



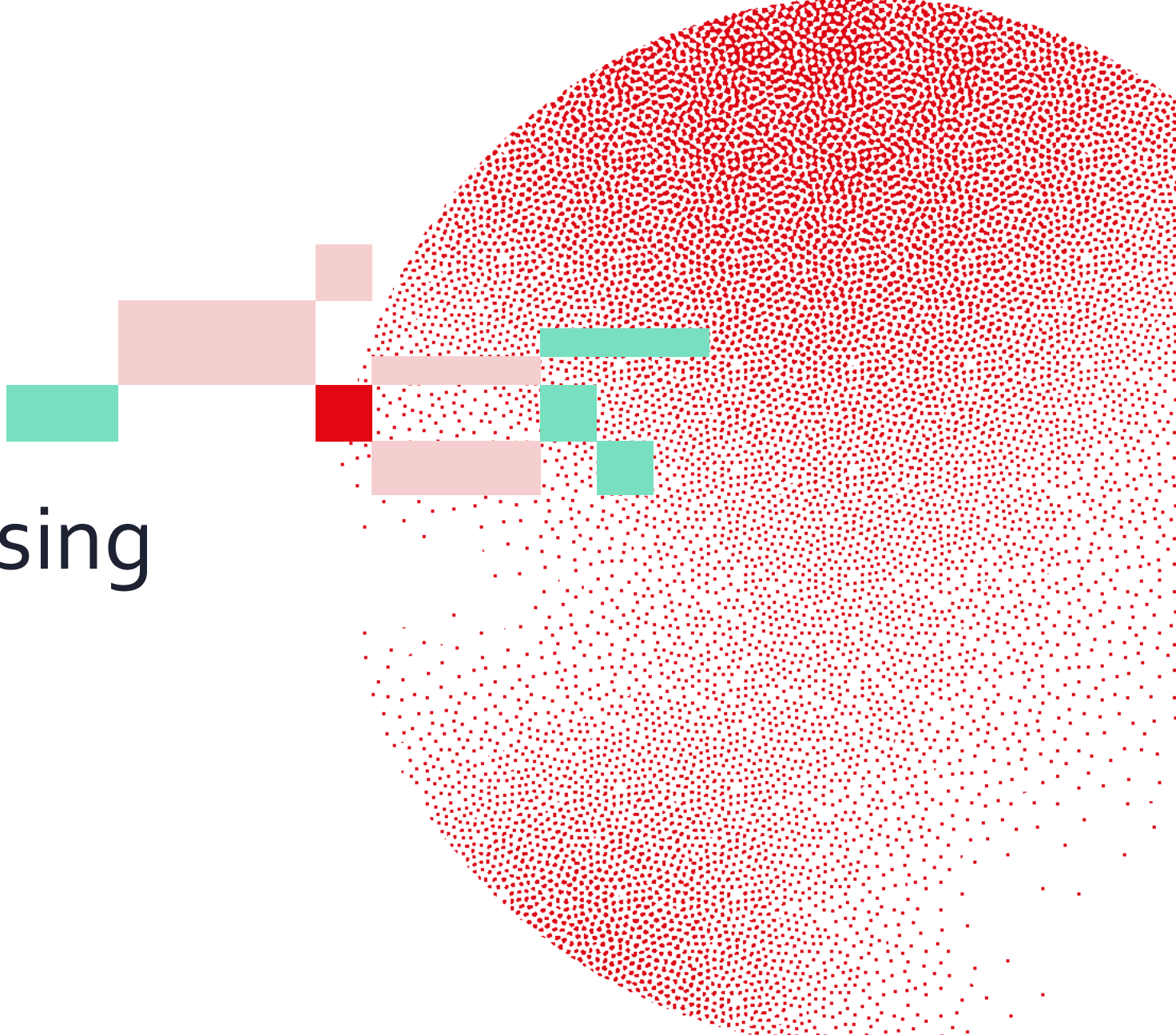
Swiss Institute of  
Bioinformatics

INTRODUCTION TO SEQUENCING-BASED  
TRANSCRIPTOMICS DATA ANALYSIS

# Intermediate processing

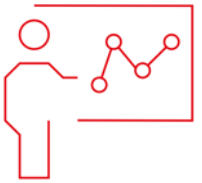
Joana Carlevaro Fita

December 9-10, 2025





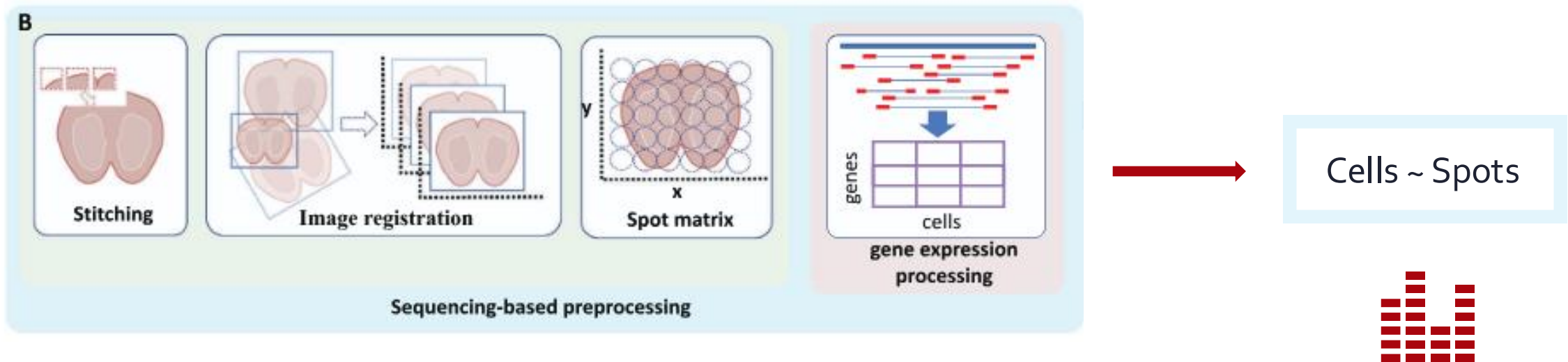
# Learning objectives



- » Describe intermediate processing steps applied to spatial transcriptomics data analysis
- » List spatially-aware methods used at each step, and explain the importance of testing them over standard scRNAseq methods
- » Explain what a spatial domain is, and key aspects in their identification

