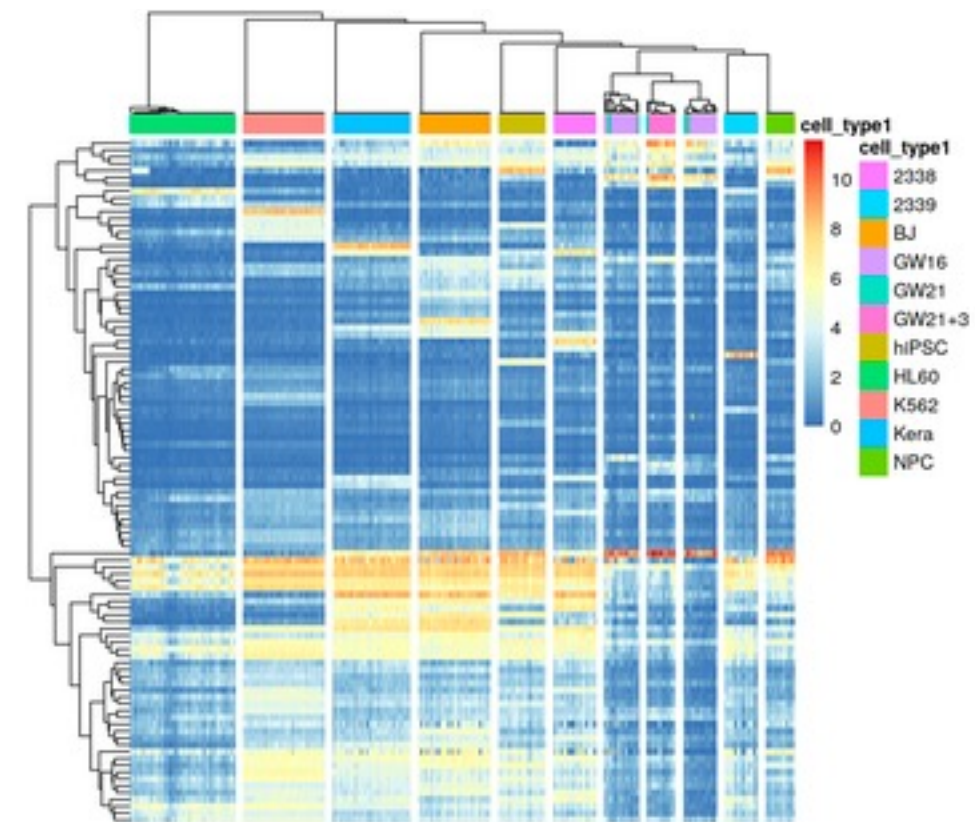
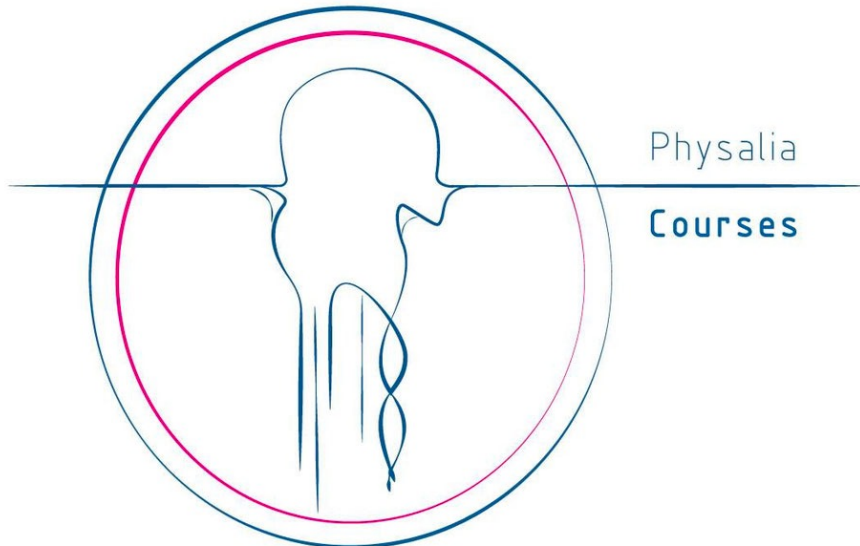
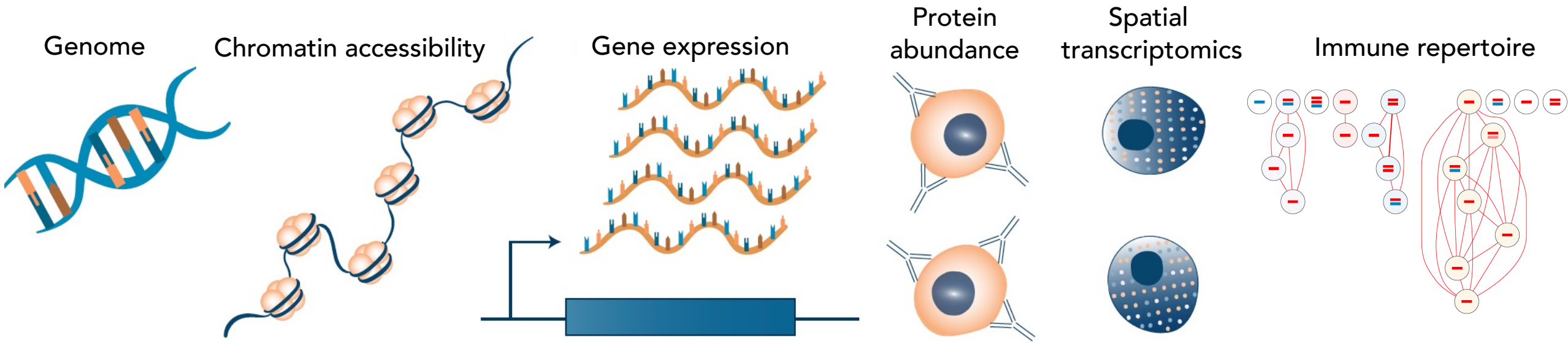


