



SciLifeLab

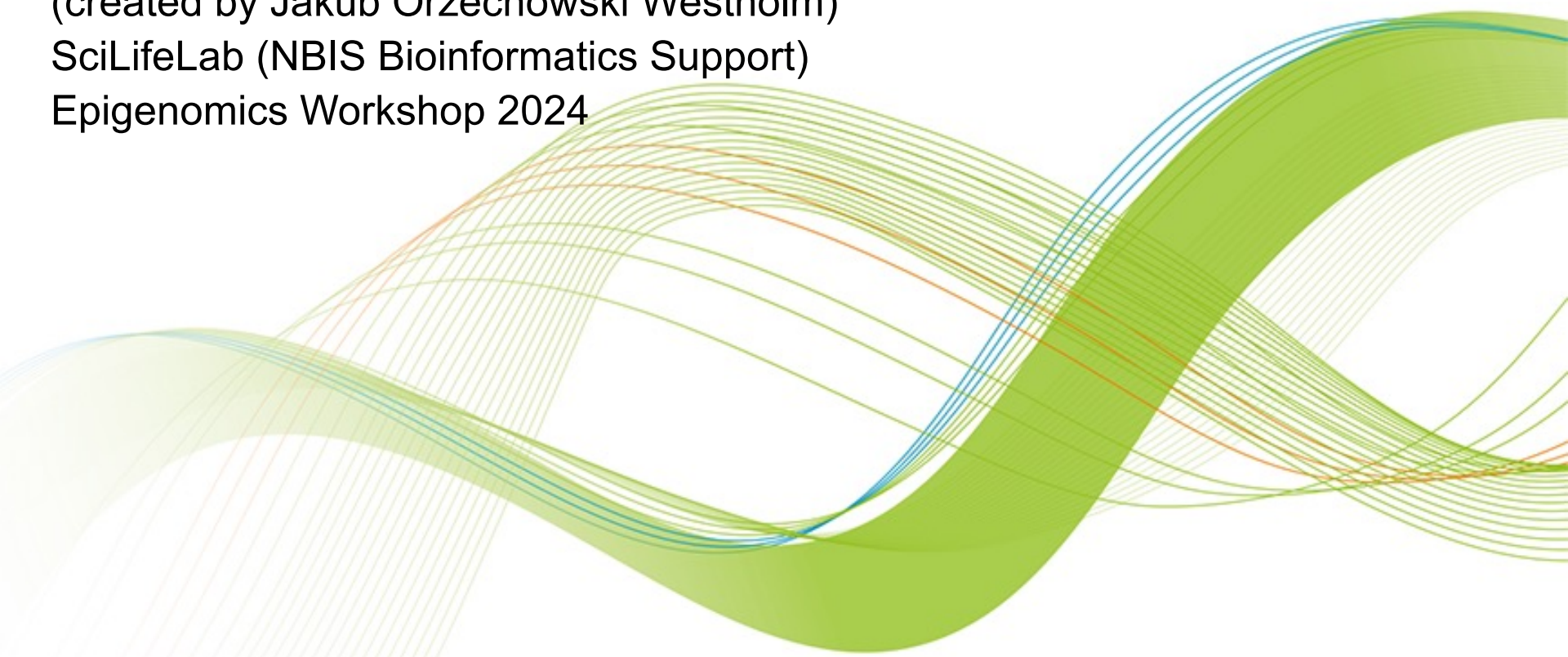
Single cell methods in epigenomics

Markus Ringnér

(created by Jakub Orzechowski Westholm)

SciLifeLab (NBIS Bioinformatics Support)

Epigenomics Workshop 2024



Single cell methods

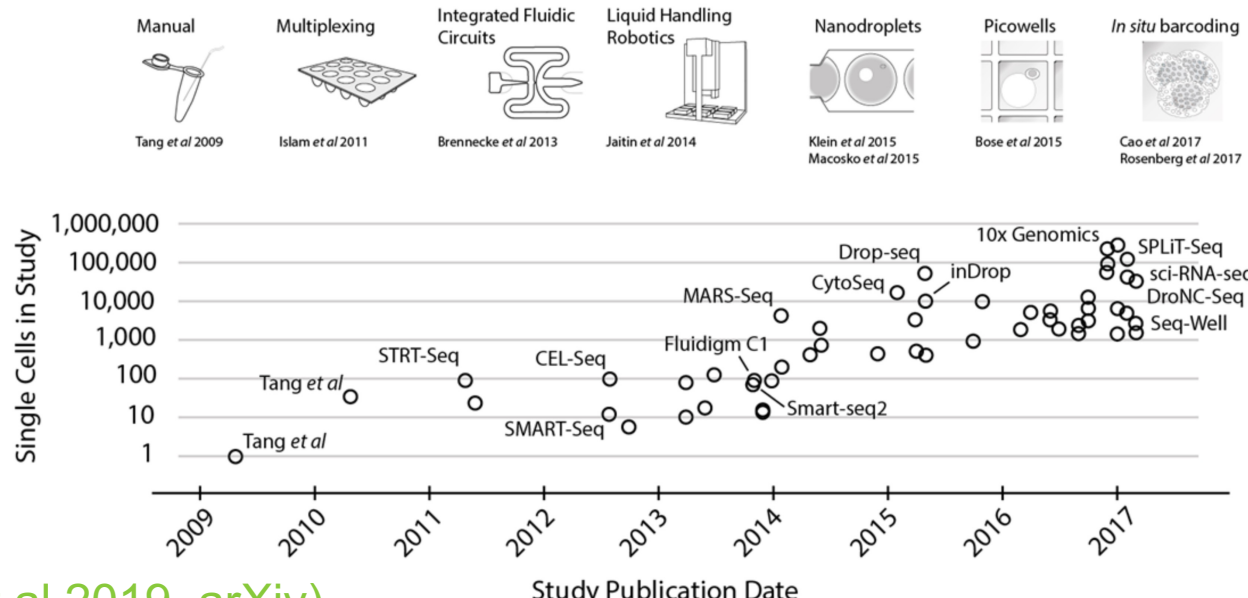
- Analyzing bulk samples will give results that represent an average of the cell population, over thousands or millions of cells
- Single cell methods give results for each cell
 - Differences between individual cells (due to stochasticity, cell cycle etc.)
 - Differences between cell types
- More noise in data
- Require new analysis steps
- This is a big, and rapidly changing field. We have a whole course on single cell RNA-seq. In this talk we will give a short introduction to single cell methods in epigenomics.

Todays talk

- Background on single cell methods
- Single cell ATAC-seq
- Single cell data analysis, common steps
- Single cell ChIP-seq & CUT&TAG
- Single cell DNA-methylation
- Spatial methods
- Broad overview
- Focus on concepts over details.