# Normalisation, feature selection, dimensionality reduction

Several scRNAseq methods are used for intermediate processing of spot-based ST



# Standard scRNAseq methods applied

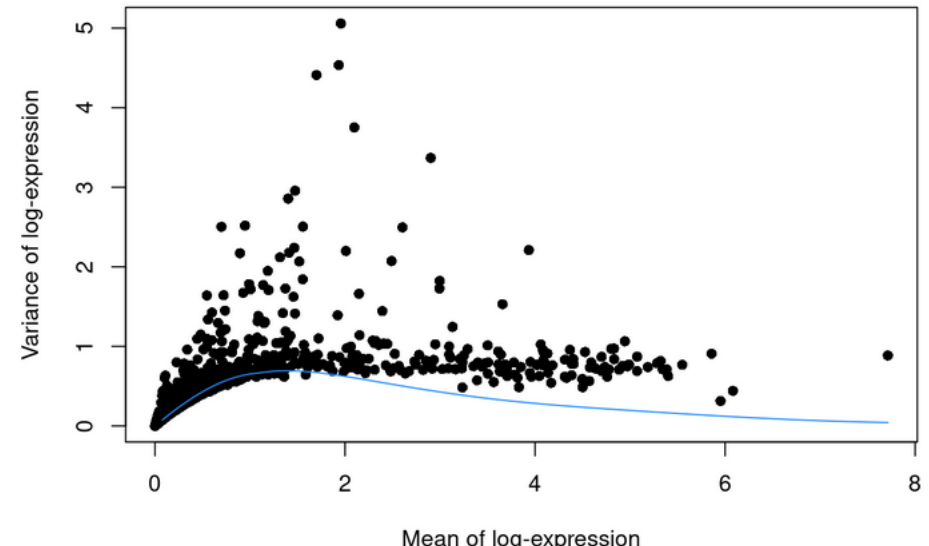
## 1. Select Features:

Identify highly variable genes (HVG) to screen a proportion of genes

- Reduce noise from random variation from biologically uninformative genes
- Improve computational efficiency

Two-step procedure:

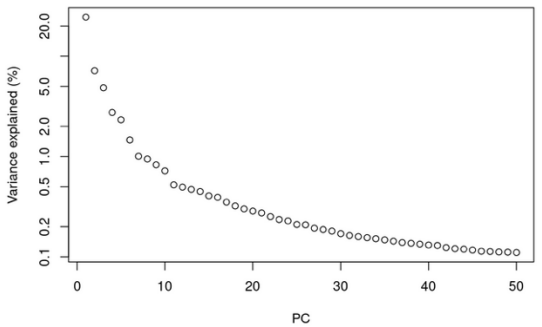
- 1) Model the mean-variance relationship, which decomposes variance into a technical component (smooth fit) and biological component (deviation thereof).
- 2) Select top HVG (fixed number or proportion of genes)



