common coordinate framework.⁶⁵ We downloaded preprocessed biomarker expression matrices, cell types, and CCF coordinates from the source publication. Two sections (ABCA-1.071, ABCA-1.079) were used to assess SCGP and SCGP-Extension. In the experiments, the 300 most variable genes of the reference section (ABCA-1.079) were used as features.

TR Kidney^{[63](#page-14-19)}: Kidney samples were obtained from patients who underwent allograft nephrectomy. Briefly, a TMA was constructed using 2 mm cores of cortical tissue. The TMA comprised 7 samples of normal, peritumoral renal cortex (from patients undergoing native nephrectomy for tumor removal), and 43 samples of cortex from patients undergoing allograft nephrectomies. After sectioning of the tissue block, only 41 cores from allograft nephrectomies remained. An FFPE embedded tissue microarray core of human kid-ney samples were stained and acquired on the PhenoCycler Fusion using a 51-plex biomarker panel [\(Figure S3C](#page-12-4)) by Enable Medicine.

Lung IMC^{[15,](#page-13-4)66}: Lung samples were acquired from three healthy donor lung specimens. In total 26 samples were imaged using IMC with 28 biomarkers (see columns in [Figure S4](#page-12-4)D). Tissue samples were collected with a particular focus on airways extending from proximal bronchi and succeeding divisions to terminal and respiratory bronchioles. Each image was manually annotated with organ-specific microanatomical domains: airways, connective tissue, submucosal glands, vessels, cartilages, and alveolar space. These manually annotated domains were used as labels for unsupervised annotations. Additionally, cells in these samples were phenotyped into seven broad clusters of cell identity: CD8 T cells, macrophages, mast cells, smooth muscle cells, endothelial cells, epithelial cells, and connective tissue cells. Cell type information was used in CN and Spatial-LDA, other methods only used biomarker expression data. We downloaded preprocessed biomarker expression matrices and domain/cell type annotations from the source publication.^{[59](#page-14-17)}

UCSF Derm⁶⁰: Tissue samples were acquired as 2–4 mm punch or shave skin biopsy specimens from healthy control and atopic dermatitis patients. In total, 44 skin samples were obtained and stained in 4 experiments with varying biomarker panels. See [Table S1](#page-12-4)

for a detailed breakdown of experiments and patient skin disorders. 35 shared protein biomarkers were used in the unsupervised annotation analysis with UTAG and SCGP.

Stanford-PC and UPMC-HNC^{[16](#page-13-5)}: These two datasets were derived from studies of pancreatic cancer and head-and-neck cancer. Tumor cores (1 mm diameter) were sampled and profiled with CODEX using immune-cell focused biomarker panels. [Figure 6D](#page-11-0) shows some of the representative protein biomarkers in the panel. We adapted one coverslip from each study to assess SCGP and SCGP-Extension. Stanford-PC contained 148 cores in total, which were further divided into two subsets: a higher-quality reference set (63 cores) and a lower-quality query set (85 cores). UPMC-HNC contained 36 cores. Clinical data related to these samples were not used in this study. All cores were manually annotated for tumor regions by an expert, and other foreground tissues were treated as nontumor/stroma regions.

Preprocessing

DLPFC, Lung IMC, and ABCA-1 datasets were downloaded and normalized as specified above.

For the CODEX datasets, we followed the preprocessing pipeline established in the prior work by Wu et al.^{[16](#page-13-5)} Briefly, a neural network-based cell segmentation tool DeepCell^{[20](#page-13-9)} was applied to DAPI images to identify nuclei, which were further dilated to obtain whole-cell segmentation.