# Analysis of single-cell ATAC-seq data

Orr Ashenberg, Jacques Serizay, Fabricio Almeida-Silva  
November 2025

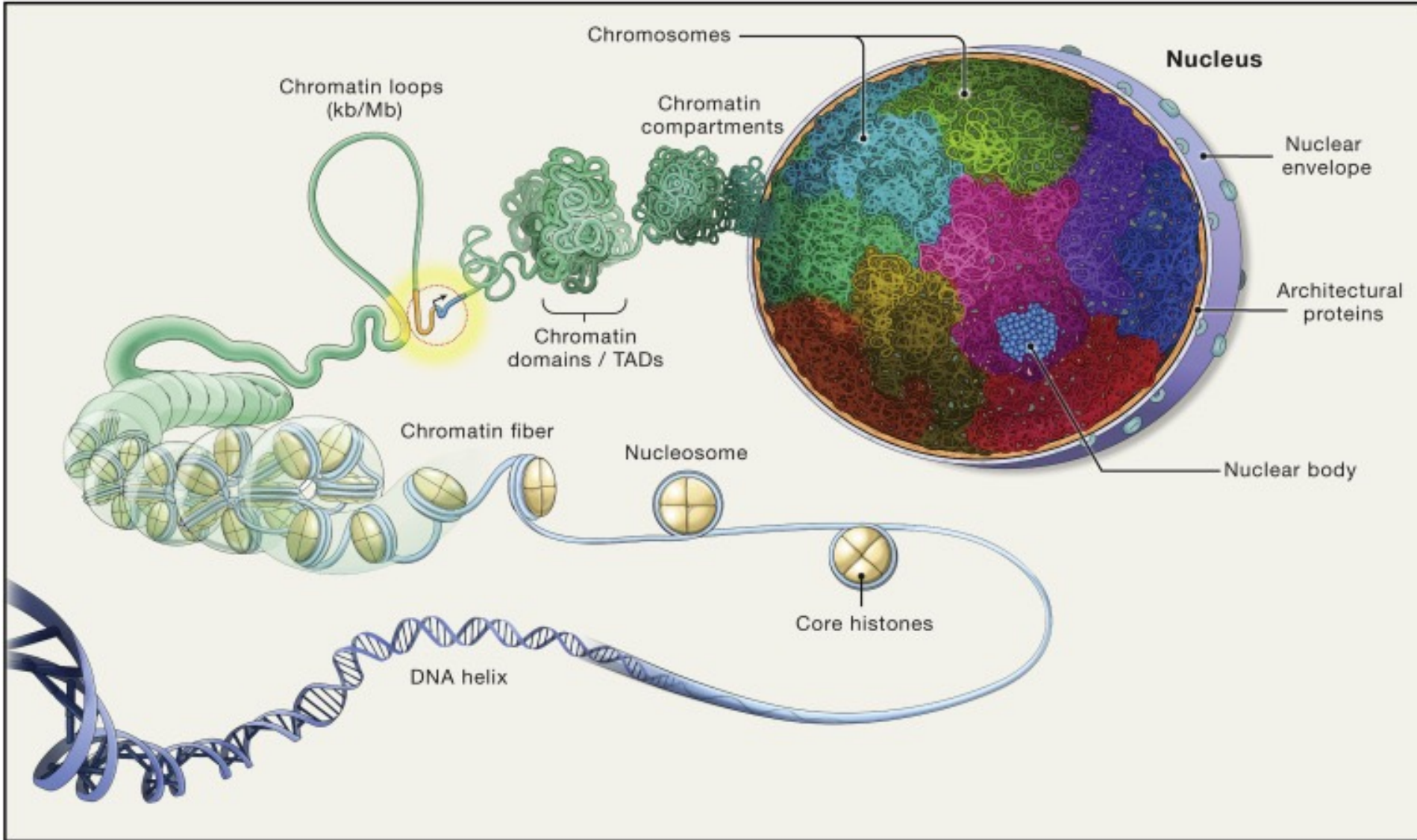


# Multimodal measurements



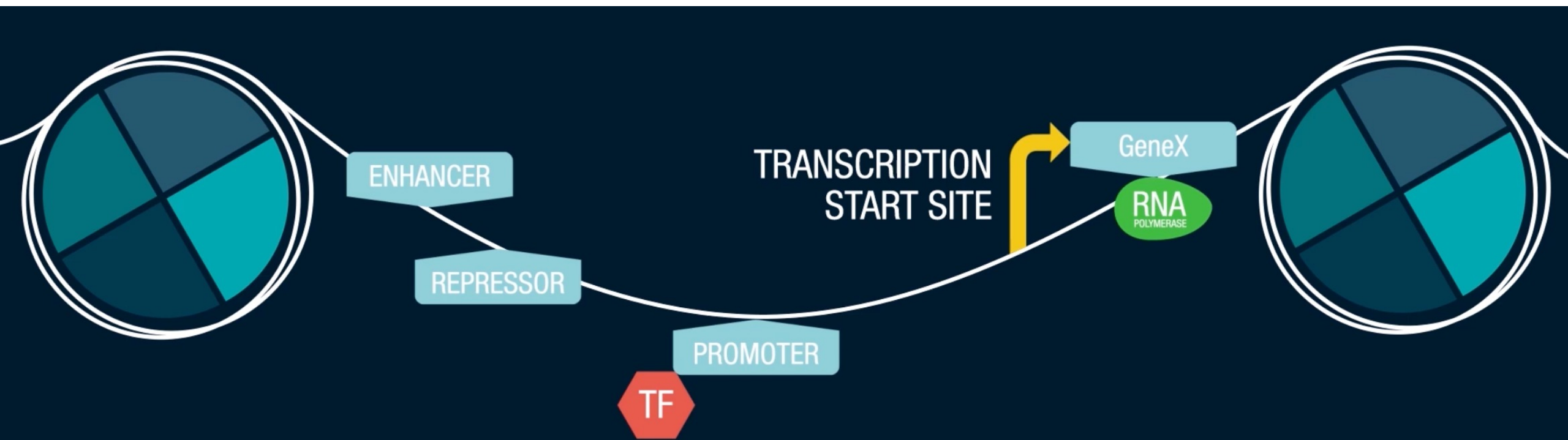
*Combining single-cell transcriptomic measurements with other data modalities can reveal gene function and gene regulation.*

# 3D organization of DNA



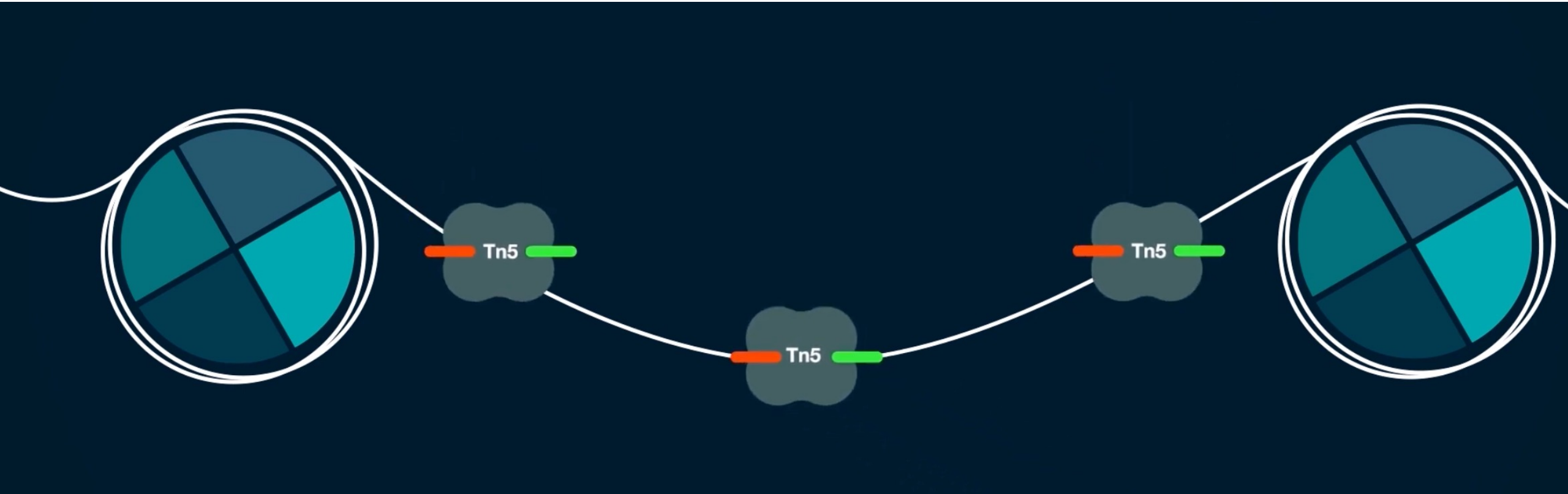
# ATAC-Seq detects accessible chromatin regions

In the cell nucleus, the chromosomes contain tightly packed chromatin material. Part of the chromatin is open and accessible to many regulatory factors who control the expression and suppression of a variety of genes.



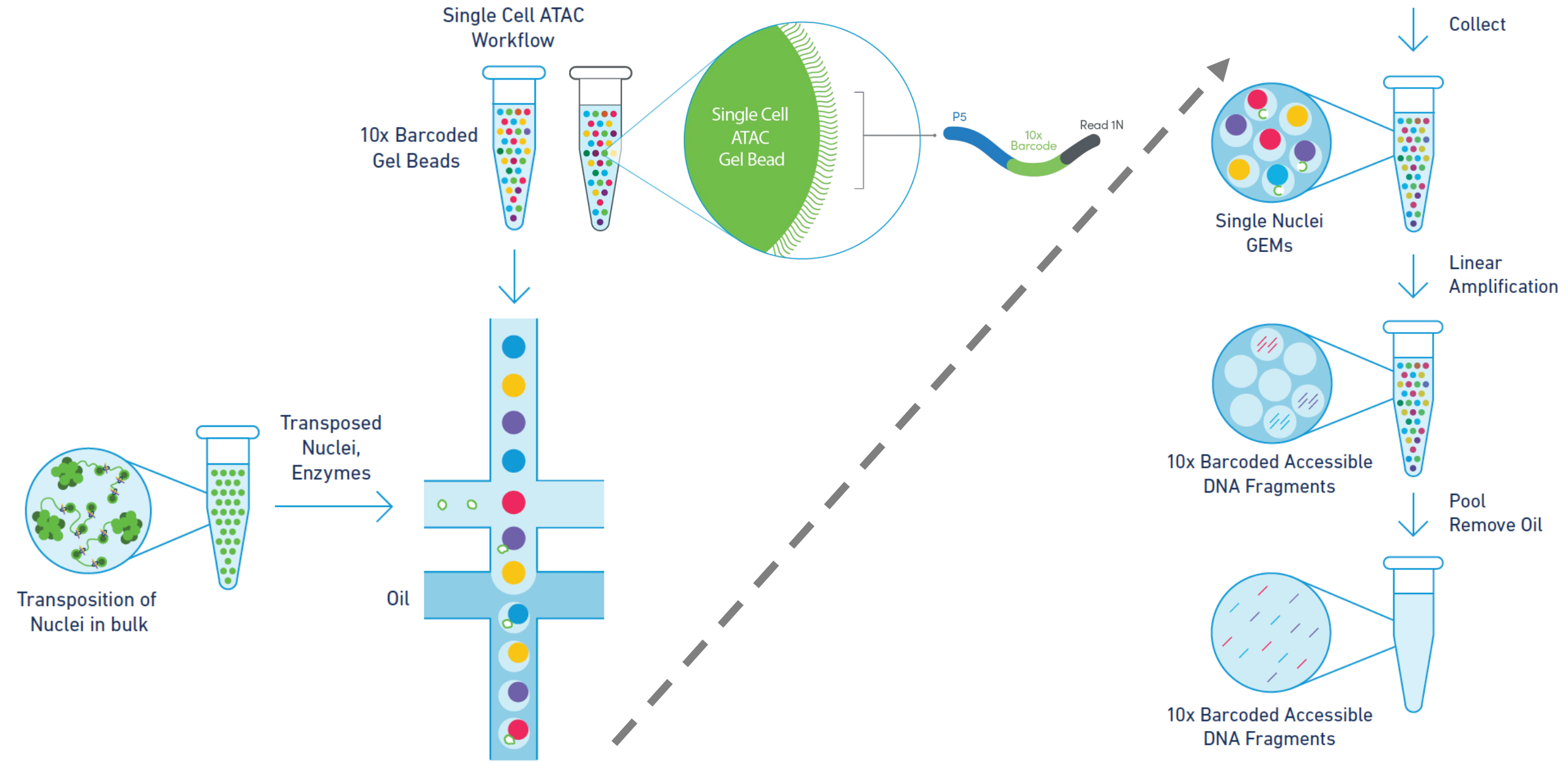
# ATAC-Seq detects accessible chromatin regions