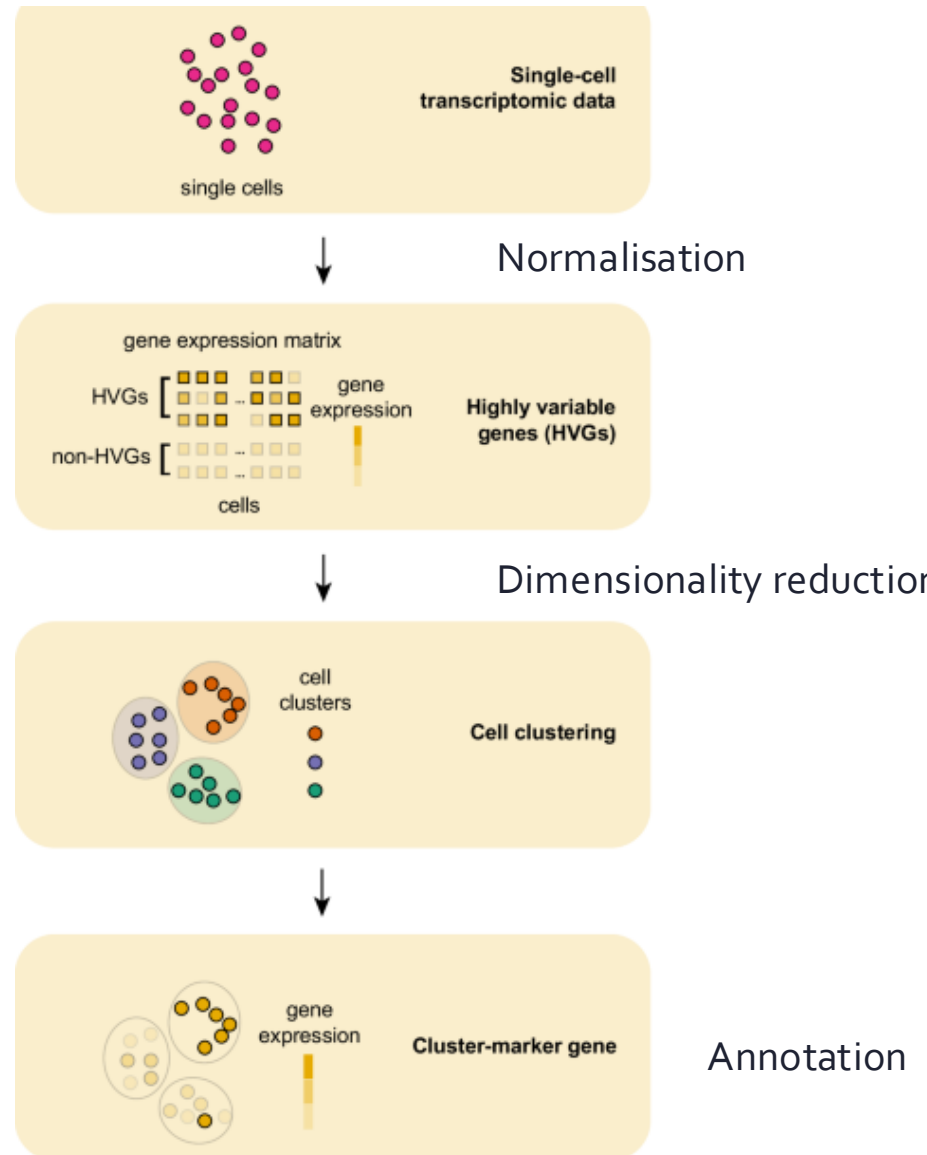
# Standard scRNAseq methods applied

## 2. PCA

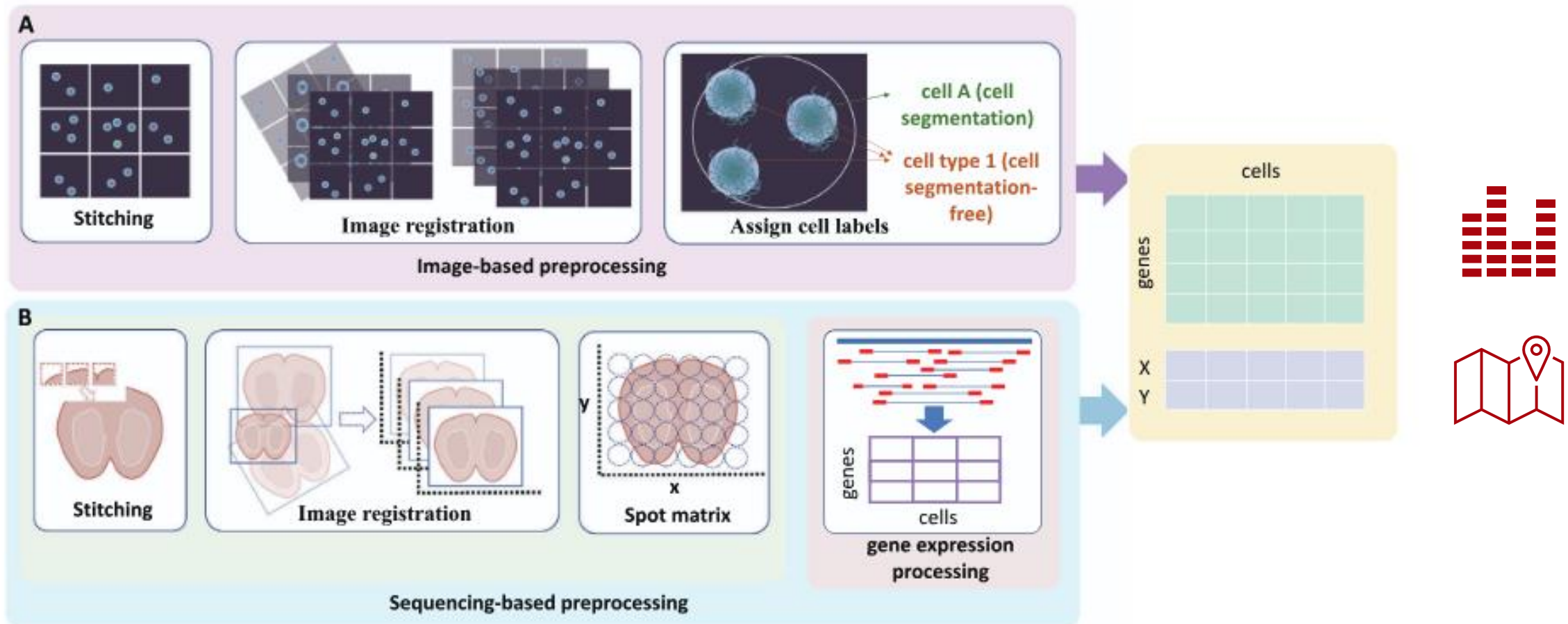
PCA to the set of top HVG  
Retain informative PCs



# Standard scRNAseq methods applied



# Spatial information



# Feature selection



## HVG: Highly variable genes

Defined only based on molecular features (gene expression).

Do not incorporate spatial information



## SVG: Spatially variable genes

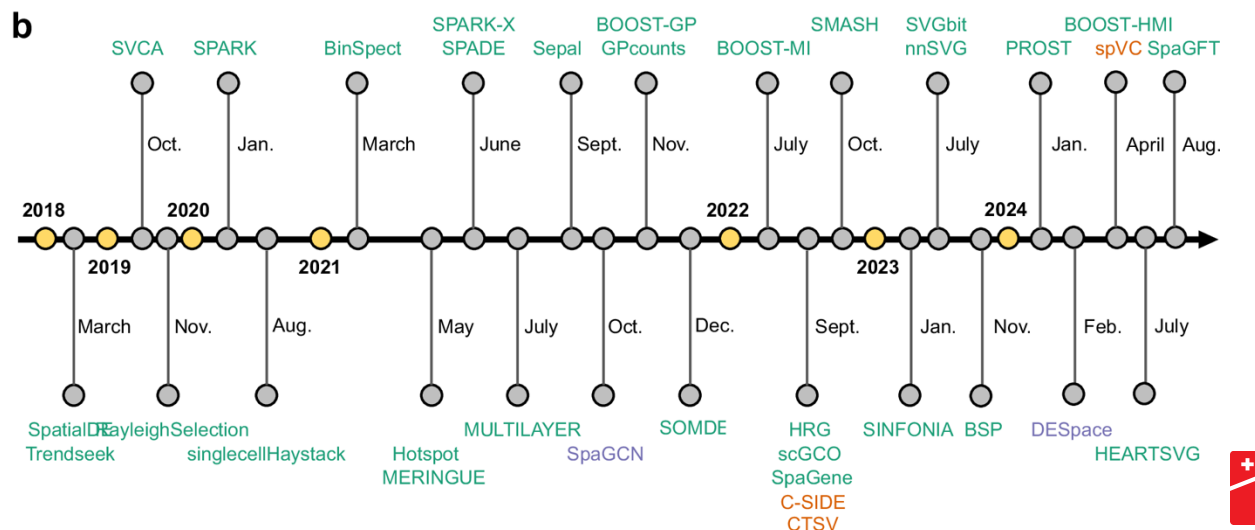
Take spatial coordinates of the measurements also into account

Select genes with non-random, informative spatial patterns → more biologically informative ranking of genes

Used either instead of or complementary to HVGs in subsequent steps.