Next, the biomarker expression for biomarker *j* in cell *i* was computed following the strategy below^{[67](#page-14-23)}:

- (1) For channel *j*, mean pixel intensity within the cell segmentation mask of cell *i* was calculated and denoted as $x_i^{(j)}$. The set of expression values for all cells in the same sample $\{x_1^{(j)}, x_2^{(j)}, ...\}$ was denoted as $X^{(j)}$.
Normalized expression value for channel *i* was calculated using quantile permalization
- (2) Normalized expression value for channel *j* was calculated using quantile normalization and arcsinh transformation:

$$
f(x_i^{(j)}) = arcsinh\left(\frac{x_i^{(j)}}{5Q(0.2;X^{(j)})}\right)
$$

in which *Q*(0.2; *X⁽ⁱ⁾*) represents the 20-th quantile of *X⁽ⁱ⁾* and *arcsinh* is the inverse hyperbolic sine function. The set of all normalized
oxpression values $(f(\mathbf{v}^{(j)}) - f(\mathbf{v}^{(j)}) = 1$ was depoted as $(Y^{(j)})$ expression values $\{f(x_1^{(j)}), f(x_2^{(j)}), ...\}$ was denoted as $(X^{(j)})$.

(1) *Z* score of normalized expression value was calculated:

$$
z(x_i^{(j)}) = \frac{f(x_i^{(j)}) - \text{MEAN}\big(f\big(X^{(j)}\big)\big)}{SD\big(f\big(X^{(j)}\big)\big)}
$$

It should be noted that SCGP does not require any cell clustering or classification inputs, it infers partitions using biomarker expression of cells.

Spatial cellular graph partitioning (SCGP)

SCGP is an unsupervised annotation tool that recognizes tissue structures by partitioning graphs constructed based on the spatial organization of cells (or other units) in the subject tissue sample(s). The SCGP pipeline comprises the following steps.

Nodes

Nodes represent small spatial regions in the tissue, and are indivisible units throughout the partitioning process. In this study, we employed three different strategies for defining nodes.

- (1) Cells: Nodes are defined based on individual cells, which are identified via the cell segmentation preprocessing step specified above. Biomarker expression values are calculated and normalized accordingly, and these values are set as the node features.
- (2) (Visium) Spots: Nodes are defined based on the barcoded spots used in the Visium platform, each measuring gene expression in a circular area 55 μ m in diameter. The normalized gene expression values or top principal components are set as node features.
- (3) Patches: In the patch-based SCGP experiment ([Figure S10\)](#page-12-4), nodes are defined based on small square patches with 12 μ m side lengths on the mIF images, sampled using a sliding window mechanism (stride equals patch side length). Only patches within the spatial range of tissue are retained (>85% overlap), and the average fluorescence intensities for each biomarker channel are used as node features.

Spatial edges

Spatial edges are constructed between spatially adjacent nodes to embed the spatial closeness relationship into the graph.

- (1) Cells: A Delaunay triangulation is conducted on centroid coordinates of all cells from the same sample. Node pairs that share edges in the triangulation output are connected, excluding any edge exceeding $35 \mu m$ in length.
- (2) (Visium) Spots: Nodes (i.e., spots) are spatially arranged in a close-packing manner. Each node is connected to its six closest neighboring nodes.
- (3) Patches: Nodes (i.e., patches) are spatially arranged in a regular 2D grid. Each node is connected to its four immediately adjacent nodes.

Feature edges

Feature edges are constructed between nodes with similar biomarker expression profiles. For node *i*, its *k* nearest neighboring nodes in the expression space are identified based on Euclidean distances between node features (e.g., z-scored protein expression values, principal components of the gene expression). We used the nearest neighbor descent^{[68](#page-14-24)} approximate queries implemented in PyNNDescent for better computational efficiency.

In practice, *k* is a hyperparameter that controls the balance between spatial coherence and expression consistency within partitions. We typically set it to an integer between 1 and 4 so that feature edges account for about 25%–50% of all edges ([Figures S3](#page-12-4)E and S3F). In general, less feature edges will generate spatially smoother partitions, though minor changes in *k* do not significantly change partition outcomes in our experiments. However, a larger *k* (>5) will lead to spatially fragmented partitions, which resemble the results of direct Leiden clustering on feature vectors. This design choice of sparse feature edges differentiates SCGP from other Leiden-based unsupervised annotation tools.^{[15,](#page-13-4)[27](#page-13-14)}

Graph community detection

Nodes, spatial edges, and feature edges define the spatial cellular graph input for SCGP. We used the Leiden algorithm^{[39](#page-14-0)} to detect graph communities.