ATACseq (as well as scATACseq) measures how open this piece of DNA is. This openness is a proxy of how easily a transcription factor can bind these parts of the genome. ATACseq measures by using an enzyme called Tn5 transposase which binds open chromatin and inserts DNA sequencing adapters.



The Tn5 transposase ideally cuts DNA just once between the neighboring nucleosomes.

# Chromium Single Cell ATAC-Seq (10x)



# Single cell resolution reveals cell-type specific regulatory elements

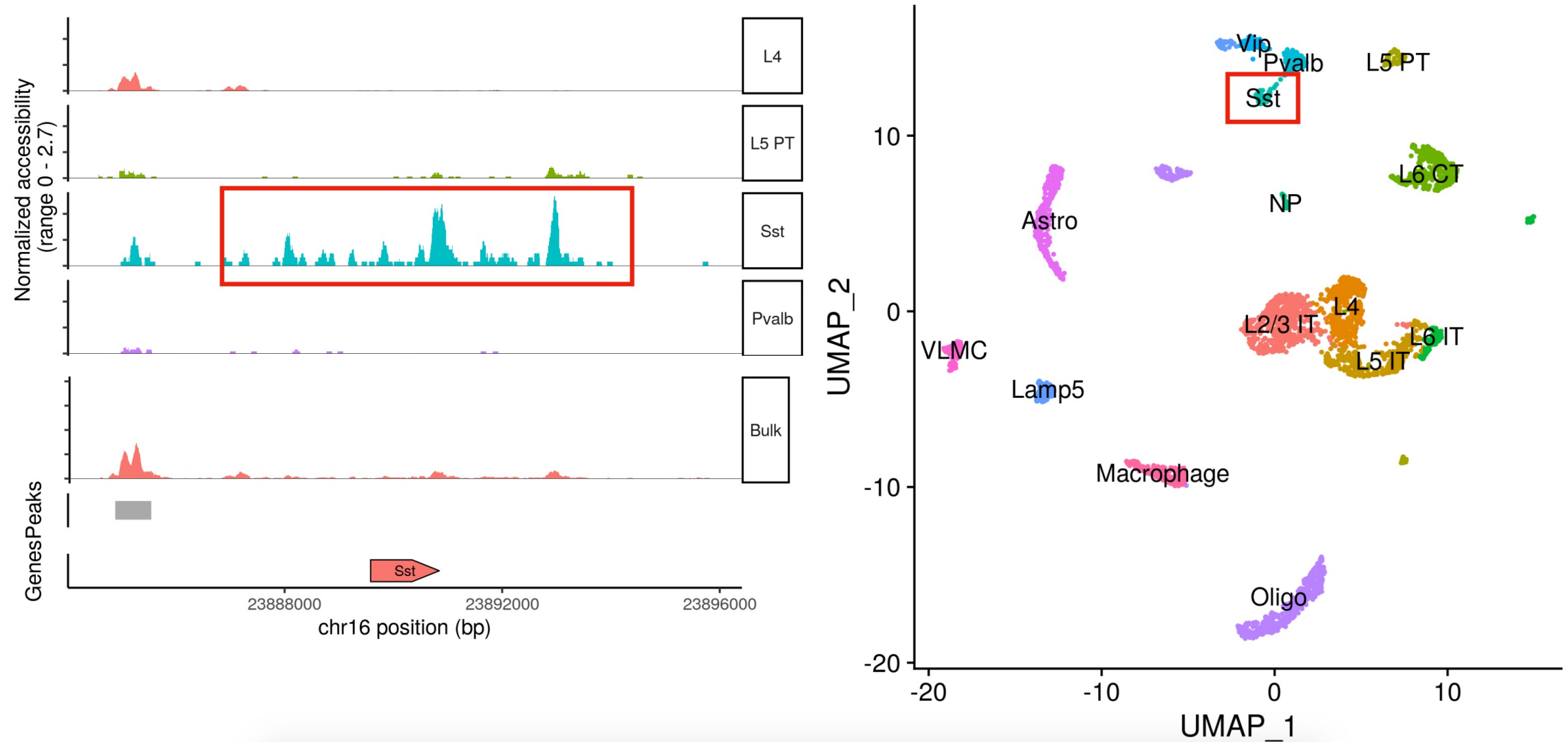


Figure adopted from "Analyzing single-cell ATAC-seq datasets" lecture by Tim Stuart

# Pre-processing generates a fragment file and a peak/cell matrix

A full list of **all** unique fragments across all single cells, as opposed to only reads that map to peaks.

## 1. Indexed fragment file

chrom	start	stop	barcode	reads
chr1	3000141	3000517	GGTTGCGAGCCGCAAA-1	3
chr1	3000159	3000373	CTCAGCTAGTGTCACT-1	1
chr1	3000431	3000621	GAAGTCTGTAACACTC-1	1

## 2. Large sparse matrix

AAACGAAAGAGTTGA-1	AAACGAAAGCGAGCTA-1
.	.
.	.
.	2
.	.
.	.
.	4

Each value in the matrix represents the number of Tn5 cut sites for each single barcode (i.e. cell) that map within each peak

# scATAC-Seq data is highly sparse

## 1. Indexed fragment file

chrom	start	stop	barcode	reads
chr1	3000141	3000517	GGTTGCGAGCCGCAAA-1	3
chr1	3000159	3000373	CTCAGCTAGTGTCACT-1	1
chr1	3000431	3000621	GAAGTCTGTAACACTC-1	1