Several SVG methods:

- de novo (nnSVG)
- pre-computed spatial clusters (by morphology or clustering methods (DESpace))

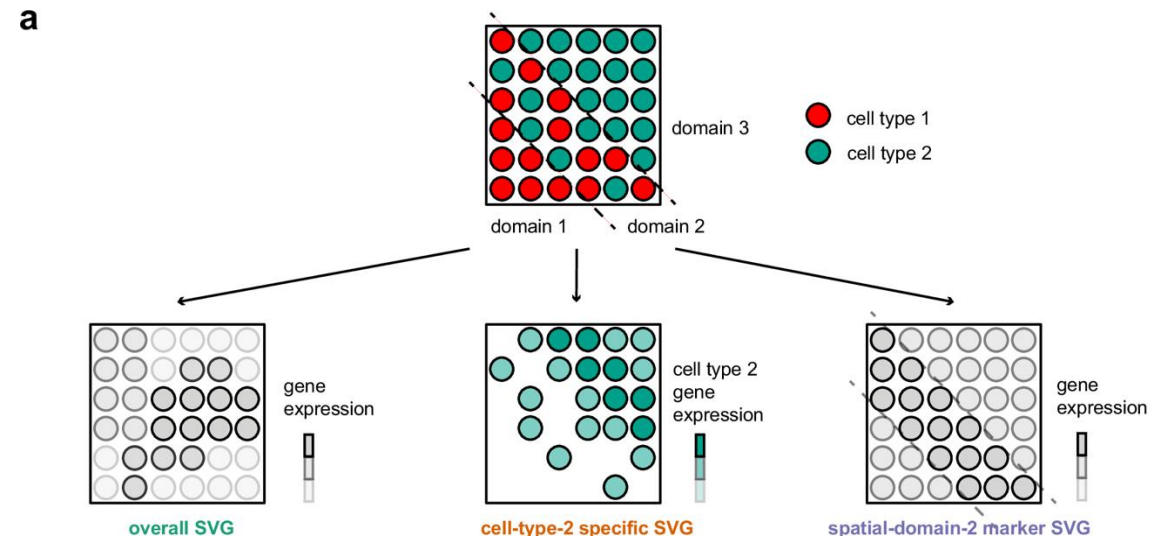




# SVG methods

## Classification of SVG methods

[Yan, Hua, and Li. Nat Comm, 2025](#) → Categorise 34 SVG methods in 3 categories:



### 1. Overall SVG:

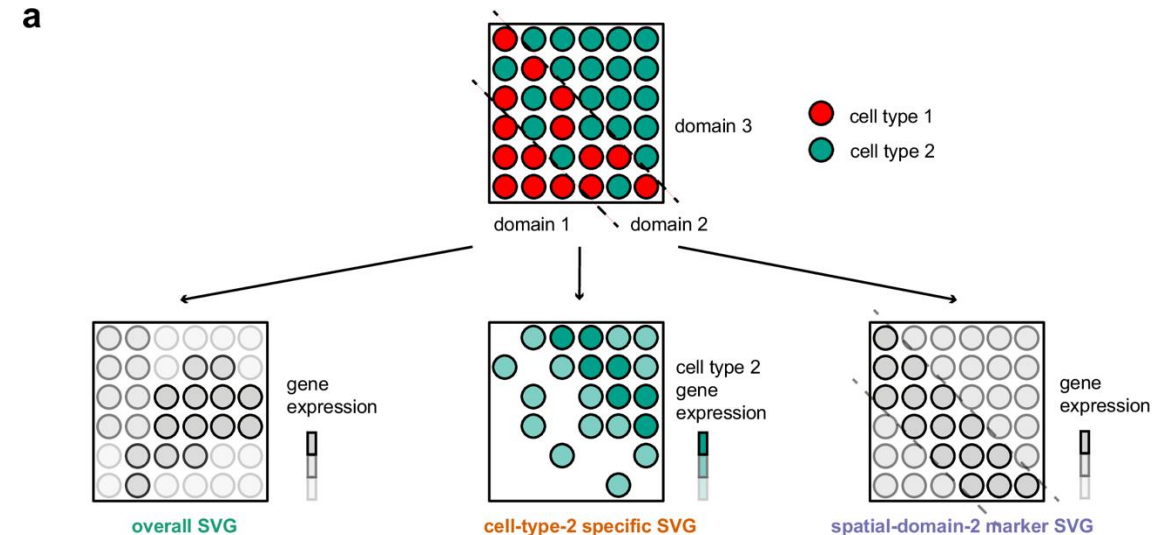
- Based on Gaussian process model: spatialDE, nnSVG
- Rank genes based on spatial autorcorrelation: Moran's I, Geary's C
- Non-parametric test of covariance matrices: SPARK
- Used as feature selection step (screen for informative genes)
- To identify spatial domains. Cluster spots (graph-based clustering) using SVGs expression and location information.
- Not necessary for all spatial domain detection methods (BayesSpace based on HVG)
- Identification of spatial-gene modules



# SVG methods

Classification of SVG methods

[Yan, Hua, and Li. Nat Comm, 2025](#) → Categorise 34 SVG methods in 3 categories:



## 2. Spatial domain-specific:

DESpace, SpaGCN

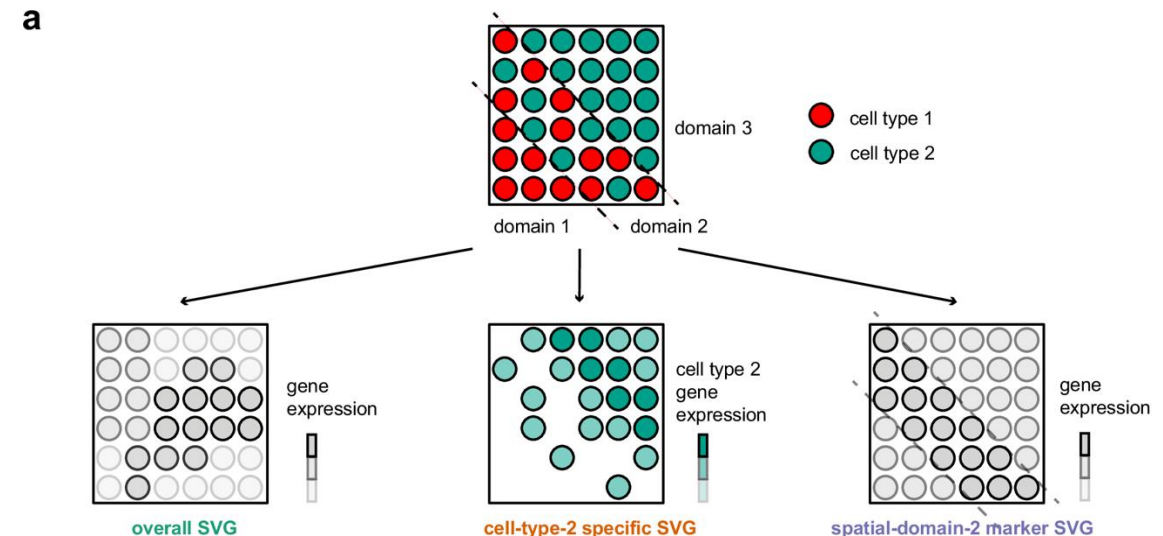
- Genes that change significantly between domains and summarise spatial information
- Identified using spatial domains
- Insights to molecular mechanisms of spatial domains/tissue layers
- Spatial domain marker SVGs can help with other domains annotation.