Background

- Single cell methods started with RNA-seq
 - First paper: ([Tang et al 2009, Nature Methods](#))
 - Output has increased a lot, from 1 cell in 2009 to millions of cells today
- Later this was adapted for ChIP-seq, ATAC-seq, DNA methylation and other assays
- Some experimental steps and analysis are similar, some are unique



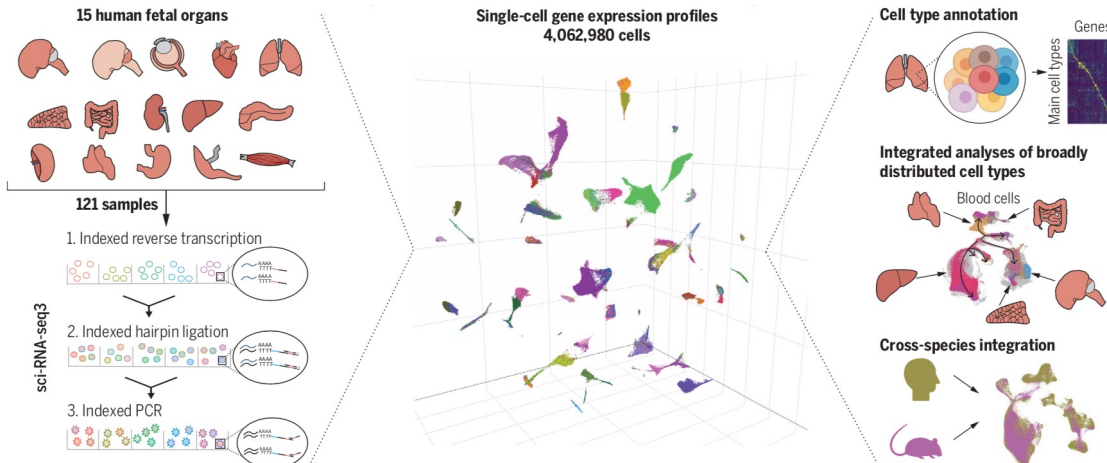
Example study

RESEARCH ARTICLE

HUMAN GENOMICS

A human cell atlas of fetal gene expression

Junyue Cao^{1*}, Diana R. O'Day², Hannah A. Pliner³, Paul D. Kingsley⁴, Mei Deng², Riza M. Daza¹, Michael A. Zager^{3,5}, Kimberly A. Aldinger^{2,6}, Ronnie Blecher-Gonen¹, Fan Zhang⁷, Malte Spielvogel⁸, James Palis⁴, Dan Doherty^{2,3,6}, Frank J. Steemers⁷, Ian A. Glass^{2,3,6}, Cole Trapnell^{1,3,10,†}, Jay Shendure^{1,3,10,11,†}



RNA-seq: 4 million cells
ATAC-seq: 800,000 cells

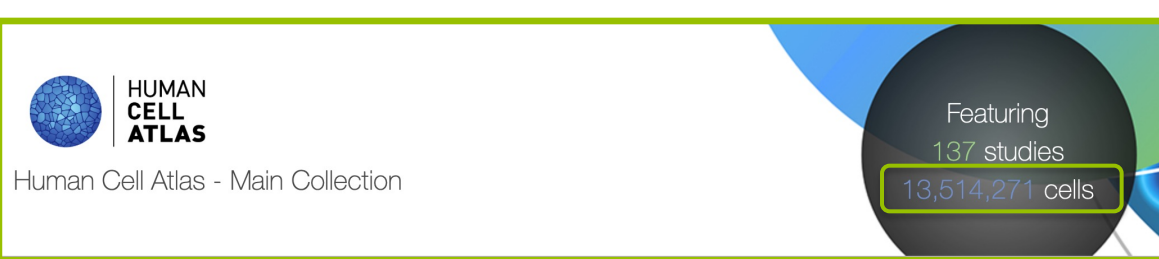
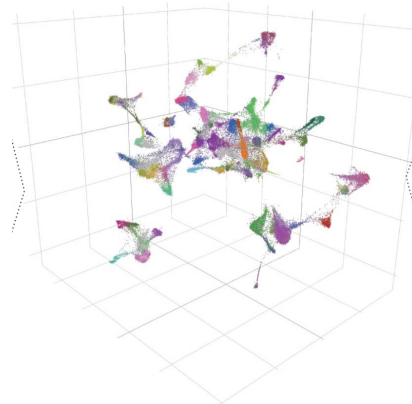
RESEARCH ARTICLE SUMMARY

HUMAN GENOMICS

A human cell atlas of fetal chromatin accessibility

Silvia Domcke^{*}, Andrew J. Hill^{*}, Riza M. Daza^{*}, Junyue Cao, Diana R. O'Day, Hannah A. Pliner, Kimberly A. Aldinger, Dmitry Pokholok, Fan Zhang, Jennifer H. Milbank, Michael A. Zager, Ian A. Glass, Frank J. Steemers, Dan Doherty, Cole Trapnell[†], Darren A. Cusanovich[†], Jay Shendure[†]

Single-cell chromatin accessibility profiles
790,957 cells

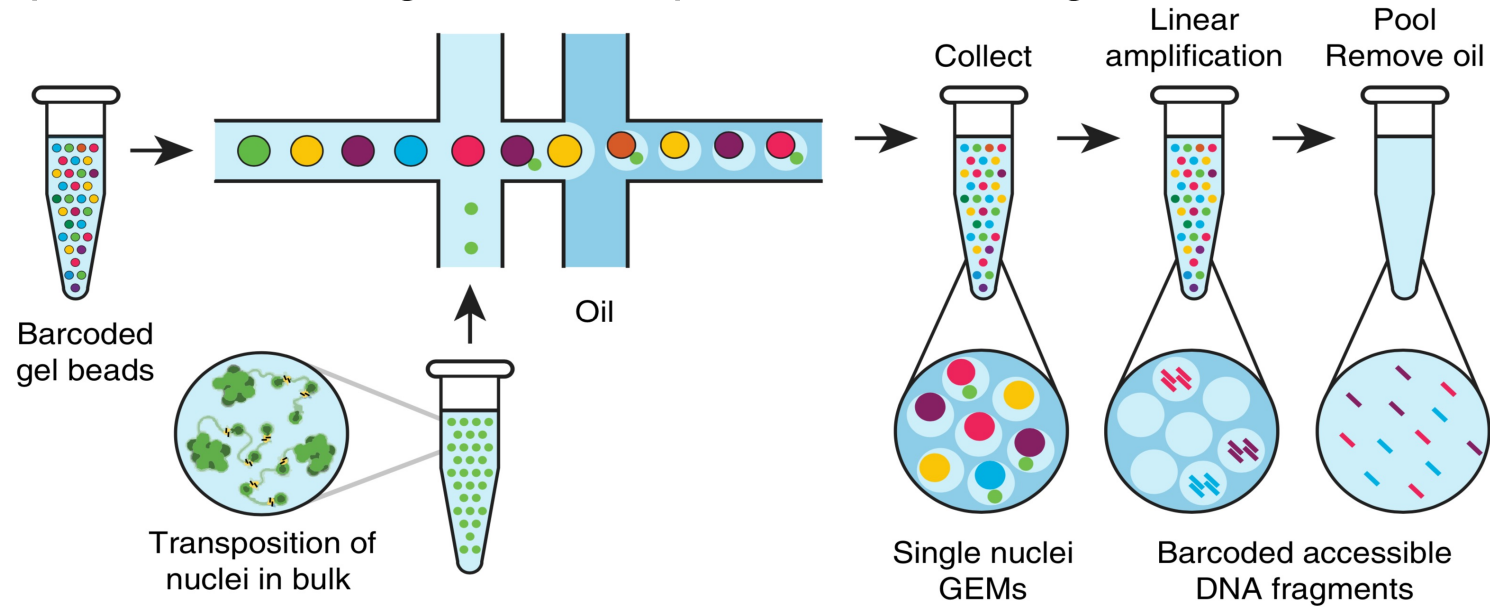


Featuring
137 studies

13,514,271 cells

Single cell ATAC-seq

- First paper, from Greenleaf lab: (Buenrostro et al. 2015, Nature)
- Now available as a kit from 10X Genomics
 - Each cell is attached to a bead containing a different barcode, inside an oil droplet.
 - These barcodes are attached to the DNA fragments, making it possible to assign each sequenced DNA fragment to a cell.