We adapted the python implementation in leidenalg, and we used the Constant Potts Model^{[69](#page-14-25)} as the quality function for community detection (leidenalg.CPMVertexPartition). Additional arguments include.

- (1) Edge weights: For each edge, its weight is defined as the inverse of Euclidean distance between the node features of the two nodes it connects. We further normalized all edge weights by their median value.
- (2) Resolution parameter (γ) : γ controls the density of the output communities.

Note that the resolution parameter γ is the second major hyperparameter of SCGP, regulating the granularity of the output partitions. We empirically tuned γ to generate the desired amount of partitions (e.g., 2 for TME samples, 10+ for whole brain samples). This process was informed by the understanding and expectations of the subject tissues ([Figure S3G](#page-12-4)).

Note that variances observed among nodes may originate from their underlying tissue structures or sample-specific biases (i.e., batch effect). Given the unsupervised nature of SCGP, scenarios in which biases overwhelm the biological differences between structures can lead to a situation where increasing γ will cause structures from different samples to disintegrate and thus be assigned to distinct partitions. This caveat is addressed in practice by curating a well-integrated reference set for SCGP followed by applying SCGP-Extension.

Post-processing

Upon acquiring the initial partition outcomes, optional post-processing steps can be executed to refine the results.

- (1) Size Filtering: Partitions accounting for less than 0.2% of all nodes are discarded.
- (2) Spatial smoothing: For any node who holds a different partition assignment from its spatial neighbors, we reassigned it to the partition held by the majority of its spatial neighbors (>50%) if applicable.

SCGP-extension

SCGP-Extension extends a given set of reference partitions to unseen query samples. Query samples are processed in the same manner as specified in the SCGP pipeline for graph construction. Reference partitions are defined on the reference nodes, which should be in the same format as nodes in the query samples. These partitions are usually generated through a primary SCGP run on the reference samples. It is worth noting that technically any form of discrete labels can be employed as reference partitions. [Figures S4B](#page-12-4) and S4C demonstrates experiments that extend ground truth labels or noisy labels defined by arbitrary annotation tools.

Pseudo-nodes

SCGP-Extension introduces pseudo-nodes as guidance for unseen query sample partitioning. Based on the reference nodes and their reference partitions, pseudo-nodes can be created for each partition via two strategies.

(1) Selection: A median node feature vector is calculated using all reference nodes affiliated with the partition. The Euclidean distance between each node's feature vector and the median vector is calculated. Pseudo-nodes are sampled from all reference nodes based on their distances to the median vector.

(2) Random sampling: The mean and covariance of the node feature vector are calculated using all reference nodes affiliated with the partition. Pseudo-nodes are generated by sampling multivariate normal random variables based on the mean and covariance.

In practice, we typically generate 100 pseudo-nodes for each partition, but the size can be adjusted based on the number of nodes in the reference and query samples. The two strategies tend to yield comparable results, we empirically prefer the selection strategy.

Prior to their integration into the query graph, dense feature edges are added to the pseudo-nodes – 20 nearest neighbors for each pseudo-node if in total 100 nodes per partition are used. This is to guarantee well-structured communities within the pseudo-nodes.

Reference-query edges

Pseudo-nodes are integrated into the query graph via additional reference-query edges. For each node in the query graph, its *k'* nearest neighbors in the pseudo-nodes based on Euclidean distances between node features are identified. An additional ratio parameter $r(0 < r \le 1)$ is included to regulate the strengths of reference-query edges. Given a total of *N* nodes in the query graph, the initial nearest neighbor search will yield *Nk'* reference-query edges. These edges are sorted by edge weights (i.e., Euclidean distances), with only the top rNk' edges with the highest weights or closest distances retained.

The completed query graph comprises spatial edges, intra-sample feature edges (*k*-nearest neighbors within the query nodes), and reference-query feature edges (r-downsampled *k*'-nearest neighbors between query nodes and pseudo-nodes). *r* and *k'* are two additional hyperparameters in SCGP-Extension, which control the degree of matching between query nodes and reference partitions. In practice we adjusted *k*, *k'* and *r* so that the total number of feature edges matches the number of spatial edges, with a comparable number of intra-sample feature edges and reference-query feature edges. *r* is typically set to a value between 0.1 and 0.5 based on expectations of matching level between query and reference samples. By default we set $k = 2$, $k' = 3$, and $r = 0.3$.