# SVG methods

Classification of SVG methods

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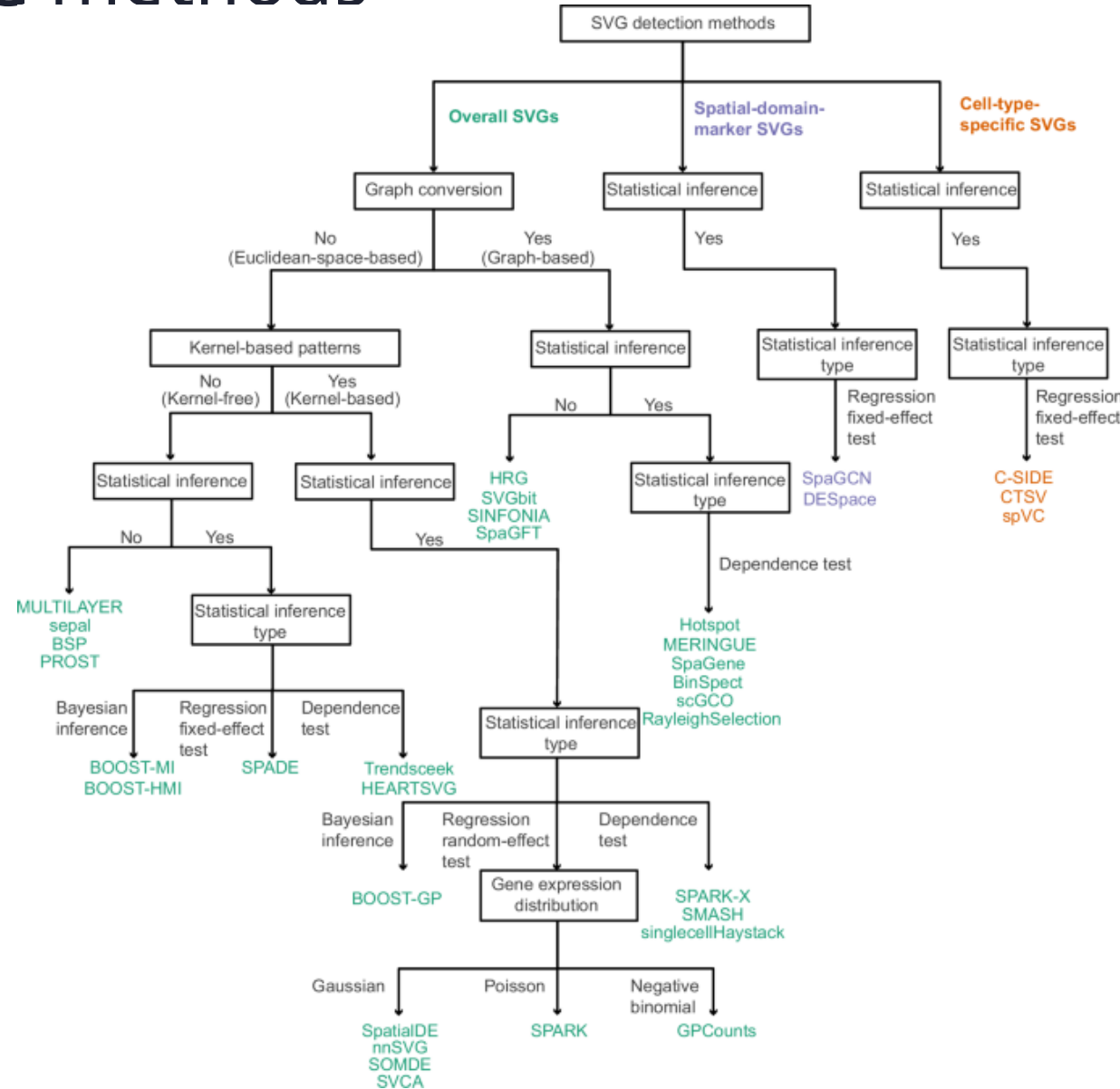
## 3. Cell type-specific:

CTSV, C-SIDE, spVC

- Genes that exhibit non-random spatial expression patterns within a cell type.
- Identified using external cell type annotations
- Identify cell subpopulations or cell states across the tissue section



# SVG methods

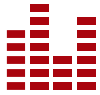


Supplementary Table 1: Summary of three existing benchmark studies for SVG detection methods

	Charitakis et al., 2023 [1]	Chen et al., 2024 [2]	Li et al., 2023 [3]
SVG Detection Methods	1. SpatialDE 2. SPARK-X 3. SOMDE 4. SpaGCN 5. SpaGFT 6. Sepal 7. SpatiaIDE 8. SpatialDE2 9. SPARK 10. SPARK-X 11. BOOST-GP 12. GPcounts 13. nnSVG 14. SOMDE	1. SpatialDE 2. SPARK-X 3. SOMDE 4. Seurat ("markvariogram") 5. SpaGCN 6. MERINGUE 7. Seurat ("moransi")	1. Moran's I 2. Spanve 3. scGCO 4. SpaGCN 5. SpaGFT 6. Sepal 7. SpatiaIDE 8. SpatialDE2 9. SPARK 10. SPARK-X 11. BOOST-GP 12. GPcounts 13. nnSVG 14. SOMDE

[...] Datasets, Evaluation Metrics, Conclusions

# Dimensionality reduction (DR)



- Non spatially-aware

DR based on the cell's molecular profile only: PCA, NMF, LDA..