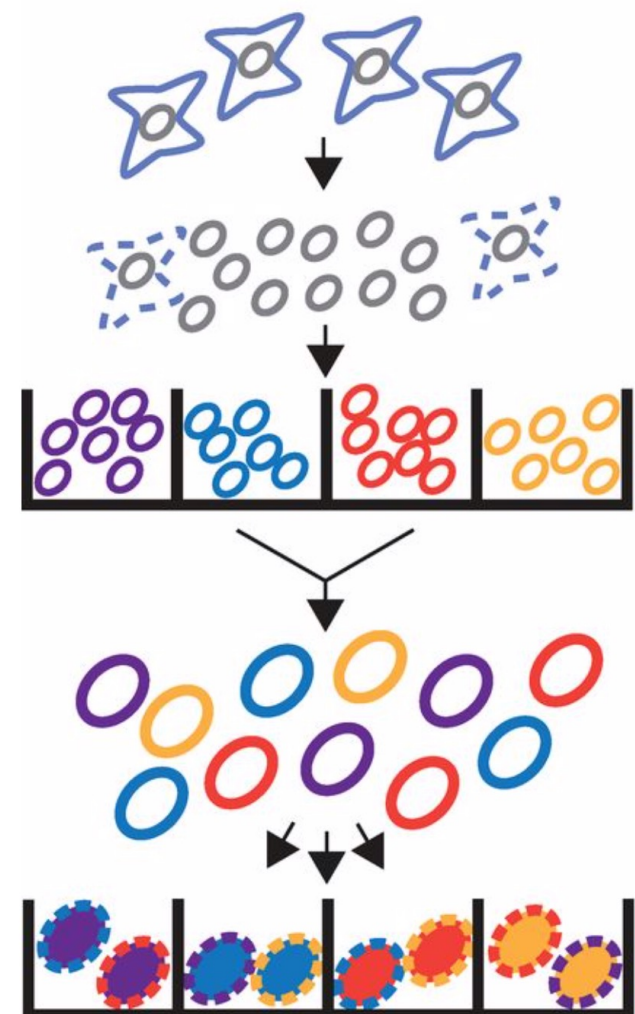


(Satpathy et al. 2019 Nature Biotechnology)



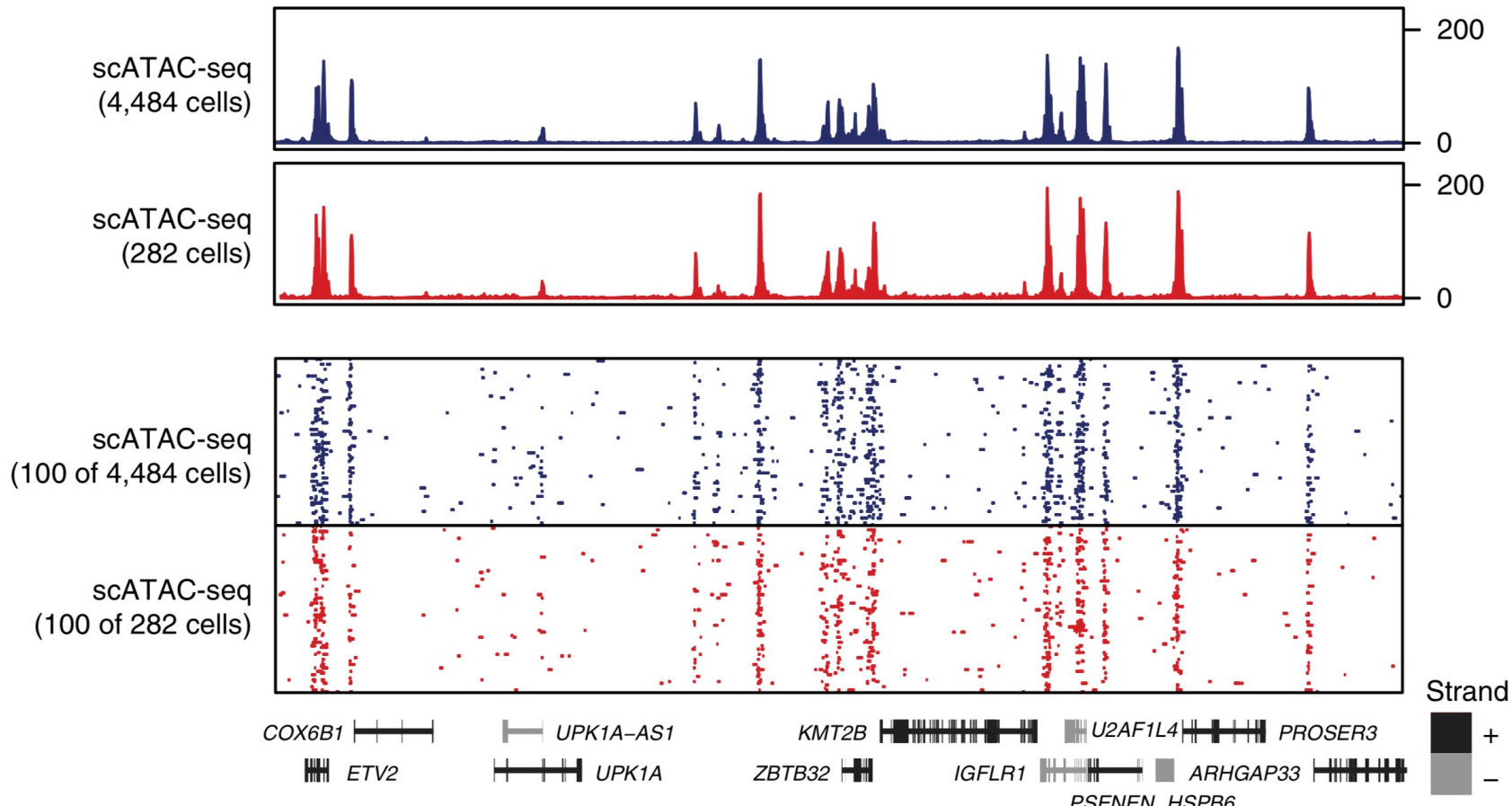
Single cell ATAC-seq II

- An alternative to the droplet based method from 10X genomics is **sci-ATAC-seq** (single-cell combinatorial-indexing with ATAC-seq).
(Cusanovich et al. 2015, Science)
 - Here, cells are split up into e.g. 96 wells, and each well has a different short barcode.
 - Cells are then pooled and re-distributed into wells again, adding another short barcode.
 - This is repeated enough times so that each cell will eventually have it's own (almost) unique combination of short barcodes.
-
- + Low cost per cell, enables high throughput
 - Lower cell recovery, important when there is limited starting material