## 2. Large sparse matrix

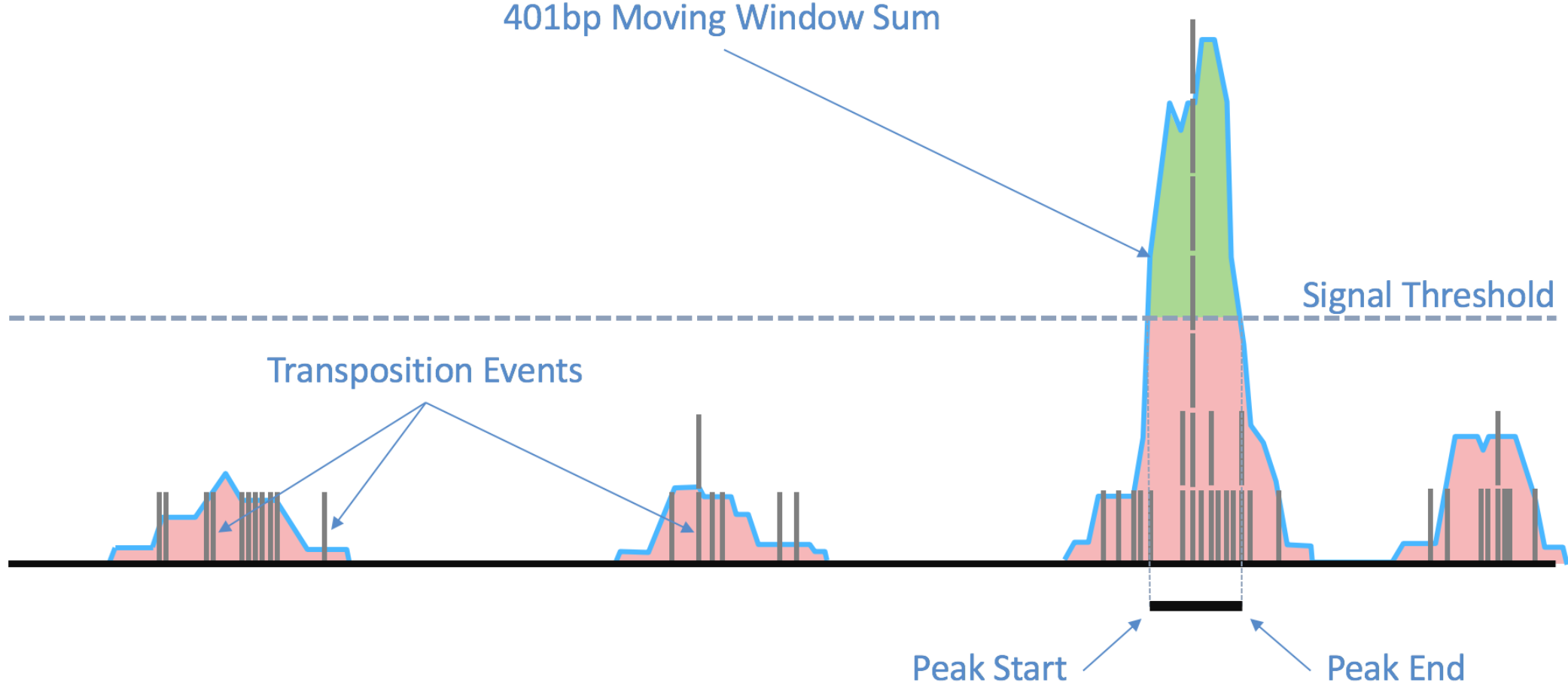
AAACGAAAGAGTTGA-1	AAACGAAAGCGAGCTA-1
.	.
.	.
.	.
2	.
.	.
.	.
4	.

chr1:565107-565550	.
chr1:569174-569639	.
chr1:713460-714823	.
chr1:752422-753038	.
chr1:762106-763359	.

## *Challenges in comparison to scRNA:*

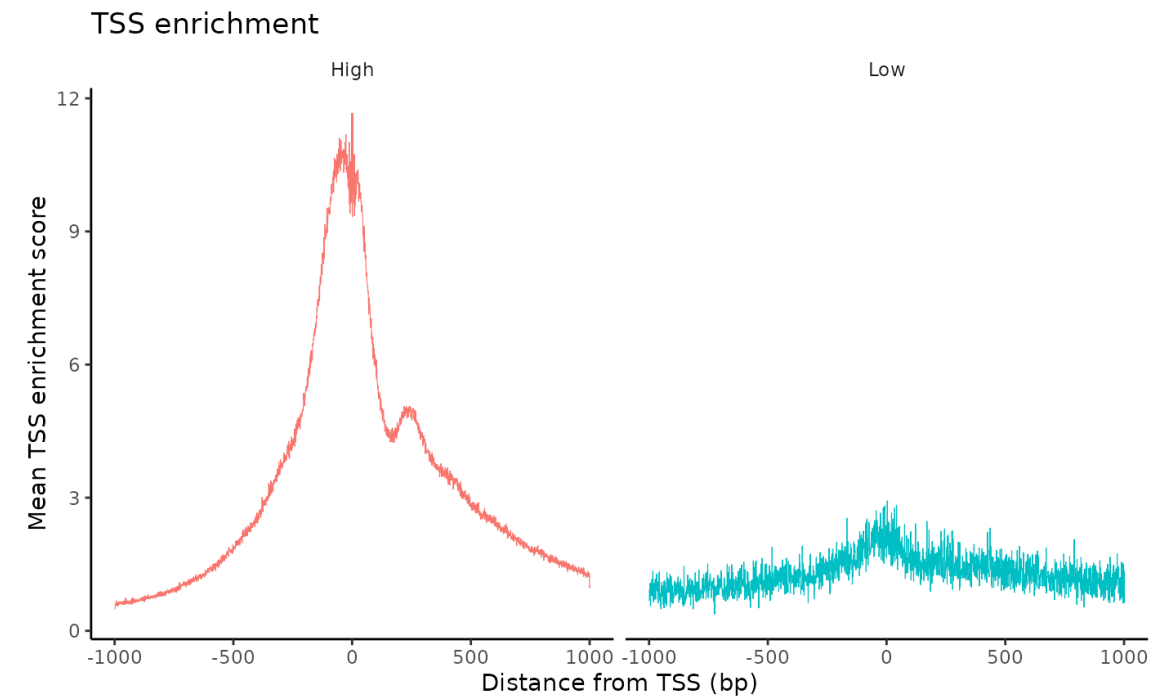
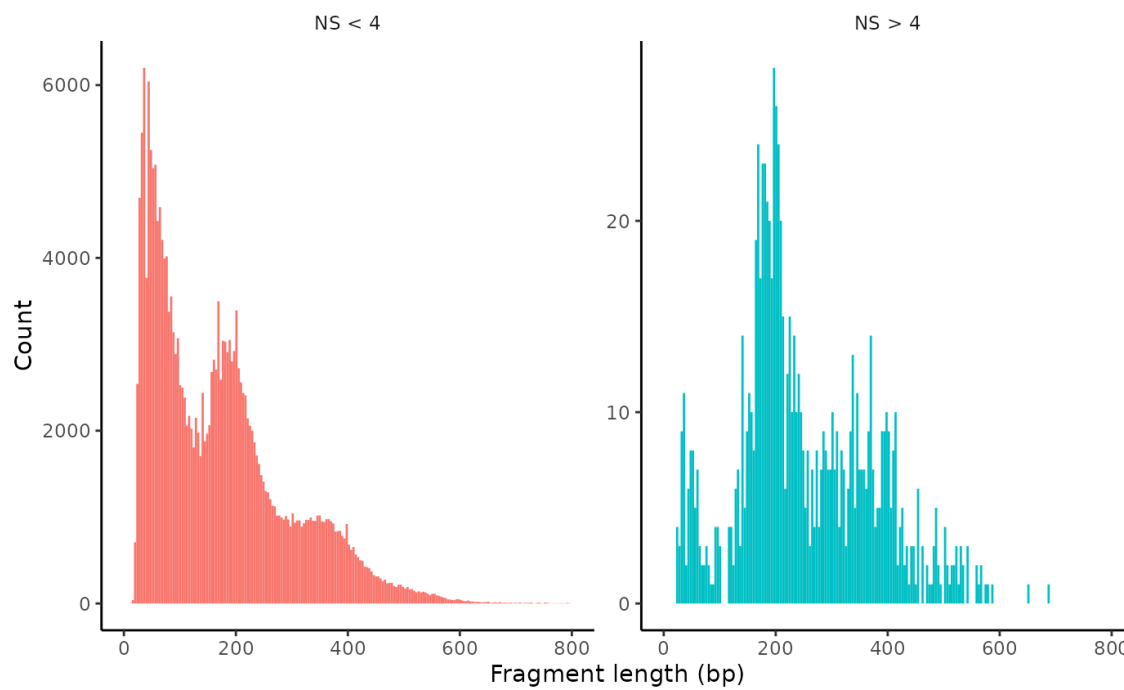
1. More sparse
2. Near-binary data
3. Non-fixed feature set
4. Order of magnitude more features

# Peak calling: from chromatin accessible fragments to peaks



# Quality control metrics for scATAC-seq data

1. Nucleosome banding pattern
2. Transcriptional start site (TSS) enrichment
3. Total number of fragments in peaks
4. Fraction of fragments in peaks



