3000788 Intro to Comp Molec Biol

Lecture 13: Single-cell transcriptomics

September 27, 2022

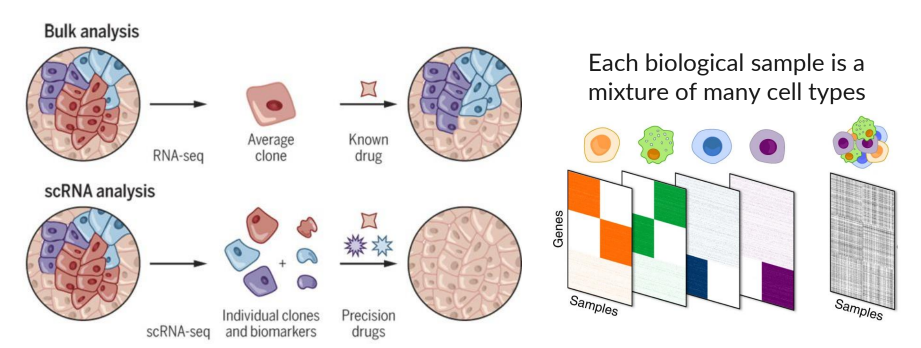


Sira Sriswasdi, PhD

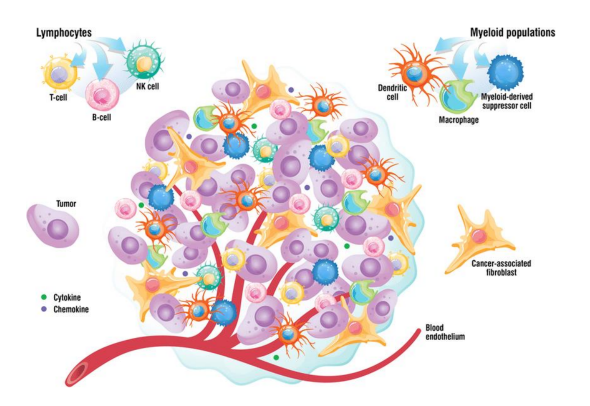
- Research Affairs
- Center of Excellence in Computational Molecular Biology (CMB)
- Center for Artificial Intelligence in Medicine (CU-AIM)

Why single-cell?

Tissue consists of multiple cell types



Tumor microenvironment



Zhang, J. and Veeramachaneni, N. Biomarker Research 10:5 (2022)

Cancer stem cell

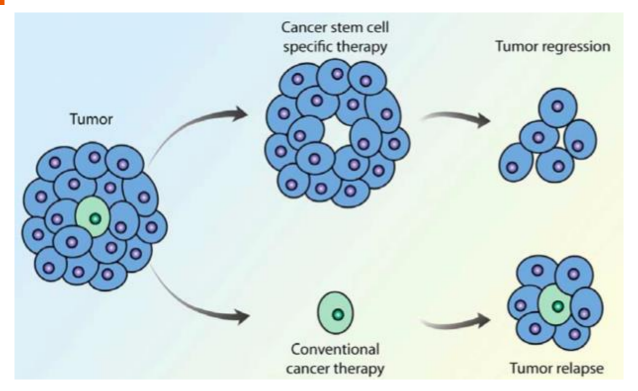
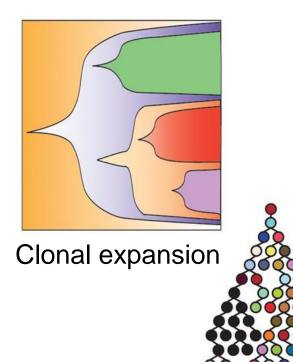