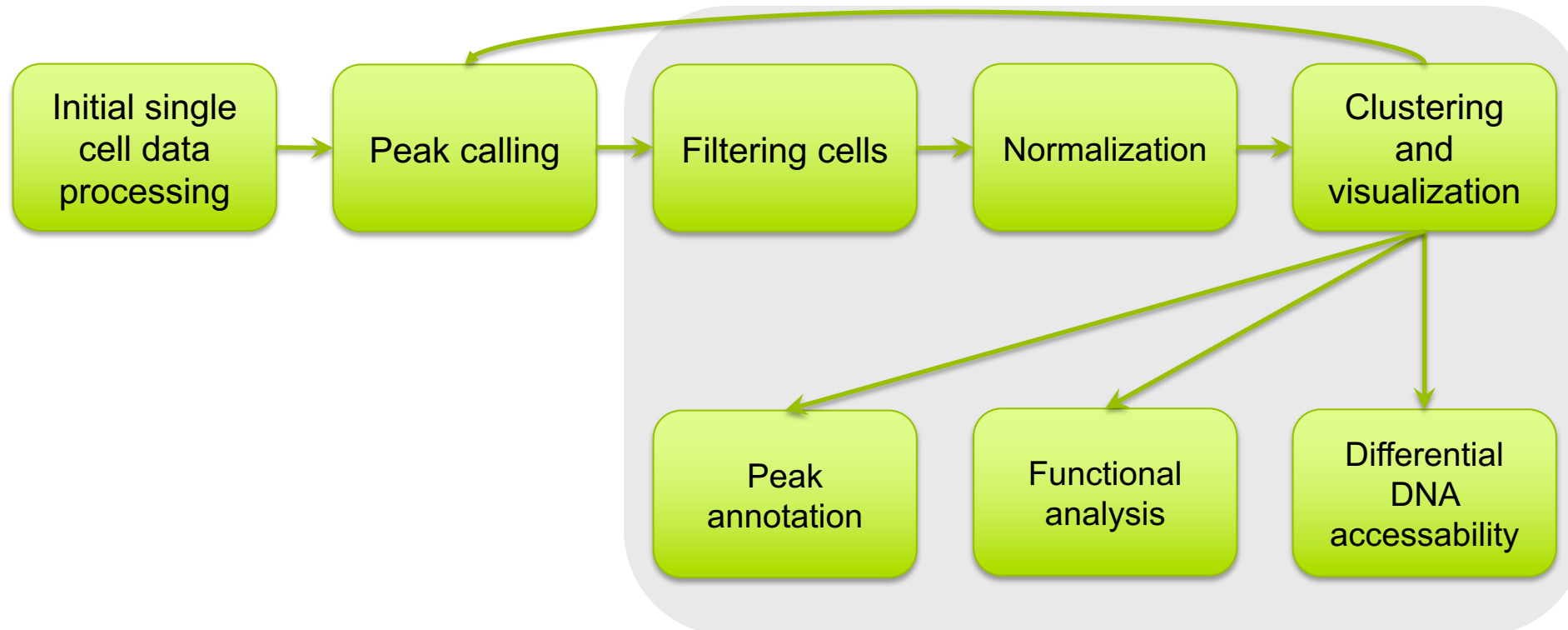


Single cell ATAC-seq data

- Looking at each individual cell, scATAC-seq data are **sparse** and **noisy**.
- But combining data from lots of cells gives meaningful signals.



Single cell ATAC-seq data analysis



Exercise later today

1. Initial single cell data processing



-
- De-multiplex: Using the cell specific barcodes, assign each read to a cell.
 - (Remove primer sequences.)
 - Map reads to the genome, e.g. with **BWA-MEM**.
 - Remove duplicates: If several read pairs map to exactly the same coordinates, only one is kept. Such duplicates are assumed to be PCR artifacts.
 - Filter out some bad cells already at this stage.

2. Peak calling

- Similar to peak calling for bulk data.
- Done on aggregated data from all cells. (There is not enough data in a single cell to call peaks.)
- If we have a rare cell type with e.g. 50 out of 2000 cells, peaks specific to this cell type can be missed when we use the aggregated data for peak calling.
 - We can go back and redo the peak calling later, only looking at specific groups of cells.
- We then count the reads from every cell in every peak:

	Cell 1	Cell 2	Cell 3	...	Cell M
Peak 1	0	1	1		0
Peak 2	0	0	0		0
Peak 3	0	0	0		1
...					
Peak N	1	0	0		0

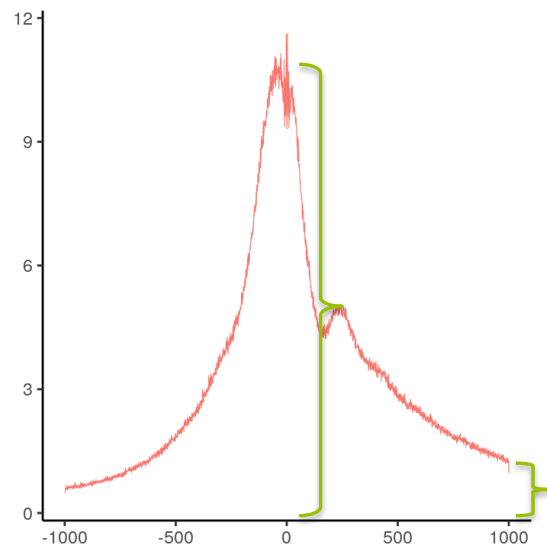
Mostly 0s

3. Filtering cells

- There are many things that could go wrong in a single cell ATAC-seq experiment
 - No cell in a droplet
 - Several cells in a droplet
 - Dead cells
 - Few reads from a cell
 - No transfection in a cell
- Therefore we use several quality measures to identify and remove problematic cells/barcodes:
 - Number of fragments in peaks: Cells with very few reads may need to be excluded due to low sequencing depth. Cells with extremely high levels may represent doublets, nuclei clumps, or other artefacts.
 - Fraction of fragments in peaks: Cells with low values (i.e. <15-20%) often represent low-quality cells or technical artifacts that should be removed.

3. Filter cells II

- Reads in blacklist regions: The ENCODE project has provided a list of blacklist regions, i.e. regions with artefactual signal. Cells with many reads mapping to these blacklist regions (compared to reads mapping to peaks) often represent technical artifacts and should be removed.
- Transcriptional start site (TSS) enrichment score. TSS are associated with open chromatin, so a low level of chromatin enrichment would suggest poor ATAC-seq experiments.

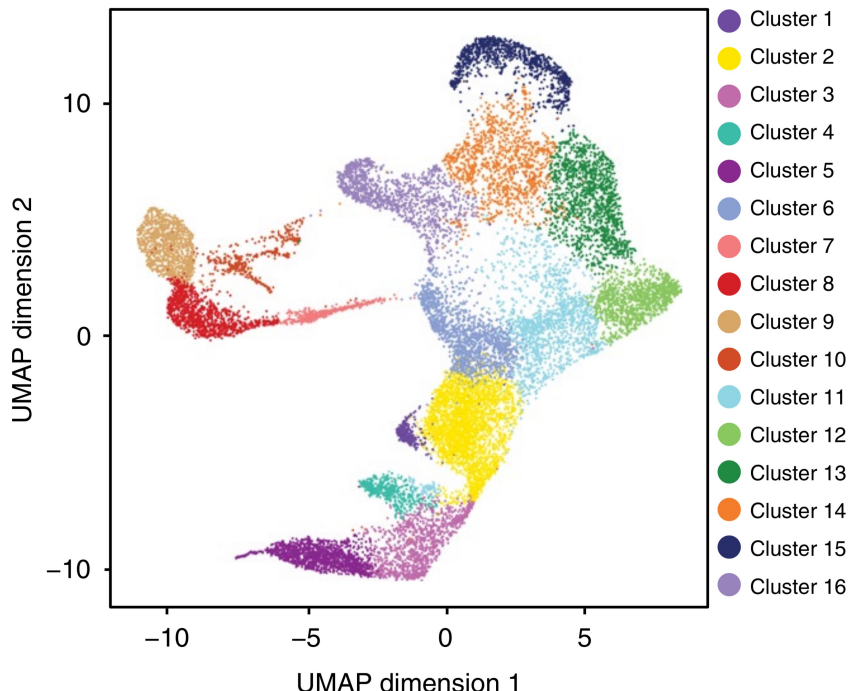


4. Normalization

- Account for different sequencing depth in different cells
- Create a simplified representation of the data, using dimension reduction (singular value decomposition). This is similar to principal component analysis (PCA).
 - The idea behind this is to reduce noise, and to select informative features to improve clustering of cells and visualization
 - Typically, the first component correlates with sequencing depth, so by removing it we get rid of artefactual signal.
 - Reducing dimensionality is often good in itself.
 - Results are often better when we select only some features (peaks)
 - Those with highest signal
 - Those with highest variability