Often combined with spatially-aware clustering methods



- Spatially-aware

Takes spatial information also into account: BANKSY, SpatialPCA, STAMP

Often combined with standard clustering approaches from scRNAseq

(SNN graph), scRNAseq trajectory inference models



# Dimensionality reduction (DR)

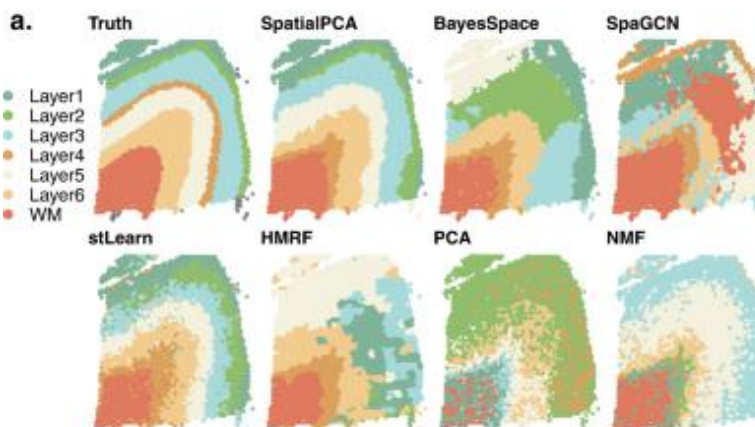
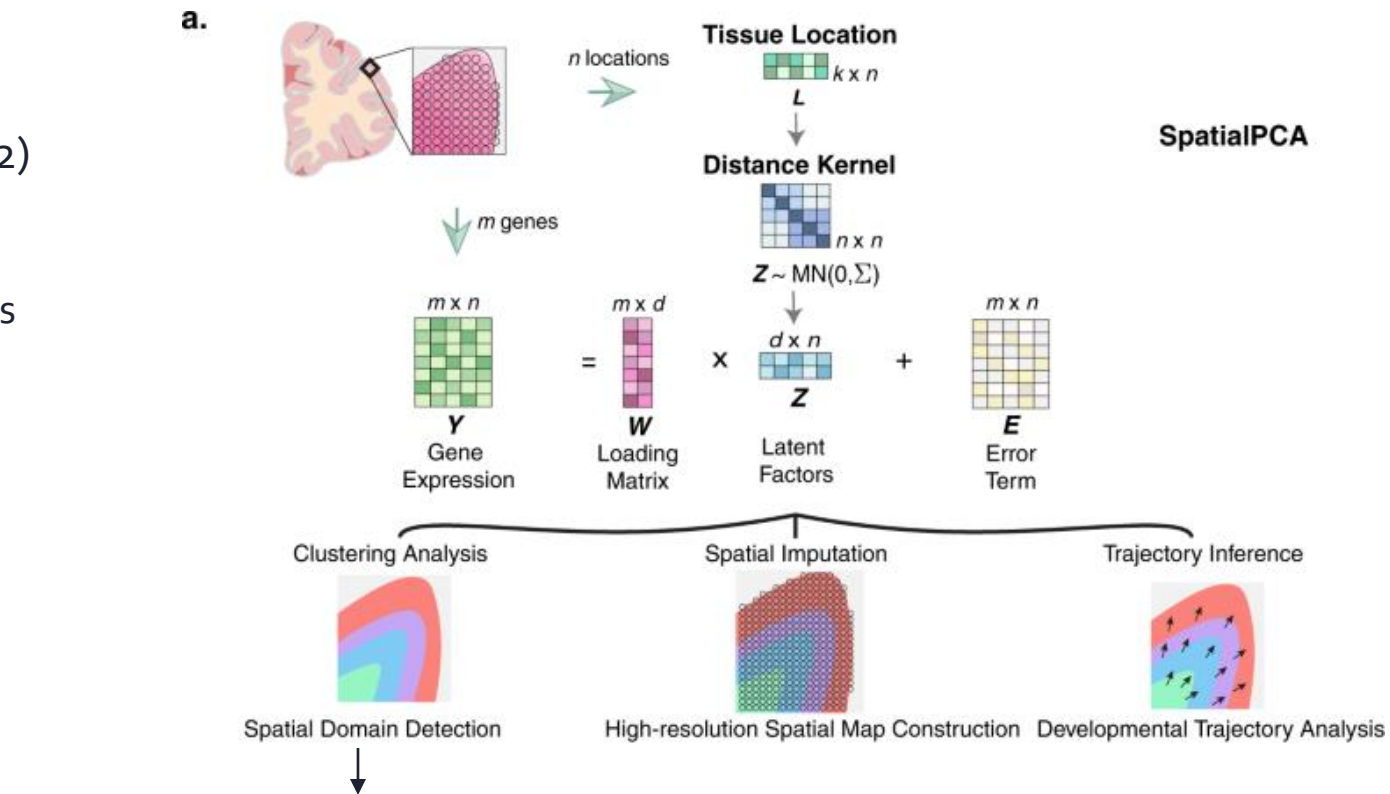
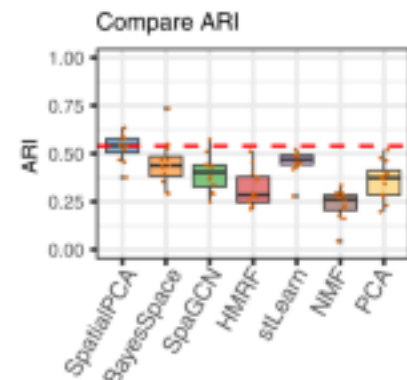
## SpatialPCA (Shang and Zhou, Nat Comm. 2022)

Neighbouring regions share similar cell type compositions

"SpatialPCA builds upon the probabilistic version of PCA, incorporates localization information as additional input, and uses a kernel matrix to explicitly model the spatial correlation structure across tissue locations."

Spatial PCs can be paired with clustering analysis on the low-dimensional components that contain spatial correlation information.