# Overview of scATAC-Seq analysis

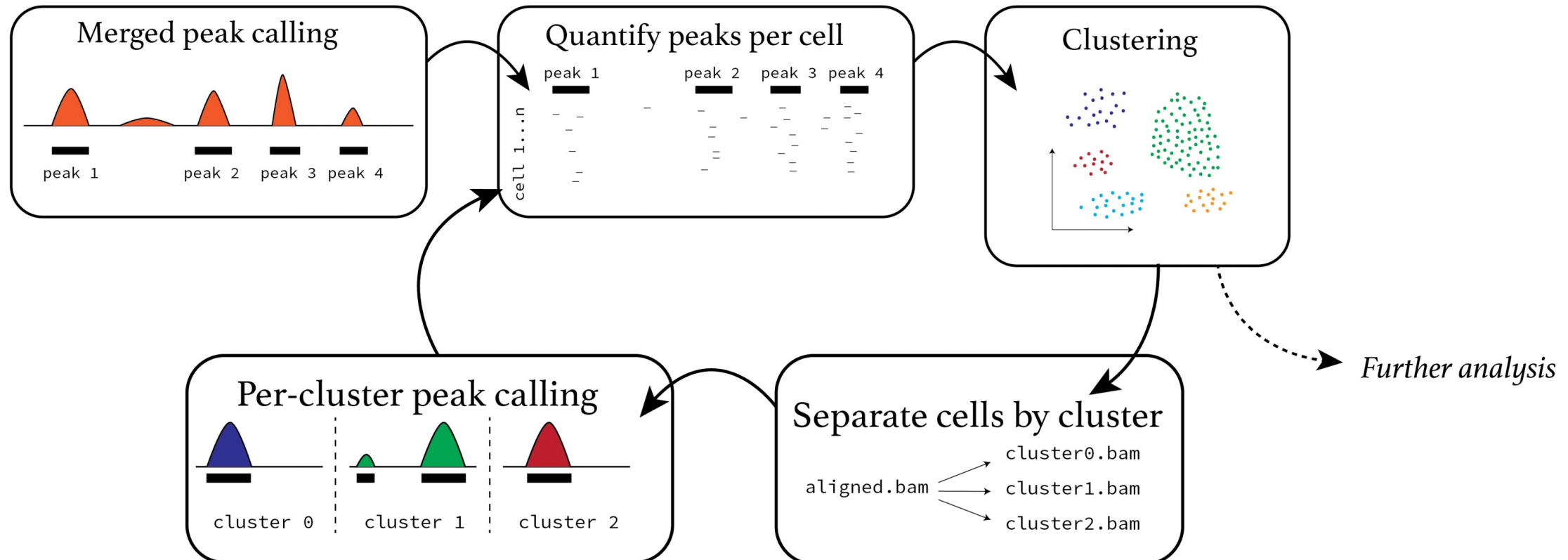


Figure adopted from "Analyzing single-cell ATAC-seq datasets" lecture by Tim Stuart

# scATAC-Seq often uses latent semantic indexing (LSI) for dimensionality reduction

- Originally developed for topic modeling / natural language processing (Deerwester et al. 1990) and first applied to scATAC-Seq in 2015 (Cusanovich et al. *Science*)
- term frequency-inverse document frequency (TF-IDF) normalization followed by singular value decomposition (SVD)
- Cell = document and peak = term
- Term frequency: normalize across cells to correct for differences in sequencing depth
- Inverse document frequency: give higher values to more rare peaks

```
pbmc <- RunTFIDF(pbmc)
pbmc <- FindTopFeatures(pbmc, min.cutoff = 'q0')
pbmc <- RunSVD(pbmc)
```

# Visualize clusters in scATAC-Seq

```
pbmc <- RunUMAP(object = pbmc, reduction = 'lsi', dims = 2:30)
pbmc <- FindNeighbors(object = pbmc, reduction = 'lsi', dims = 2:30)
pbmc <- FindClusters(object = pbmc, verbose = FALSE, algorithm = 3)
DimPlot(object = pbmc, label = TRUE) + NoLegend()
```

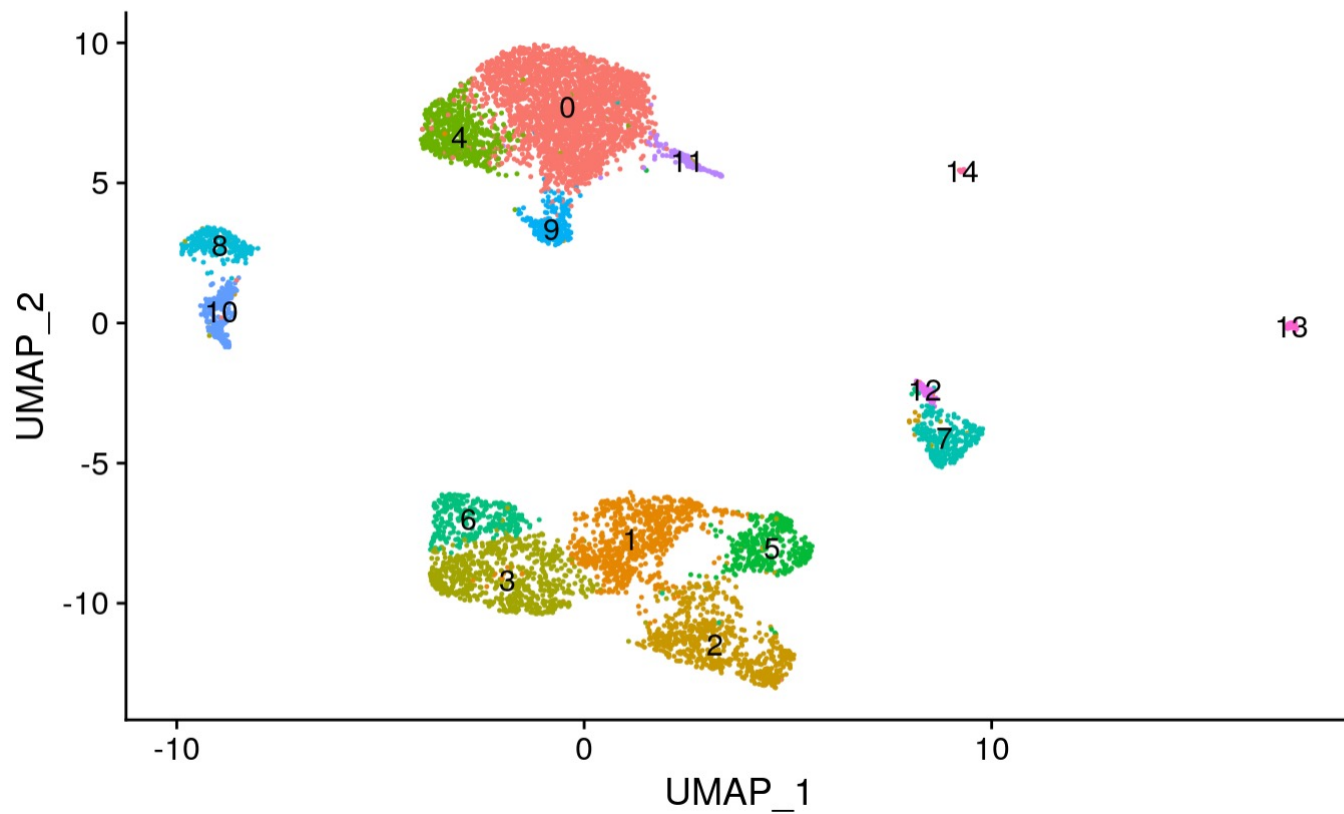
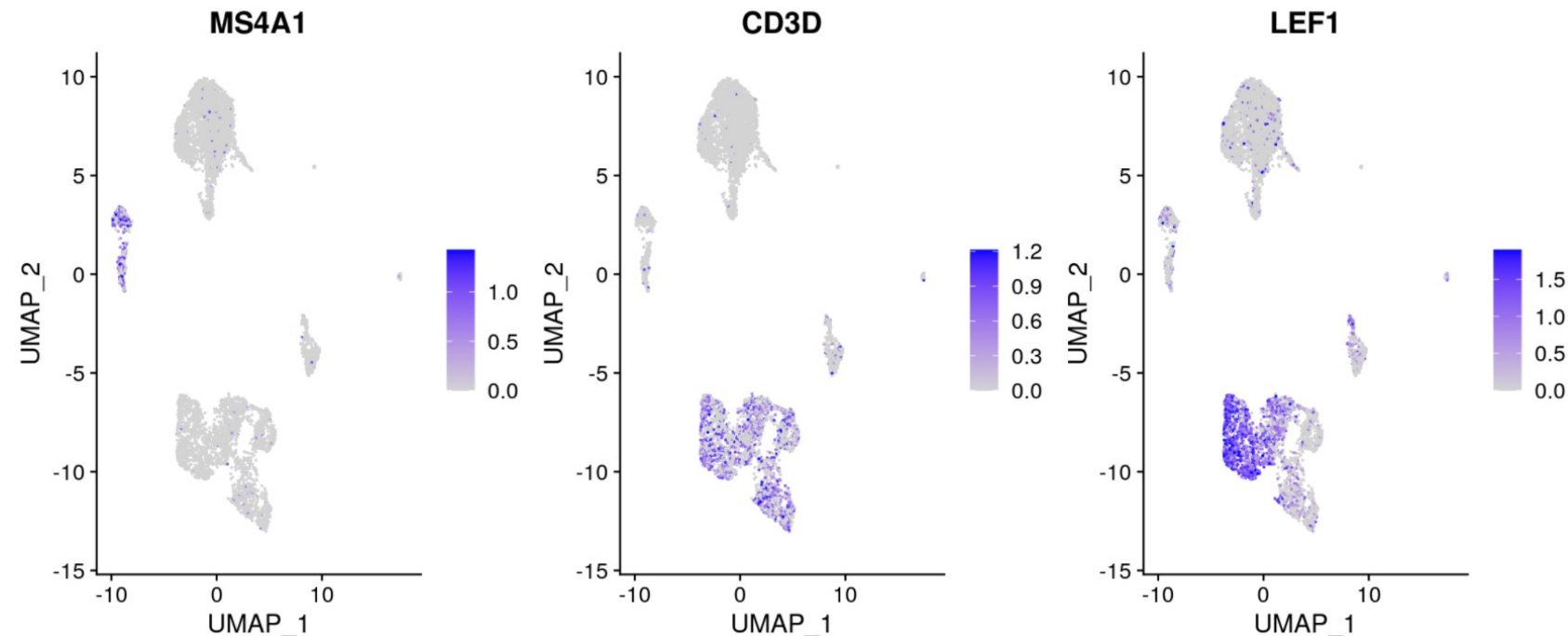