Graph community detection with fixed membership assignment

The same community detection strategy is used to identify partitions in the query sample. Additional arguments including edge weights and resolution parameter γ are specified in the same manner as specified in the SCGP pipeline.

Notably, as pseudo-nodes are created for reference partitions, their assignments are predetermined and fixed throughout the partition optimization process^{[70](#page-15-0)} using the is_membership_fixed argument of the leidenalg.Optimiser().optimise_partition method. As a result, query nodes that are similar to any of the existing reference partitions will be assigned to the corresponding group, while nodes distinct from the reference will be assigned to novel partitions.

Predictive modeling for partition extension

An alternative approach to extend existing partitions to unseen samples is through constructing a predictive model and applying it for inference. This is demonstrated in two experiments in [Figures 4A](#page-7-0) and [S4B](#page-12-4). In these experiments, we used manual annotations of reference samples to train gradient boosted tree classifiers, which were subsequently applied to unseen query samples.

The training dataset was constructed using the reference nodes and used manual annotations as labels. The inputs contained reference node features, as well as 1-hop aggregated node features, which were computed by averaging features from the center nodes and their immediate spatial neighbors (defined by distance thresholding). This augmentation was inspired by the UTAG and SpaGCN methods and allowed the model to have larger fields of view. The test inputs for the query nodes were formulated similarly, and the trained models were employed to infer their cluster/partition assignments.

We evaluated a range of common machine learning methods including logistic regression, linear SVR, k-nearest neighbor classifier, and random forest. Gradient boosted trees implemented via $XGBoost^{\dagger}$ yielded the best performance.

Evaluation metrics

On datasets with ground truth annotations, we assessed the performances of SCGP, SCGP-Extension and other unsupervised annotation tools using the following metrics.

- (1) Adjusted Rand Index^{[72](#page-15-2)} (ARI) and Rand Index (RI): ARI and RI are measures that evaluate the similarity between two data clusterings, in which ARI also takes into account the probabilities of random agreement between two clusterings. ARI ranges from -1 to 1, where 1 indicates perfect agreement and 0 indicates a random agreement. We adapted the implementation in scikit-learn^{[58](#page-14-16)}: sklearn.metrics.adjusted_rand_score. RI ranges from 0 to 1, where 1 indicates perfect agreement. RI is only used in the Lung IMC dataset to reproduce metrics reported in its source publication. We adapted the implementation in scikit-learn: sklearn.metrics.rand_score.
- (2) V Measure: V Measure is the harmonic mean of homogeneity and completeness, two intrinsic metrics used to evaluate the quality of clustering. Homogeneity measures if all points within one unsupervised cluster are members of a single label class, while completeness verifies that all members of a given label class are assigned to the same cluster. All three metrics range from 0 to 1, where higher values indicate better clustering performance. We adapted the implementation in scikit-learn:

sklearn.metrics.homogeneity_completeness_v_measure. Homogeneity score is used in the Lung IMC dataset to reproduce metrics reported in its source publication.

- (3) F1 Score: F1 is an accuracy measure calculated as the harmonic mean of precision and recall. For each label compartment, the F1 score is calculated through the following process:
	- \circ Labels: 1s are assigned to all nodes affiliated with the target label compartment, 0s are assigned to the rest.
	- \circ Predictions: For a given partition, predictions are calculated using the indicator function of whether a node is assigned to that specific partition. A series of predictions will be derived for all partitions identified by the unsupervised annotation tool.
	- \circ Metrics: Multiple F1s are calculated based on the labels and the series of predictions. The highest F1 score corresponds to the partition that has the best match with the target label compartment, and this score is taken as the final score.

To compare performances of different methods, we used the following statistical test.

(1) Wilcoxon signed-rank test: this test is employed to compare paired metrics (ARI, F1) calculated between ground truths and partitions defined by unsupervised annotation methods, assessing the significance of differences between groups of metrics.