Adjusted Rand Index (ARI)  
Statistical measure to quantify  
similarity between clusterings

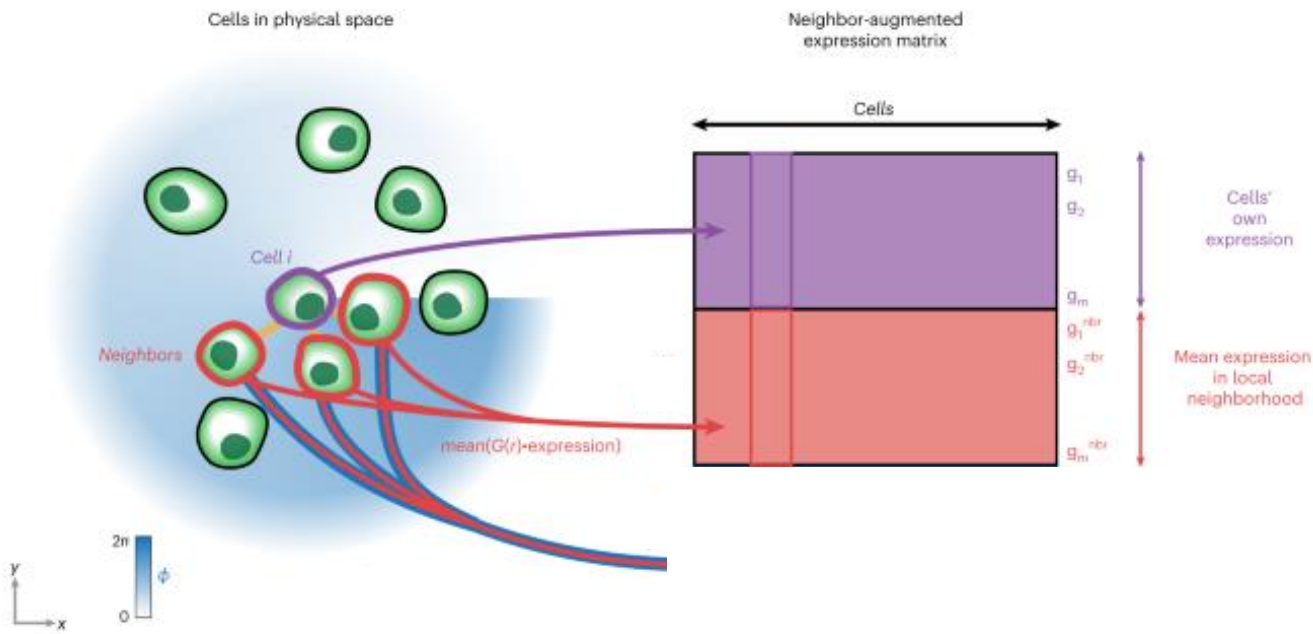




# Dimensionality reduction (DR)

## Building Aggregates with a Neighborhood Kernel and Spatial Yardstick (BANKSY)

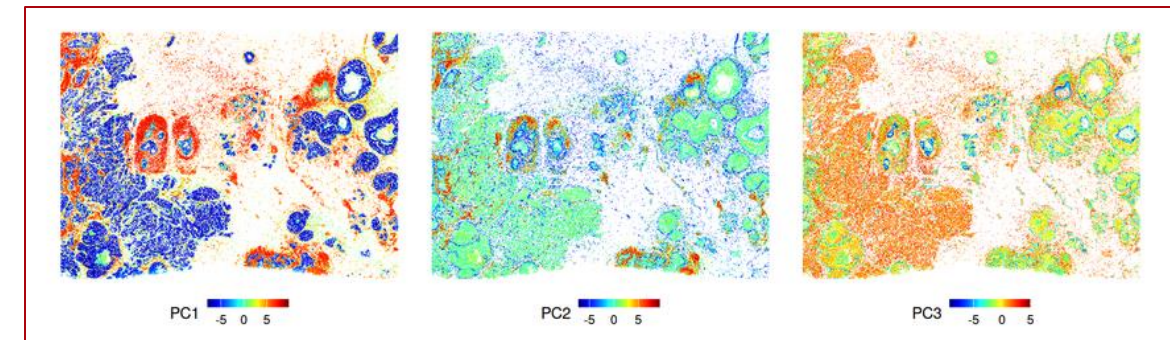
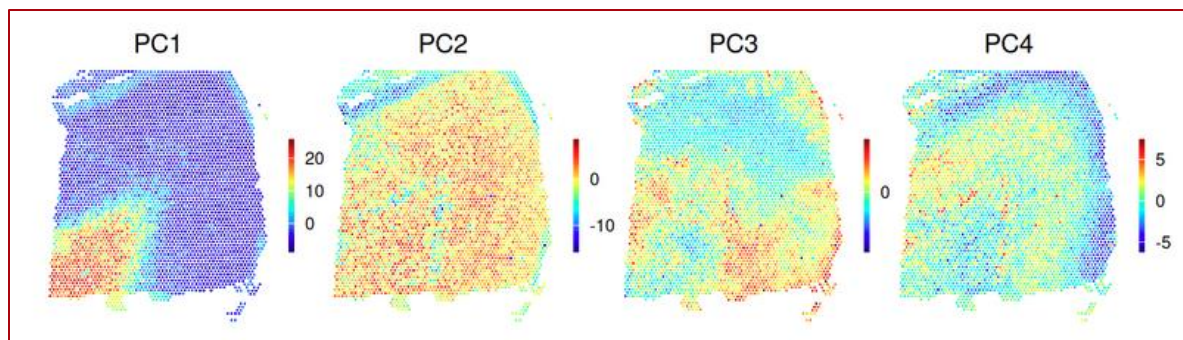
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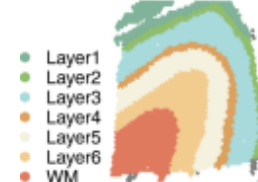
# Dimensionality reduction (DR)