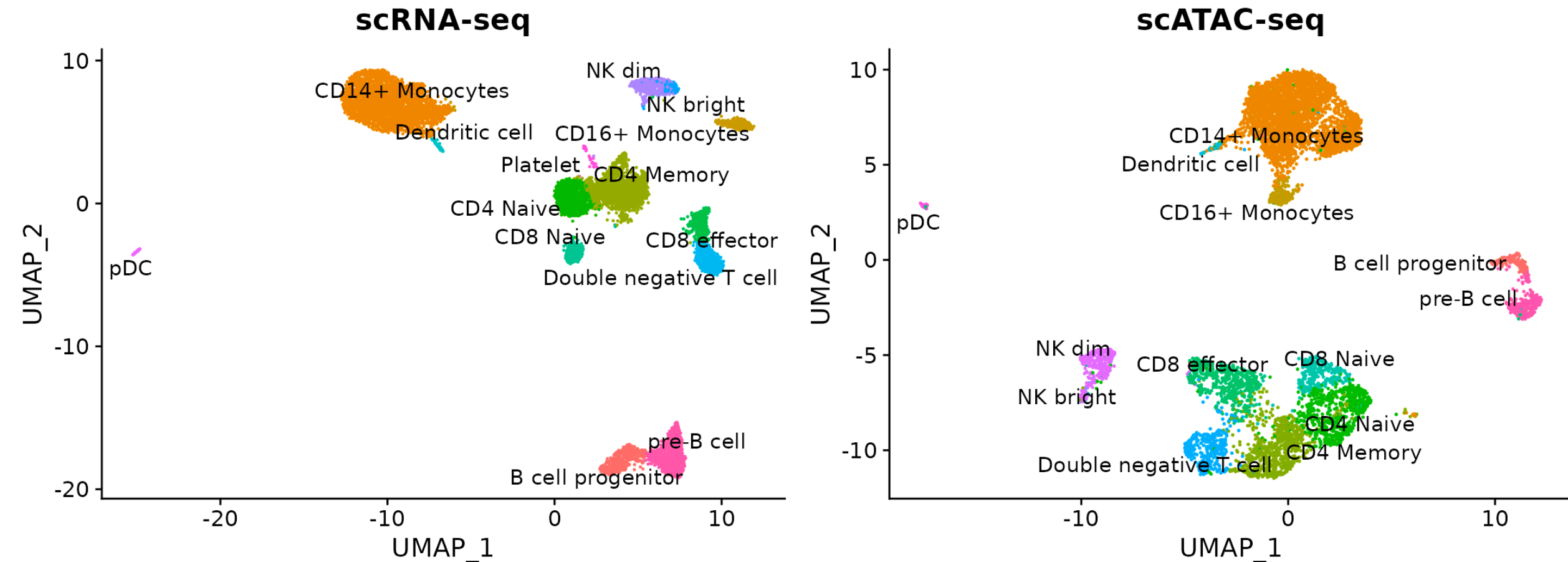
Figure adopted from signac tutorial, [https://satijalab.org/signac/articles/pbmc\\_vignette.html](https://satijalab.org/signac/articles/pbmc_vignette.html)

# Fragment file helps infer gene “activity” and annotate clusters

Quantify the activity of each gene in the genome by assessing the chromatin accessibility associated with each gene: count the number of fragments for each cell that map to the promoter + gene body



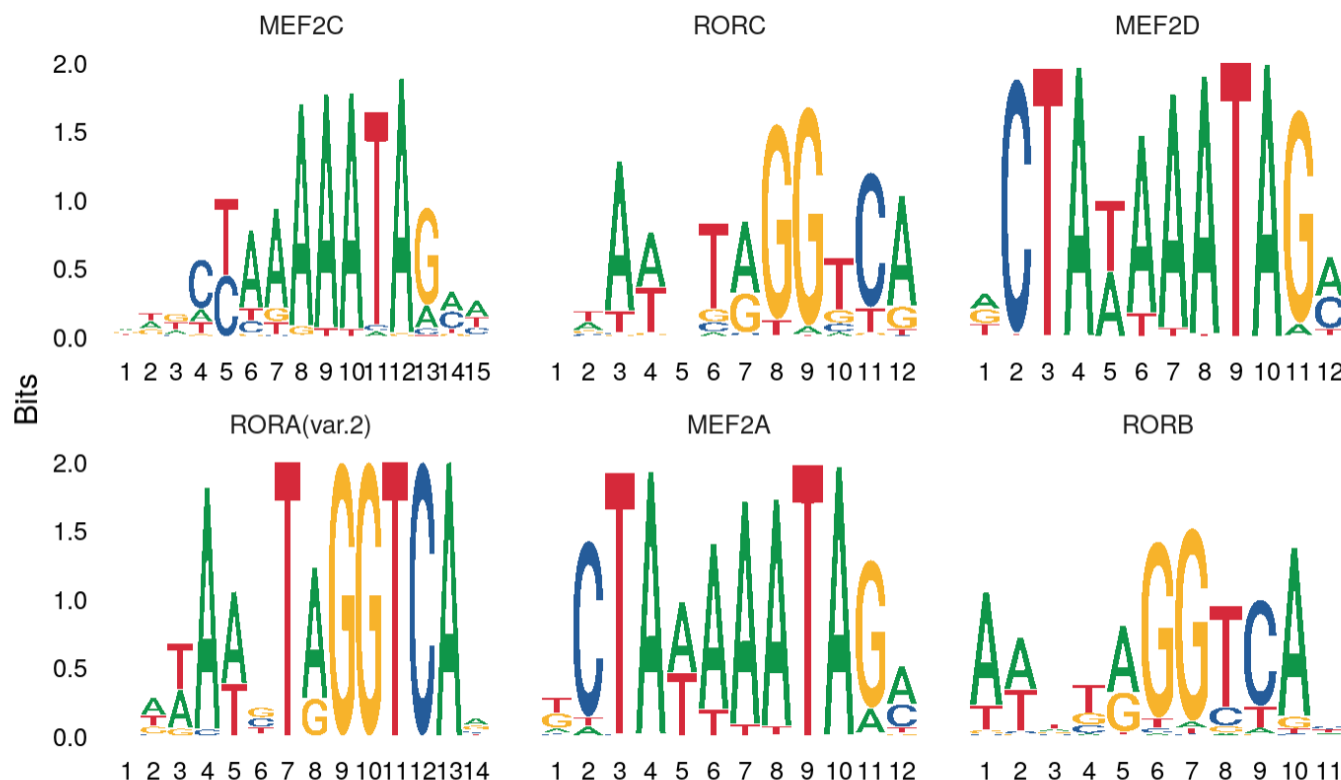
# Integrating with scRNA-seq data using CCA + MNN (Seurat v4)



*Critical assumption: there is generally a positive correlation between chromatin accessibility and gene expression!!!!*