In the analysis of disease states, we used the following statistical tests.

- (1) Two-sample t-test: this test is employed to determine if there is a significant difference between the means of two independent groups. It is used to compare characteristics of samples with different disease conditions.
- (2) Jonckheere-Terpstra test: this test is employed to assess ordered differences among multiple independent groups. It is used to determine if certain characteristics of tissue structures have ordered changes with respect to disease stages.

Running time and memory usage

We profiled running time and peak memory usage for all unsupervised annotation methods using two major tasks.

- (1) Joint clustering/partitioning of 17 samples from the DKD kidney dataset, containing 137,654 cells;
- (2) Joint clustering/partitioning of 4 samples from the same specimen (Br8100) from the DLPFC Visium dataset, containing 14,364 spots.

The profiling is performed on an amazon cloud service ec2 instance (r6i.16xlarge or g4dn.16xlarge if GPU is required). See [Table S2](#page-12-4) for full results.

CN, UTAG, SCGP, and SCGP-Extension had the shortest running times. KMeans-based methods including CN required minimum memory usage, while memory consumption of Leiden-based methods relied on the number of nearest neighbors in the graph construction, in which SCGP and SCGP-Extension were less demanding.

Hyperparameter optimizations

For unsupervised annotation methods benchmarked in this work, we searched the hyperparameter space for each method and chose the best results. Below we briefly described the pipelines for these methods, please refer to their source publications and code bases for full details. Note that parameters that are optimized during the hyperparameter search are marked as ''searched''.

- (1) KMeans.
	- \circ Input: Biomarker expression vectors of cells/spots;
- o Clustering: KMeans algorithm (K∈[2,10], searched), implemented by scikit-learn.^{[58](#page-14-16)}
- (2) Leiden. 39
	- o Input: *k*-nearest neighbor graph ($k \in [10, 100]$, searched) constructed using the biomarker expression vectors of cells/spots;
	- o Clustering: Leiden algorithm, implemented by leidenalg (<https://github.com/vtraag/leidenalg>). Resolution parameter is searched.
- (3) Cellular Neighborhood (CN). [13,26](#page-13-2)
- \circ Input: Cell types are first identified through leiden clustering. For each cell, a composition (frequency of cell types) vector is calculated based on a window of 10–20 nearest neighboring cells (searched) as measured by Euclidean distance between X/Y coordinates.
- ^B Clustering: KMeans algorithm (*K*∊½2;10, searched), implemented by scikit-learn.

(4) UTAG.[15](#page-13-4)

- \circ Developed and validated on spatial proteomics (mIF) data: IMC, CyCIF.
- \circ Input: For each cell/spot, an average biomarker expression vector is calculated over all neighboring cells/spots and the center cell/spot within a window surrounding the center cell/spot thresholded by Euclidean distance (10–20 μm, searched), referred to as the spatially aggregated feature matrix. *k*-nearest neighbor graph (*k*∈[10,100], searched) is then constructed using the aggregated expression vectors.
- o Clustering: Leiden algorithm, implemented by leidenalg. Resolution parameter is searched.
- o Results on the LungIMC dataset are directly downloaded from [https://zenodo.org/records/6376767.](https://zenodo.org/records/6376767)