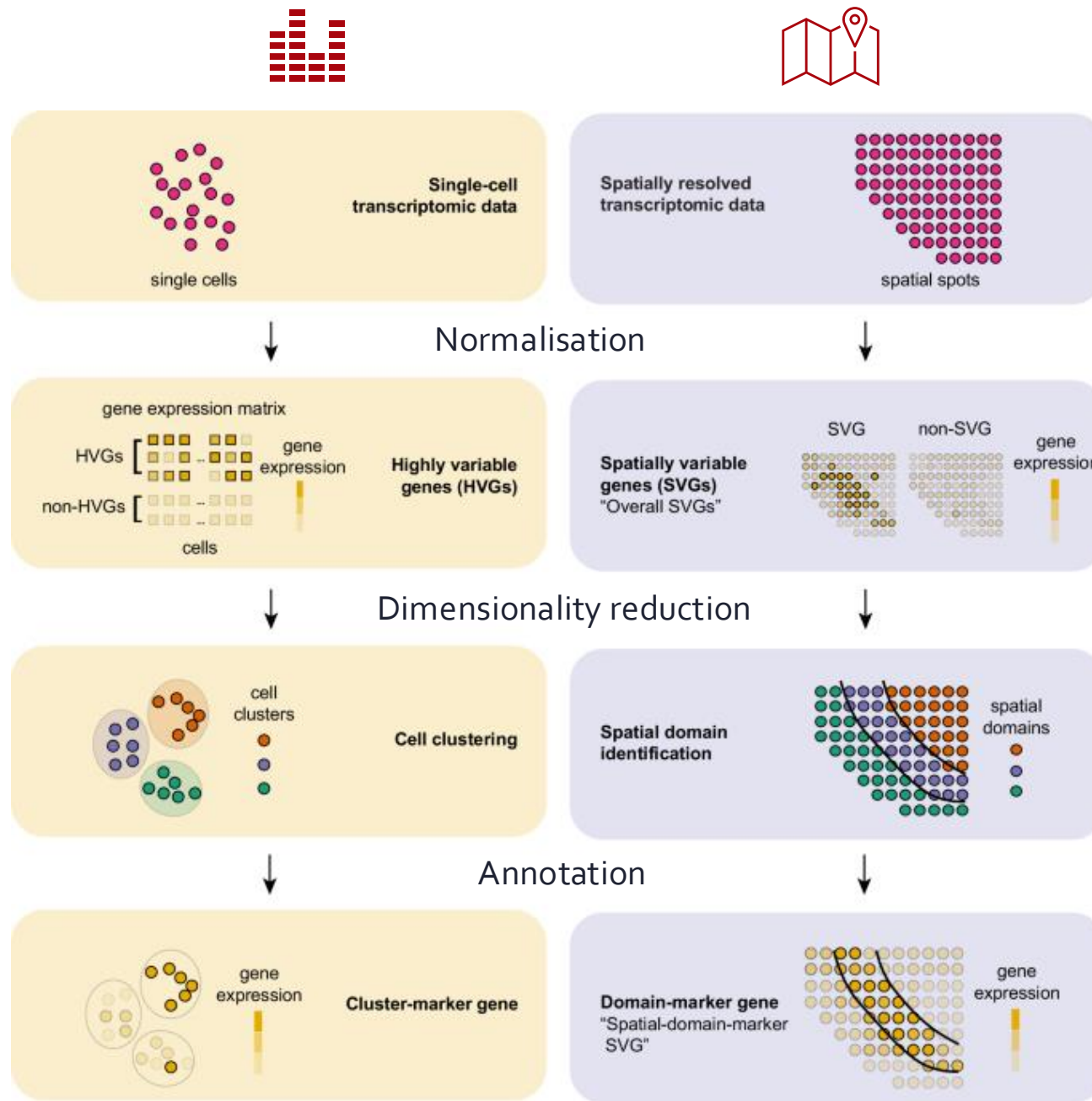
## Visualise PCs in spatial context

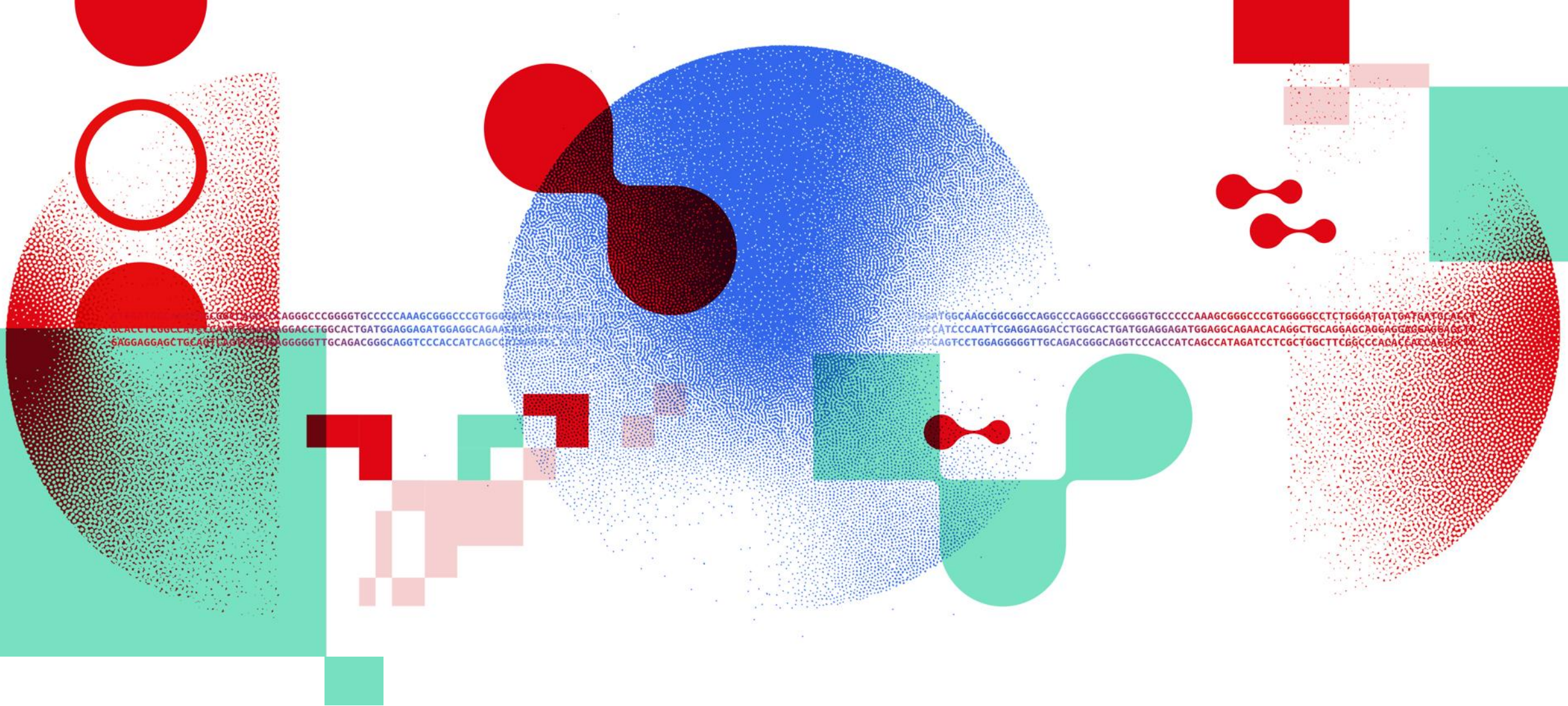
Visualise PCs in the tissue slice in order to observe if the main sources of variation explain distinguishable tissue structures/spatial locations



a. Truth







# Thank you

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