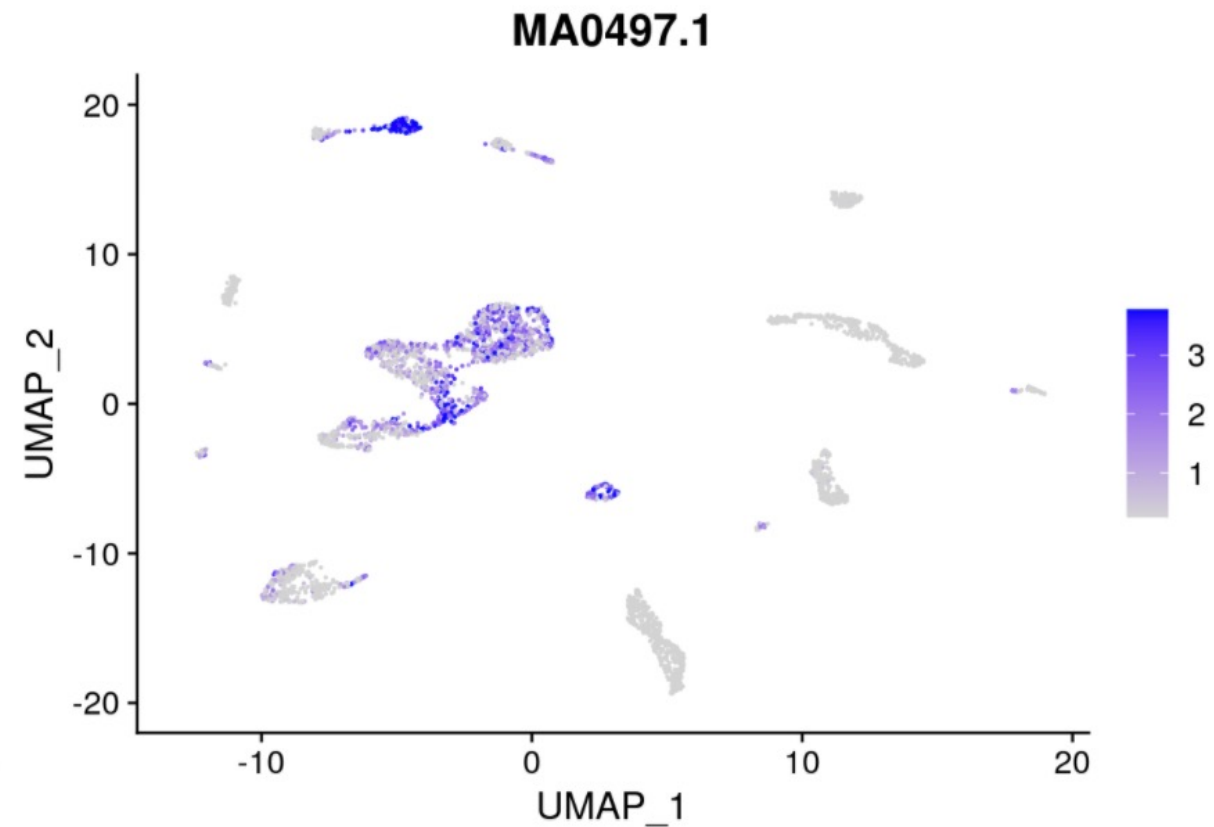
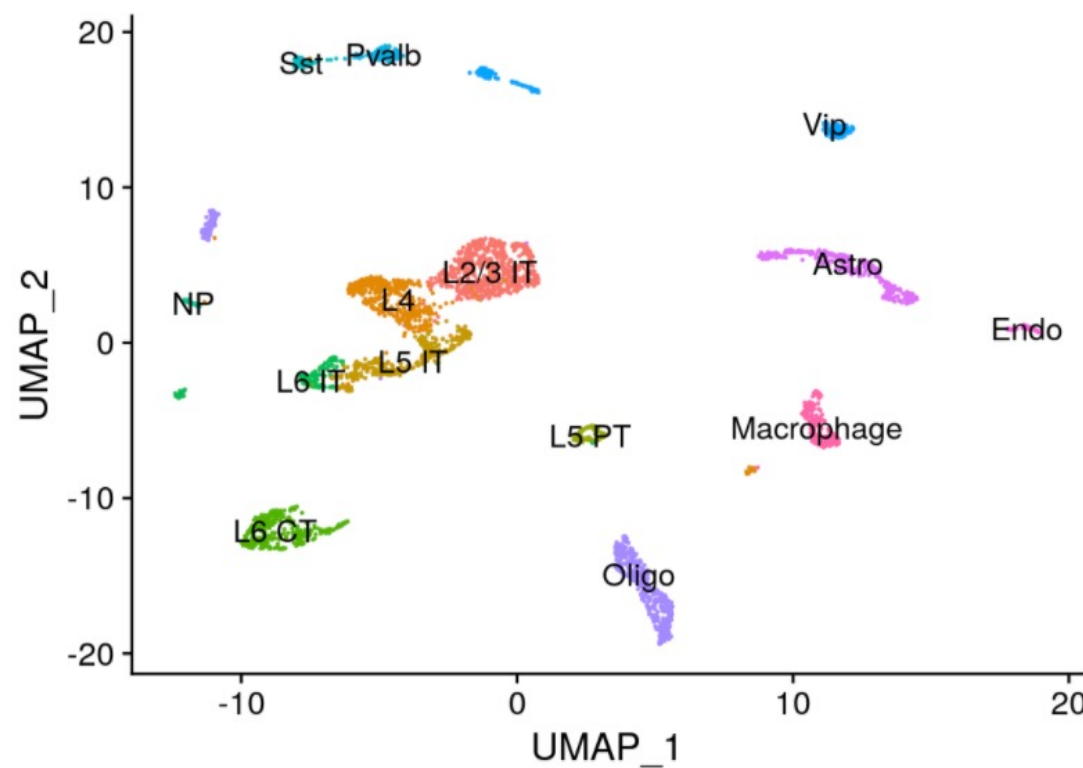
# Finding overrepresented motifs

To identify potentially important cell-type-specific regulatory sequences, signac searches for DNA motifs that are overrepresented in a set of peaks that are differentially accessible between cell types.



# Computing motif activities

ChromVAR identifies motifs associated with variability in chromatin accessibility between cells.



# Software for scATAC-Seq analysis

<b><i>Signac</i></b>	<a href="https://satijalab.org/signac/index.html">https://satijalab.org/signac/index.html</a>
<b><i>SnapATAC</i></b>	<a href="https://github.com/r3fang/SnapATAC">https://github.com/r3fang/SnapATAC</a>
<b><i>ArchR</i></b>	<a href="https://www.archrproject.com/">https://www.archrproject.com/</a>
<b><i>cisTopic</i></b>	<a href="https://github.com/aertslab/cisTopic">https://github.com/aertslab/cisTopic</a>
<b><i>chromVAR</i></b>	<a href="https://github.com/GreenleafLab/chromVAR">https://github.com/GreenleafLab/chromVAR</a>
<b><i>CICERO</i></b>	<a href="https://cole-trapnell-lab.github.io/cicero-release/">https://cole-trapnell-lab.github.io/cicero-release/</a>
<b><i>episcanpy</i></b>	<a href="https://episcanpy.readthedocs.io/en/latest/">https://episcanpy.readthedocs.io/en/latest/</a>

# Software for scATAC-Seq analysis

	ArchR	Signac	SnapATAC	
Pre-processing	NR	NA	✓	
Data import / base file type creation	✓	NA	✓	
QC filter cells	✓	✓	✓	
Matrix creation	✓ (Tile)	✓ (Peak)	✓ (Tile)	
Doublet removal	✓	NP	NP	Doublet Removal
Data imputation with MAGIC	✓	NP	✓	
Genome-wide gene score matrix	✓	✓	✓	
Dimensionality reduction and clustering	✓	✓	✓	Clustering
UMAP and tSNE plotting	✓	✓	✓	
Cluster peak calling	✓	NP	✓	
Cluster-based peak matrix creation	✓	NP	✓	
Motif enrichment	✓	✓	✓	Standard ATAC-seq Analyses
chromVAR motif deviations	✓	✓	✓	
Footprinting	✓	NP	NP	
Feature set annotation	✓	NP	NP	
Track plotting	✓	✓	NP	
Co-accessibility	✓	NP	NP	Data Visualization
Interactive genome browser	✓	NP	NP	
Cellular trajectory analysis	✓	NP	NP	Advanced ATAC-seq Analyses
Project bulk data into scATAC embedding	✓	NP	NP	
Integration of RNA-seq and ATAC-seq	✓	✓	✓	Integration of RNA-seq and ATAC-seq
Genome-wide peak-to-gene links	✓	NP	NP	

NR = Not Required

NA = Not Applicable

NP = Not Possible

# Some approaches to multiome data (scRNA-seq and scATAC-seq)

ArchR: [https://greenleaflab.github.io/ArchR\\_2020/Ex-Analyze-Multiome.html](https://greenleaflab.github.io/ArchR_2020/Ex-Analyze-Multiome.html)

Signac (same people who built Seurat): [https://satijalab.org/signac/articles/pbmc\\_multomic.html](https://satijalab.org/signac/articles/pbmc_multomic.html)

FigR: <https://buenrostrolab.github.io/FigR>

# Methods to overcome sparsity in ATAC-seq data

Computing meta-cells is one methodology used to overcome sparsity in scATAC-seq data