(5) Spatial-LDA.^{[28](#page-13-15)}

- o Developed and validated on spatial proteomics data: CODEX, MIBI-TOF.
- \circ Input: Cell types are first identified through leiden clustering. For each cell, its local environment is encoded as the count of cell types (bag-of-cell) within a window surrounding the center cell thresholded by Euclidean distance (10-30 µm, searched). Spatial prior (i.e., adjacency between cells) is first constructed by computing the Voronoi partitioning of cell positions, in which pairs of cells that share a facet in the Voronoi partitioning are connected, then reduced to a minimum spanning tree based on the edges.
- o Clustering: Latent Dirichlet Allocation with spatial prior, implemented in https://github.com/calico/spatial_lda. Major parameters (i.e., difference penalty) are searched.
- (6) BayesSpace. 29
	- \circ Developed and validated on spatial transcriptomics data: Visium.
	- \circ Input: Top principal components of the log transformed and normalized gene expression counts.
	- \circ Clustering: Spots are modeled using a fully Bayesian model with a Markov random field prior, specified by the Potts model. Model parameters are estimated using a Markov chain Monte Carlo method. We adapted codes (in R) from [https://](https://edward130603.github.io/BayesSpace/articles/BayesSpace.html) edward130603.github.io/BayesSpace/articles/BayesSpace.html. Major parameters (i.e., number of clusters) are searched.
- \circ Only used on the DLPFC data.
- (7) SpaGCN. 32
	- O Developed and validated on spatial transcriptomics data: Visium, SlideSeqV2, MERFISH, STARmap.
	- \circ Input: Biomarker expression vectors of cells/spots are reduced to their top 20 principal components and then constructed into a weighted undirected graph, in which edges are weighted by Euclidean distances (histology information is not included in this study).
	- \circ Clustering: A one-layer graph convolutional network on the input graph generates initial embeddings for cells/spots, which are clustered using the Louvain algorithm (resolution searched). Network parameters and cluster centroids are optimized by minimizing a soft assignment-based loss function using stochastic gradient descent with momentum until convergence. We adapted codes from <https://github.com/jianhuupenn/SpaGCN>.
- (8) SpiceMix.^{[24](#page-13-11)}
	- \circ Developed and validated on spatial transcriptomics data: Visium, seqFISH+, STARmap.
	- \circ Input: a Hidden Markov Random Field model is constructed based on the graphical model, which treats cells/spots as nodes and connects spatially adjacent pairs (through Delaunay triangulation) with edges.
	- \circ Clustering: Cells/spots are first clustered using the Louvain algorithm to initialize estimates of hidden states and model parameters, which are further iteratively optimized via coordinate ascent. A total of 6–15 metagenes (searched) are used. We adapted codes from <https://github.com/ma-compbio/SpiceMix>.

QUANTIFICATION AND STATISTICAL ANALYSIS

Details of statistical tests are described in the figure legends and in [STAR Methods](#page-16-0) (see the subsection "evaluation metrics").

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Supplemental information

Discovery and generalization

of tissue structures from spatial omics data

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Supplementary Figures

A. Simulated data is adopted from Chidester et al.^[s1], which modeled the structure of the mouse cortex. Each simulated layer of the cortex consists of different combinations of glial cells, excitatory and inhibitory neurons (e.g., eL1, i1, g1). SCGP identified ground truth layer structures across diverse settings (i.e., cell density, number of layers). **B.** Quantitative measurements revealed that joint clustering of multiple samples derived from the same data distribution always yield better performances than independent clustering**.** Notably, thinner layers (2-3 cells) are typically more challenging to recognize. **C.** SCGP identified ground truth tissue structures in TMA data simulated using different strategies: Condition 1 consists of a base layer and two major structures; Condition 2 consists of a base layer and a much wider variety of tissue structures; Condition 3 defines segments of equal sizes and assigns them to different tissue structures randomly. Cells are sampled for each tissue structure based on a predefined cell type composition mapping. Note that both cell types and biomarker expression are directly sampled from a real head-and-neck cancer dataset^[s2] that we have previously studied. **D.** In condition 3 simulation, we explicitly add novel structures (e.g., red, purple partitions) to the query samples. SCGP-Extension successfully recognized both known and novel tissue structures across all query samples, achieving comparable or superior accuracy to the reference samples. **E.** SCGP and SCGP-Extension achieved robust performances (ARI > 0.7) across diverse simulated TMA data regardless of shapes and complexities (i.e., 10+ tissue structures).

A. Clustering/partitioning outputs from unsupervised annotation tools on four samples of different DKD classes are illustrated. Qualitatively, Leiden clustering identified cell types that are not spatially smooth; CN defined an extra cluster (cyan) for proximal tubules; UTAG defined multiple smaller clusters (light green, gray, purple) that do not correspond to any compartments; Spatial LDA and SpiceMix had more noisy annotations for glomeruli (orange); SpaGCN misrecognized some regions as blood vessels; SCGP misrecognized some fibrotic glomeruli as blood vessels. **B-C.** Additional quantitative metrics (V measures and Rand index) show consistent results. *P* values are calculated using the Wilcoxon signed-rank test. **D.** Full heatmap for all biomarkers tested in the DKD Kidney dataset shows signature protein biomarkers for SCGP partitions, with each partition corresponding to a manually annotated compartment.