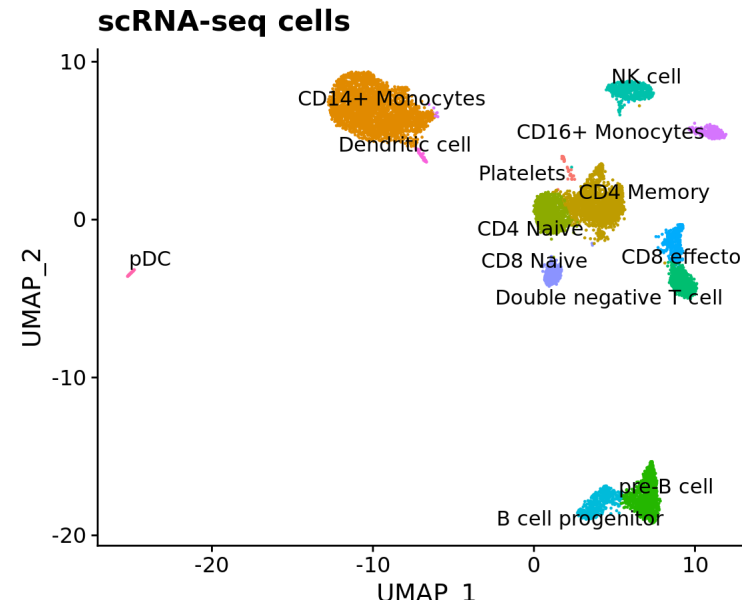
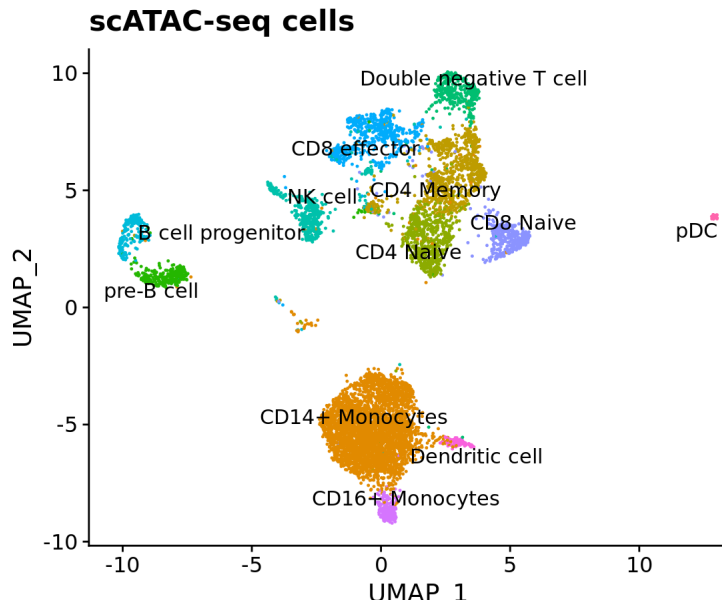
5. Cluster and visualize cells

- Visualize how cells relate to each other
 - Project many dimensions down to 2.
 - Conceptually similar to PCA, but not linear
 - UMAP algorithm
- Clustering, to identify groups of similar cells (representing different cell types or cell states).
 - There are many different clustering methods
 - Many settings for such methods
 - Trial and error



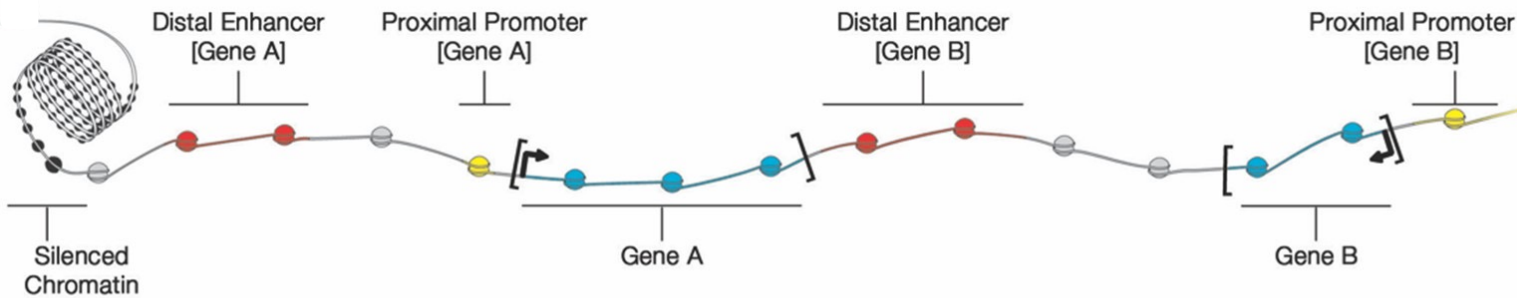
5. Cluster and visualize cells II

- It's often not clear which cell types etc. these clusters represent.
 - In single cell RNA-seq we can look at marker genes, unique to a specific cell type. In single cell ATAC-seq, this is harder.
 - If it's possible to get RNA-seq data from a similar set of cells, these can be annotated and then used to annotate the ATAC-seq clusters.
 - This is sometimes called label transfer, will be discussed in the next talk, about integrating -omics data, and in the exercise.



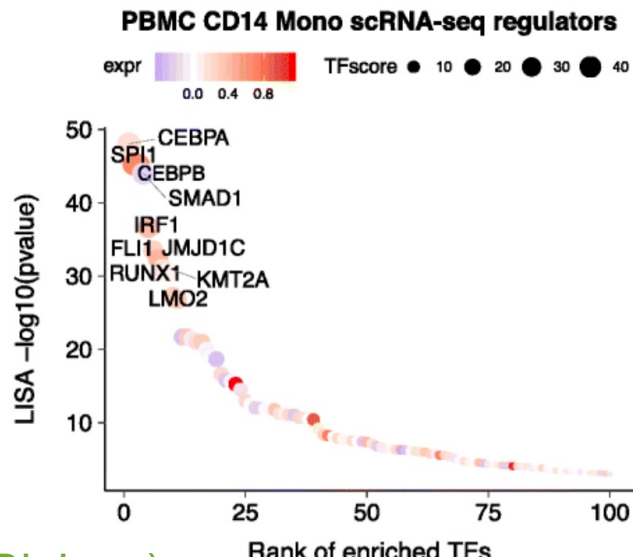
6. Peak annotation

- To easier interpret the peaks, it's useful to note their location with regard to the nearest gene (or the nearest TSS).
 - Remember that a region might not interact with the nearest gene, this is just a starting guess!
- This is similar to what was discussed for ATAC-seq and ChIP-seq data.



7. Functional analysis

- Like for bulk ATAC-seq the regions with open chromatin can be further analyzed, to see with transcription factors might bind there. This can give important information on which signaling pathways drive gene expression in different cells.
 - Looking for enriched motifs
 - Cross-referencing open chromatin regions against public ChIP-seq data on different TFs.
- This can be done for each cell or cluster of cells



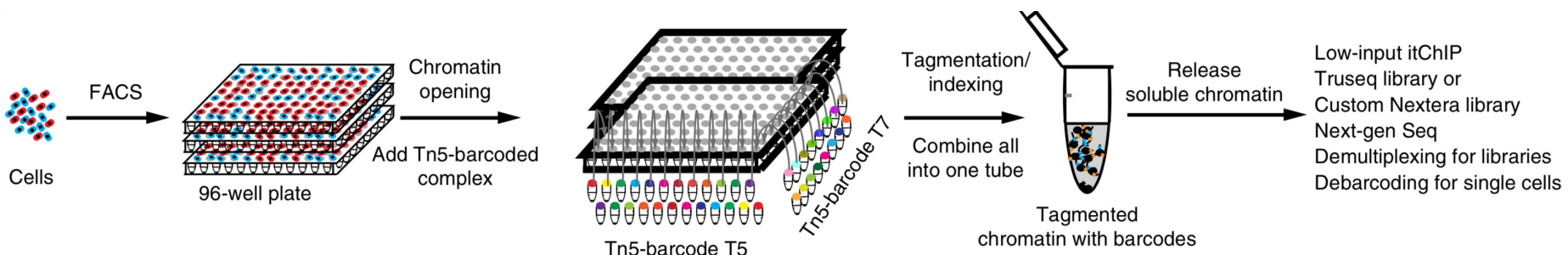
8. Differential DNA accessibility

- It's often interesting to know which chromatin regions differ in accessibility between cell types etc.
- This is a similar problem to differential gene expression (for RNA-seq data) and differential binding (for ChIP-seq data).
- But single cell ATAC-seq data have special properties that make the analysis different from bulk analysis.
 - Sparse data (low read counts, most entries are 0)
 - Many replicates/cells.
- Examples of methods:
 - Logistic regression
 - Negative binomial generalized linear model