Computing meta-cells (e.g. SEACells algorithm) can improve computation of peak-gene associations

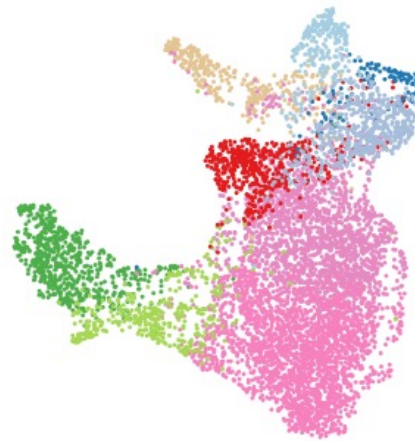
scRNA-seq

celltype



scATAC-seq

celltype



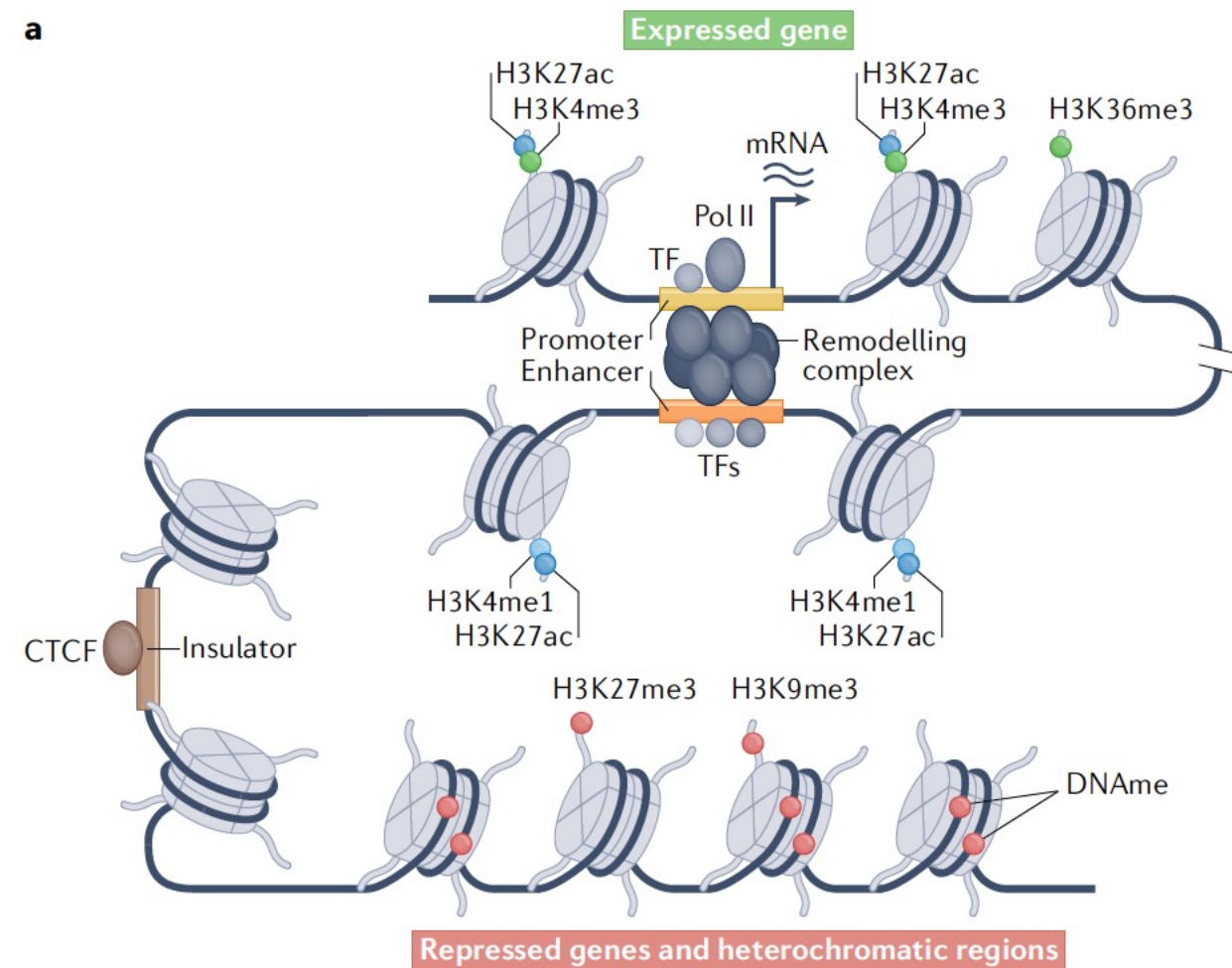
- CLP
- Ery
- HMP
- HSC
- MEP
- Mono
- cDC
- pDC
- DCPre

Metacell Assignments

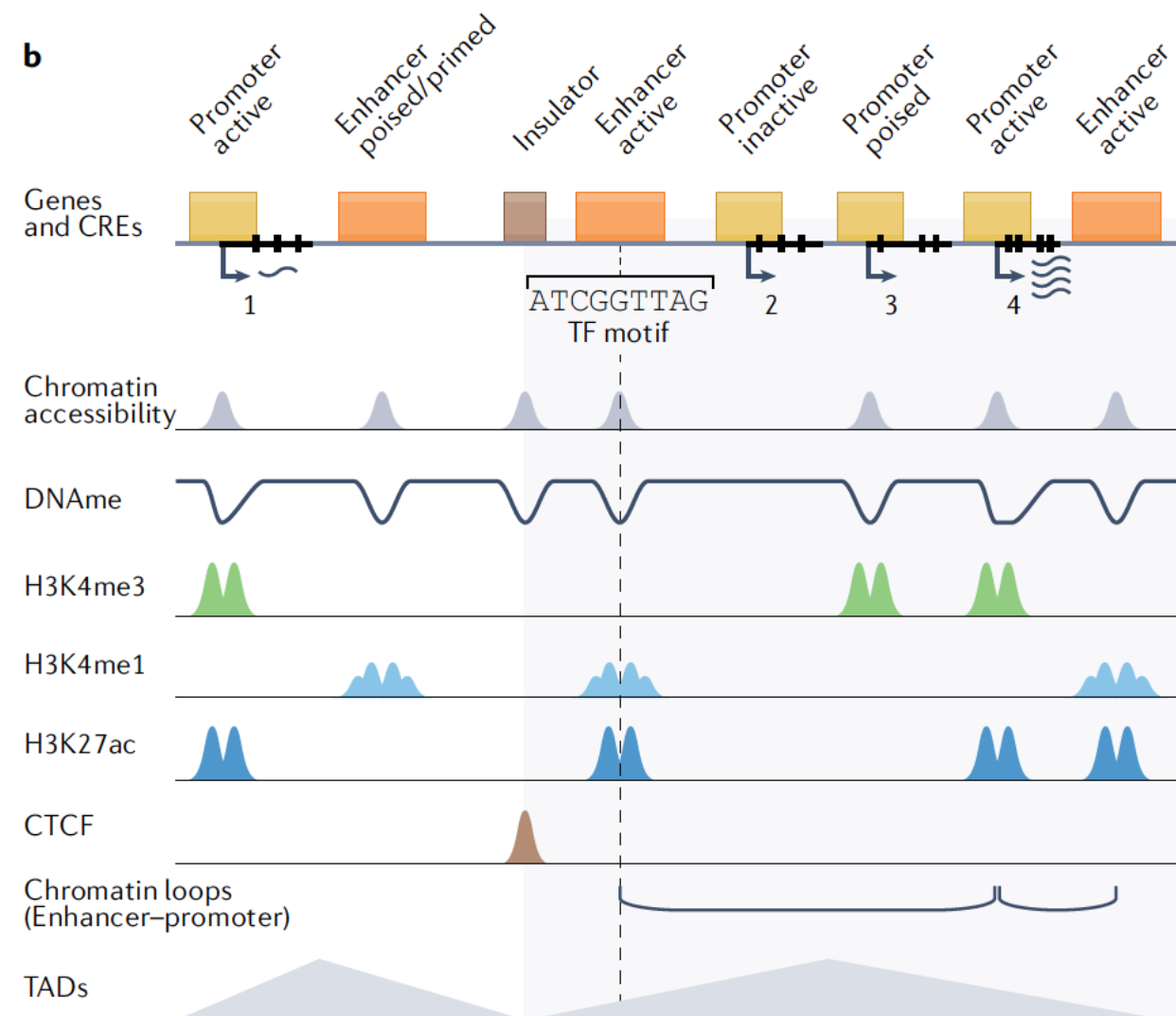


# Studying gene regulation using single-cell epigenomics

a



b



# Studying gene regulation using single-cell epigenomics