Figure S3. Ablation experiments on the DKD Kidney dataset, related to Figure 2.

A. SCGP established a hybrid graph consisting of spatial edges and feature edges. Excluding either component will cause major performance decay: Excluding spatial edges degrades SCGP to a vanilla Leiden clustering, resulting in noisy and unsmooth partitions; Excluding feature edges disconnects distant tissue structures and leads to inconsistent outputs. Switching the type of spatial edges does not significantly change outputs. **B.** ARIs of different variants show consistent results as observed in panel A. **C-D.** To evaluate how segmentation noise might affect partition outcome, we experimented with randomly excluding nodes/cells from the graph. Outcomes and performances are robust and stable even after 20% dropout. **E-F.** We examined the influence of a hyperparameter choice: num of feature edges. More feature edges (>5) resulted in fragmented partitions and worse alignment. Optimal performances were observed when the number of edges is set between 2 and 4 in this experiment. **G.** Granularity of the partition outcomes can be controlled by the resolution parameter.

Figure S4. Partitioning of DLPFC samples, related to Figure 3.

A. Joint unsupervised annotations recognized major layers in four DLPFC samples. Notably, characterization of the two thinner layers (Layer 2 and Layer 4) were worse in most methods. **B.** Approaches utilizing partial ground truth annotations were examined.(continued)

(continued) The predictive modeling approach trained a gradient boosted tree model (XGBoost) with the ground truth annotations and applied it to the rest. SCGP-Extension can be adapted with ground truth annotations as references. The Label₂-Extension variant demonstrated the best performances. **C.** SCGP-Extension can be applied with noisy references: when using UTAG annotations (fitted on Reg 151673) as references, resulting query partitions did not show better alignment with ground truth, but captured the same spatial compartmentalization patterns as the reference region. **D.** Quantitative metrics show performances of unsupervised annotation tools. Methods utilizing partial ground truth achieved best performances, in which extensions of ground truth labels performed better than XGBoost predictions. **E.** Additional quantitative metrics (V measures and F1 scores) were calculated for independent partitioning of all samples in DLPFC and show consistent results.

Figure S5. SCGP annotations of mouse brain sections, related to Figure 3.

A. Cell types were annotated for a coronal mouse brain section. Please refer to Zhang et al.[s3] for color details. **B.** Cells were further mapped to Allen CCFv3 to annotate brain parcels. **C.** SCGP accurately identified major parcels, achieving an ARI score of 0.751. **D.** Fine-grained SCGP identified more fine-grained structures. Note the layered structures in isocortex, which can be robustly extended to the query section (panel H). **E-H.** Cells in another distant section were annotated and partitioned using the reference section (panel C-D) and SCGP-Extension, which yielded consistent results, both visually and quantitatively, with an ARI score of 0.730. **I.** SCGP with higher granularity identified fine-grained structures in the hippocampal region.

Figure S6. SCGP annotations of TR Kidney dataset and UCSF Derm dataset, related to Figure 4.

A. Primary SCGP experiment on 10 samples from the TR Kidney dataset with heavy inflammation defined six major partitions corresponding to kidney tissue structures including tubules, glomeruli, and blood vessels. The red partition exhibits high expression of immune cell biomarkers, suggesting substantial immune cell infiltration. **B.** Partitions were extended to 5 normal samples from the TR Kidney dataset that have minimal inflammation. Note the absence of the immune aggregates partition and the denser arrangement of tubules and glomeruli. **C.** Primary SCGP annotations on 17 samples from experiment 1 of the UCSF Derm dataset defined major tissue structures including the epidermis layers, dermis layer, immune aggregates, and glands. **D**-**F.** Partitions were extended to samples from different experiments of the UCSF Derm dataset, yielding consistent annotations. **G.** Signature protein biomarker expression of the six kidney tissue structures are visualized in the heatmap. Note that the same set of kidney tissue structures as in Fig. 2D were derived despite the fact that TR Kidney dataset used a different biomarker panel. **H.** Signature protein biomarker expression for the major skin partitions identified by SCGP are visualized in the heatmap.