- **ATAC Cell Ranger**
 - Computational pipeline from 10X genomics, does (more or less) all of the analysis steps described here
- **Seurat/Signac**
 - R packages originally developed for single cell RNA-seq: Filtering cells, normalization, clustering, visualization, differential DNA accessibility. Data integration.
- **ArchR**
 - R package for processing and analyzing single-cell ATAC-seq data
- **episcanpy**
 - Python package, originally developed for single cell RNA-seq. Similar functionality to Seurat/Signac
- **ChromVar**
 - R package, mostly useful for motif analysis. (Can do clustering, visualization, differential DNA accessibility too..)
- **Giggle**
 - Command line tool for cross-refencing genomic regions against public data sets.

Single cell ChIP-seq like methods

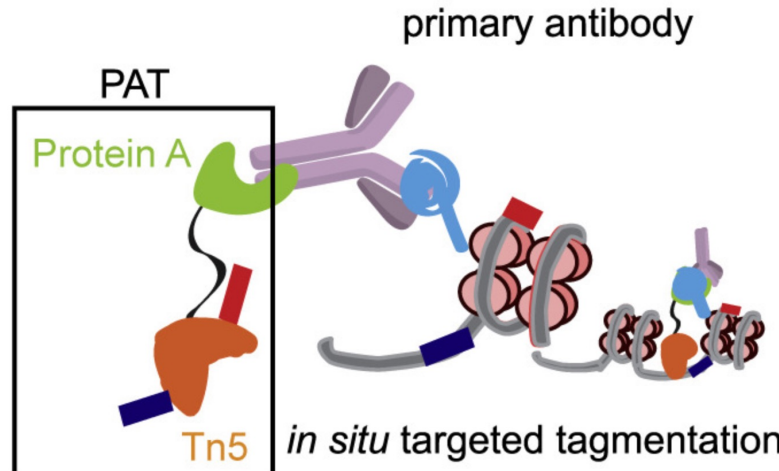
- ChIP on single cells, e.g. using droplets, is hard.
 - (Rotem et al. 2015, Nature Biotechnology) had around 800 reads/cell. Still enough to distinguish different cell types.
 - (Grosselin et al. 2019, Nature Genetics) had around 1600 reads/cell.
- Tagmentation based methods:
 - (Ai et al. 2019 Nature Cell Biology) came up with sc-itChIP-seq (single cell indexing and tagmentation ChIP-seq):
 - First use Tn5 transposase to add cell barcodes to DNA.
 - Then do ChIP in bulk.
 - 9000 reads / cell.
 - 96 well plates



(Ai et al. 2019 Nature Cell Biology)

Single cell ChIP-seq like methods II

- ChIP-free methods:
 - (Wang et al. 2019, Molecular Cell) CoBATCH
 - Antibody binds to protein of interest. → This recruits PAT complex with Tn5 → Tagmentation of DNA near protein of interest.
 - 12000 reads/cell
 - Combinatorial indexing (like for sci-ATAC-seq)
 - Quite simple protocol, no ChIP
 - (Kaya-Okur et al. 2019, Nature Communications) CUT&Tag, similar idea. (Used nanowells instead of combinatorial indexing.)



**nature
biotechnology**

ARTICLES

<https://doi.org/10.1038/s41587-021-00869-9>



Single-cell CUT&Tag profiles histone modifications and transcription factors in complex tissues

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In contrast to single-cell approaches for measuring gene expression and DNA accessibility, single-cell methods for analyzing histone modifications are limited by low sensitivity and throughput. Here, we combine the CUT&Tag technology, developed to measure bulk histone modifications, with droplet-based single-cell library preparation to produce high-quality single-cell data on chromatin modifications. We apply single-cell CUT&Tag (scCUT&Tag) to tens of thousands of cells of the mouse central nervous system and probe histone modifications characteristic of active promoters, enhancers and gene bodies (H3K4me3, H3K27ac and H3K36me3) and inactive regions (H3K27me3). These scCUT&Tag profiles were sufficient to determine cell identity and deconvolute regulatory principles such as promoter bivalency, spreading of H3K4me3 and promoter-enhancer connectivity. We also used scCUT&Tag to investigate the single-cell chromatin occupancy of transcription factor OLIG2 and the cohesin complex component RAD21. Our results indicate that analysis of histone modifications and transcription factor occupancy at single-cell resolution provides unique insights into epigenomic landscapes in the central nervous system.

- Data analysis for all of these methods is similar to single cell ATAC-seq.
- Single cell DNA-protein interaction studies, i.e. “ChIP-seq like” is still new, but developing fast. Throughput will likely increase a lot.

Single cell DNA methylation

- Methods
 - Whole genome vs reduced representation/targeted
 - Bisulphite vs bisulphite-free (methylation-sensitive restriction enzymes)
- Quite hard and expensive
- Data
 - Mostly 5mC
 - Thousands of cells
 - $10^4 - 10^7$ CpGs per cell
 - Not the same CpGs in all cells.
- Analysis still hard

Article

DNA methylation atlas of the mouse brain at single-cell resolution

<https://doi.org/10.1038/s41586-020-03182-8>

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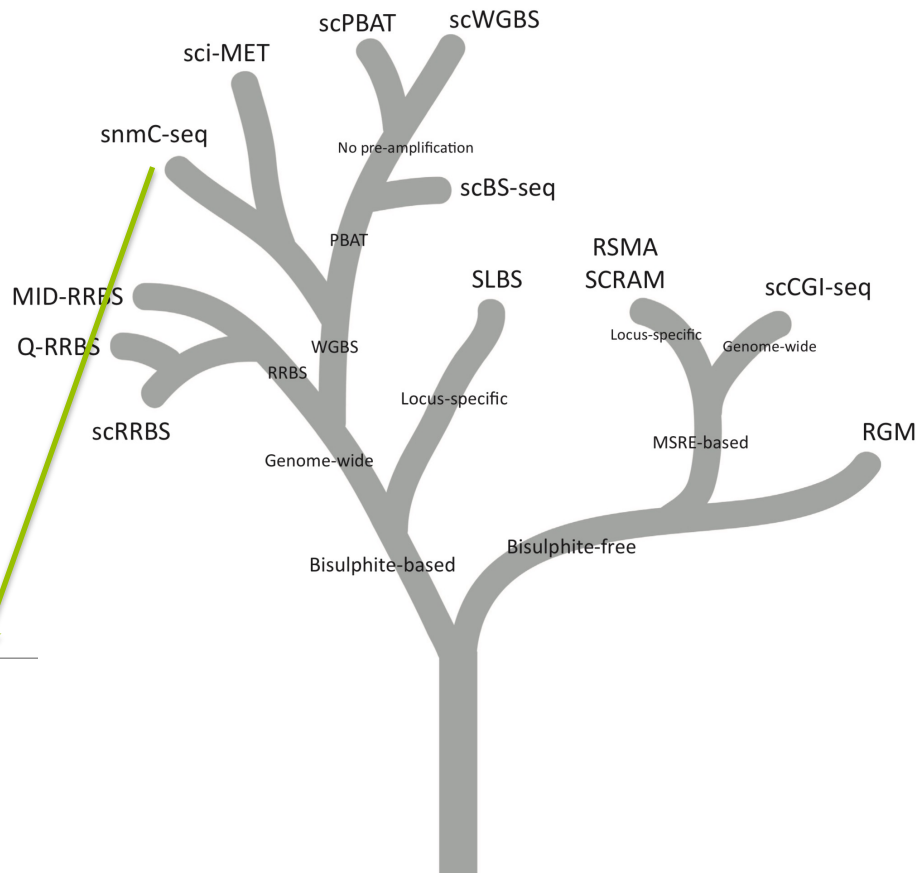
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Open access