Figure S7. Annotating representative samples from TR Kidney dataset and UCSF Derm dataset using variants of SCGP, related to Figure 4.

On a set of representative samples from the TR kidney dataset, A. UTAG defined partitions do not correspond well to tissue structures. **B.** SCGP failed to assign tissue structures from different conditions into the same partition. Note the different colors. **C.** Individual SCGP partitions of different samples were matched post hoc to reflect shared tissue structures. Note the vertical separation of partitions due to uneven background. **D.** SCGP-Extension enabled consistent recognition of tissue structures regardless of artifact and different disease conditions. **On a set of representative samples from the UCSF derm dataset, E-F.** Joint partitioning with UTAG and SCGP failed to assign tissue structures from different experiments into the same partition. Note the different colors of epidermal layers in the samples. **G.** Individual partitions of different samples were matched post hoc to reflect shared tissue structures. **H.** Reference-query extension pipeline enabled consistent recognition of tissue structures across samples from multiple experiments with different skin conditions.

Figure S8. SCGP annotations of Lung IMC dataset, related to Figure 4.

A. Clustering/partitioning outputs from various unsupervised annotation tools on representative lung samples are visualized. When jointly partitioning all 26 samples, one class of tissue structures from different samples is usually assigned to disconnected partitions, as evidenced by the diverse colors. SCGP-Extension resolved the issue by extending reference partitions defined on a well-integrated subset of seven samples. Resulting partitions on the remaining samples demonstrated the best visual alignment with manual annotations. **B-C.** Quantitative metrics^[s4] (homogeneity score and rand index) show that SCGP-Extension achieved the best performance. *P* values are calculated using the Wilcoxon signed-rank test. **D.** Signature protein biomarker expression for the tissue structures identified by SCGP-Extension are visualized in the heatmap, corresponding to the six manually annotated compartments. Note that submucosal glands and cartilage are two novel tissue structures revealed in the query samples by SCGP-Extension

Figure S9. Identification of tumor/stroma regions in tumor microenvironments, related to Figure 6.

A. Tumor and stroma regions in the representative samples from UPMC-HNC were annotated by SCGP. **B.** Structures in the representative samples from the Stanford-PC dataset were annotated by SCGP. 148 tumor cores from this dataset were divided into two subsets: 63 and 85 cores respectively as reference and query sets for validating SCGP-Extension. Note the blue partition that indicates a tertiary lymphoid structure. **C.** Quantitative metrics (ARI, F1 scores, V measures) were calculated for all unsupervised annotation methods, in which SCGP/SCGP-Extension achieved superior performances. *P* values are calculated using the Wilcoxon signed-rank test.

Figure S10. SCGP annotations of mIF images as spatial grids of patches, related to STAR Methods.

A. In the absence of cell segmentation, mIF images can be directly partitioned using SCGP by adopting a spatial-grid approach: Images were first dissected into small patches through a sliding window. Nodes were defined based on patches, with node features summarizing biomarker expression in the patches. Spatial edges and feature edges were constructed in a similar fashion to enable application of SCGP. Note that the resolution of the grid can be adjusted by varying the size of the patches. **B.** SCGP on the patch grids of three mIF images yielded similar results as cell-based SCGP, recognizing tubules, glomeruli, blood vessels, and immune aggregates in the tissue. In the severe DKD (IIB) sample, the fibrotic glomeruli were recognized as combinations of the basement membrane and normal glomeruli partitions, suggesting the degradation of normal glomerular structures into scar tissues. **C.** Signature protein biomarker expressions are visualized in the heatmap for the partitions identified by patch-based SCGP. Note the distinct immune aggregates partition enriched in CD45, CD68, and DAPI (i.e., higher cell density).

Supplementary Tables

Table S1. Details of datasets: number of samples/cells/patients, major grouping or patient characteristics, related to STAR Methods.

Table S2. Running time and memory usage of unsupervised annotation methods, related to STAR Methods.

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