Hanqing Liu^{1,2,16}, Jingtian Zhou^{1,3,16}, Wei Tian¹, Chongyuan Luo^{1,4}, Anna Bartlett¹, Andrew Aldridge¹, Jacinta Lucero⁵, Julia K. Osteen³, Joseph R. Nery¹, Huaming Chen¹, Angeline Rivkin¹, Rosa G. Castanon¹, Ben Clock⁶, Yang Eric Li⁷, Xiaomeng Hou^{8,9,10,11}, Olivier B. Poirion^{8,9,10,11}, Sebastian Preissl^{8,9,10,11}, Antonio Pinto-Duarte⁵, Carolyn O'Connor¹², Lara Boggeman¹², Conor Fitzpatrick¹², Michael Nunn¹, Eran A. Mukamel¹³, Zhuzhu Zhang¹, Edward M. Callaway¹⁴, Bing Ren^{2,8,9,10,11}, Jesse R. Dixon⁶, M. Margarita Behrens⁵ & Joseph R. Ecker^{1,15,16}

>100K cells



Karemaker & Vermeulen. 2018,
Trends in Biotechnology

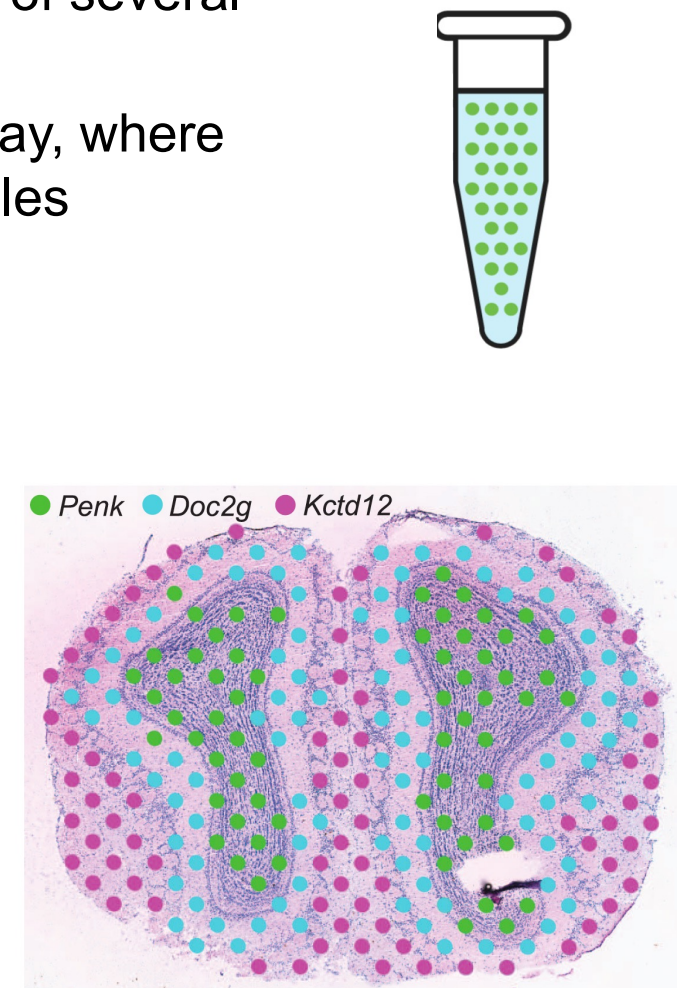
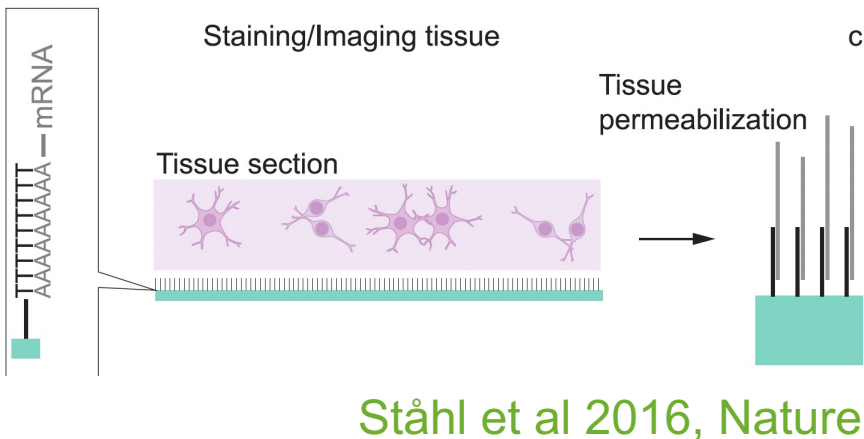
Combining assays from the same cells

- Many methods combine several assays from the same cells, e.g.
 - scRNA-seq and scATAC-seq (10X genomics, SNARE-seq, and many more)
 - scRNA-seq and sc-protein abundance (CITE-seq)
 - scRNA-seq and scDNA methylation
 - scRNA-seq and scDNA methylation and sc nucleosome (scNMT-seq)
 - scRNA-seq, scATAC-seq, sc-protein abundance and clonal info from mitochondrial DNA (DOGMA-seq)

Method	Molecular layers profiled				Throughput (low/medium/ high)	Special features (compared to techniques from same category)	Format	References				
	Epigenome		Genome	Transcriptome								
	Chromatin accessibility	Chromatin conformation										
scCAT-seq	x			x	+	usable fragments	well	Liu et al. (2019)				
Paired-seq	x			x	+++	throughput	well	Zhu et al. (2019)				
sc(ATAC + RNA)-seq	x			x	+	cost; simple workflow	well	Reyes et al. (2019a)				
sci-CAR	x			x	+++	acc. & RNA intersect coverage	well	Cao et al. (2018)				
SNARE-seq	x			x	+++	sensitivity	droplet	Chen et al. (2019)				
ASTAR-seq	x			x	++	price-performance ratio	microfluidics	Xing et al. (2020)				
SHARE-seq	x			x	+++	throughput, performance	well	Ma Sai, et al. (2020)				
ISSAAC-seq	x			x	+++	throughput, performance (esp. ATAC)	well/droplet	Xu et al. (2022)				
scDam&T-seq		x		x	+	protein-DNA interactions information	well	Rooijers et al. (2019)				
scNOMe-seq	x	x			+	estimates nucleosome phasing	well	Pott, (2017)				
scCOOL-seq	x	x	x		+	acc. & DNAme intersect coverage	well	Guo et al. (2017)				
iscCOOL-seq	x	x	x		++	accessibility coverage	well	Gu et al. (2019a)				
scMethyl-HIC		x	x		+	mapping rate	well	Li et al. (2019)				
sn-m3C-seq		x	x		+++	DNAme coverage	well	Lee et al. (2019)				
scNMT-seq	x		x	x	++	throughput	well	Clark et al. (2018)				
scNOMeRe-seq	x		x	x	+	DNAme coverage	well	Wang et al. (2021)				
scSIDR-seq			x	x	+	captures total RNA	well	Han et al. (2018)				
TARGET-seq			x	x	+++	cost; throughput	well	Rodriguez-Meira et al. (2019)				
RETrace		x	x		+	captures microsatellites	well	Wei and Zhang, (2020)				
scTrio-seq2		x	x	x	++	DNAme coverage	well	Bian et al. (2018)				

Spatial methods

- Sequencing based methods “spatial transcriptomics”
 - + Captures all genes
 - not single cell resolution (each spot consists of several cells)
- The basic idea is to hybridize a tissue to an array, where spatial barcodes are added to the RNA molecules

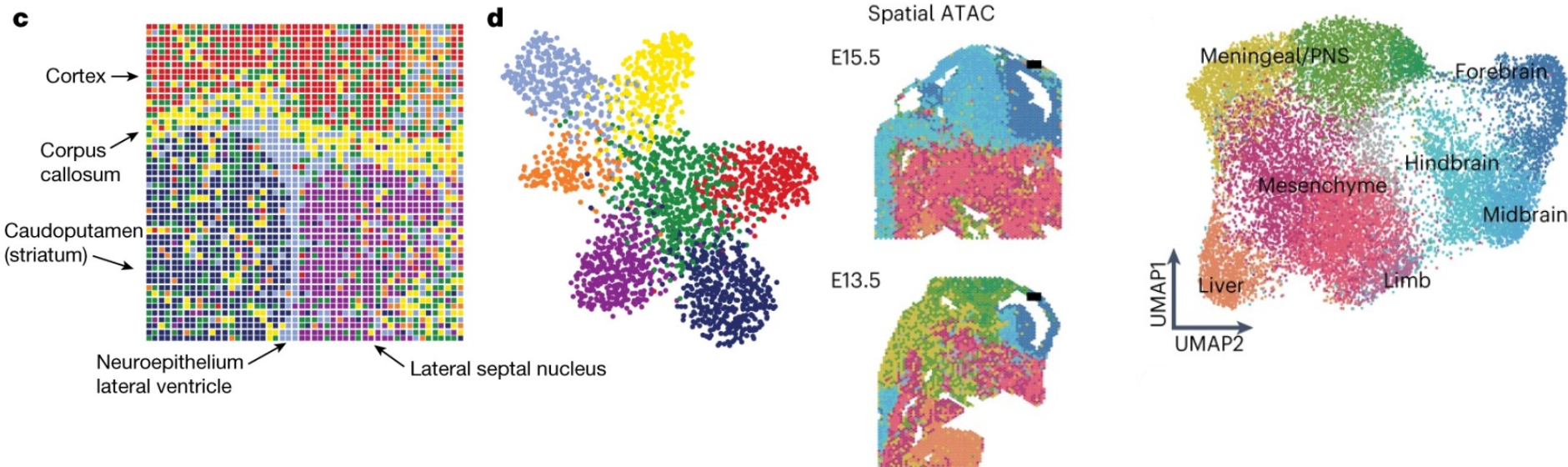


Spatial methods, II

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- For sequencing based methods, a lot of the analysis is similar to single cell data, but instead of cells, you work with “spots” (these typically cover several cells).
 - Spatial methods are often used together with single cell methods, to get information both on what different cell types looks like (from the single cell data) and where they are located (from the spatial data)

Spatial methods, III

Recently, spatial methods have been adapted to ATAC-seq.

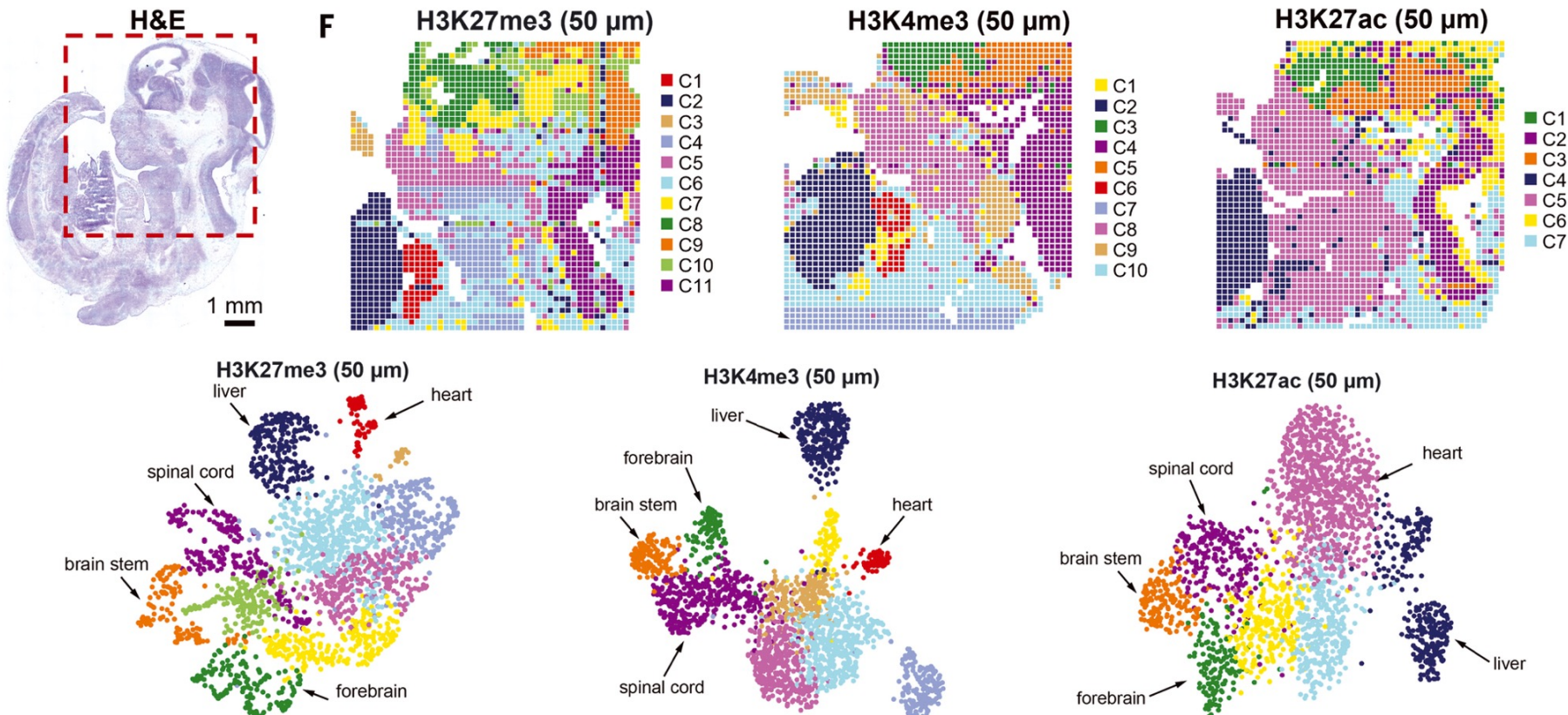


Deng et al 2022, Nature

Llorens-Bobadilla et al 2023, Nature Biotech.

Spatial methods, IV

Spatial methods have also been adapted to CUT & Tag.



Deng et al 2022, Science

-
- Single cell ATAC-seq
 - Usually works quite well
 - Commercial kits available
 - You will have a look at the data analysis in the exercise.
 - Single cell DNA methylation
 - A lot of development happening
 - Useful methods will become more widely available (already scWGBS/splat-seq at NGI).
 - Spatial Methods
 - Well established for transcriptomics
 - Adaptations for epigenomics are being developed, but still not widely used.