Image from https://hsci.harvard.edu/stem-cells-and-cancer

Knowledge at single-cell resolution

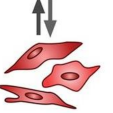
Heterogeneity



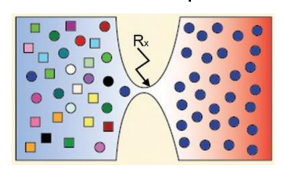




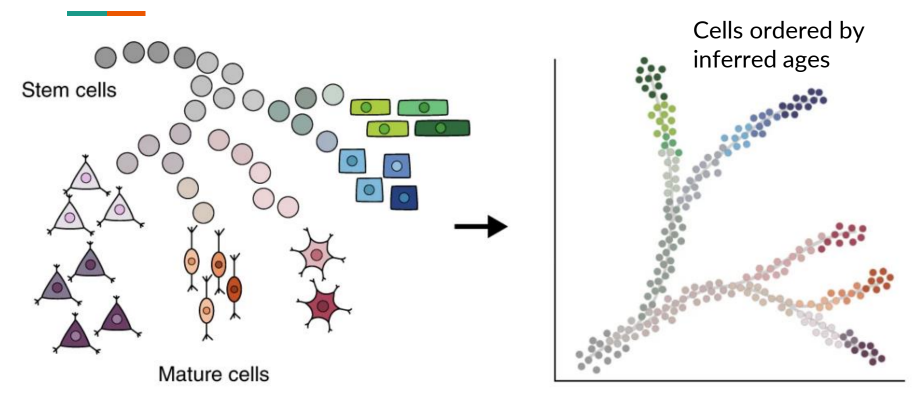




Treatment response

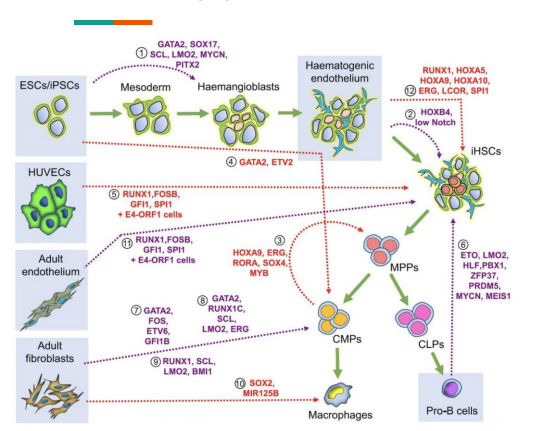


Cell development through single-cell data



Source: Moon et al. Nature Biotechnology 37:1482-92 (2019)

Detecting gene switches

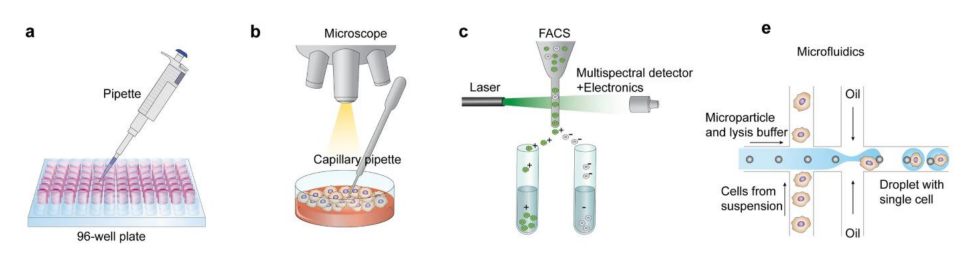


- Genes whose expressions:
- Change over inferred developmental time
- Diverge between two developmental branches
- Turn on/off across cell type

Source: Ivanovs et al. Development 144:2323-37 (2017)

Single-cell vs bulk transcriptomics

Cell isolation techniques



Hwang et al. Exp & Mol Med 50:96 (2018)

Diverse protocol choices

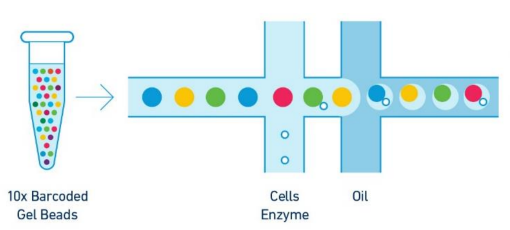
Methods	Transcript coverage	UMI possibility	Strand specific	References
Tang method	Nearly full-length	No	No	Tang et al., 2009
Quartz-Seq	Full-length	No	No	Sasagawa et al., 2013
SUPeR-seq	Full-length	No	No	Fan X. et al., 2015
Smart-seq	Full-length	No	No	Ramskold et al., 2012
Smart-seq2	Full-length	No	No	Picelli et al., 2013
MATQ-seq	Full-length	Yes	Yes	Sheng et al., 2017
STRT-seq and STRT/C1	5'-only	Yes	Yes	Islam et al., 2011, 2012

Chen et al. Front Genet. 10:317 (2019)

Quartz-Seq2	3'-only	Yes	Yes	Sasagawa et al., 2018
DroNC-seq	3'-only	Yes	Yes	Habib et al., 2017
Seq-Well	3'-only	Yes	Yes	Gierahn et al., 2017
sci-RNA-seq	3'-only	Yes	Yes	Cao et al., 2017
SPLiT-seq	3'-only	Yes	Yes	Rosenberg et al., 2018
Chromium	3'-only	Yes	Yes	Zheng et al., 2017
InDrop	3'-only	Yes	Yes	Klein et al., 2015
Drop-seq	3'-only	Yes	Yes	Macosko et al., 2015
CytoSeq	3'-only	Yes	Yes	Fan H.C. et al., 2015
MARS-seq	3'-only	Yes	Yes	Jaitin et al., 2014
CEL-seq2	3'-only	Yes	Yes	Hashimshony et al., 2016
CEL-seq	3'-only	Yes	Yes	Hashimshony et al., 2012

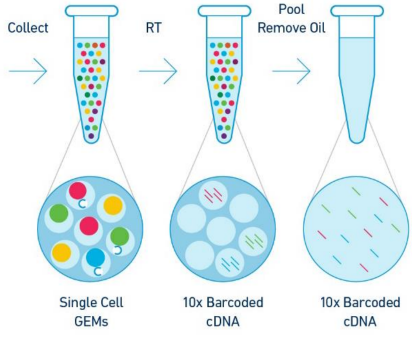
- Mostly sequence 3' ends of transcript
- Unique Molecular Identifier (UMI) = PCR barcode

10x Genomics' chromium technique

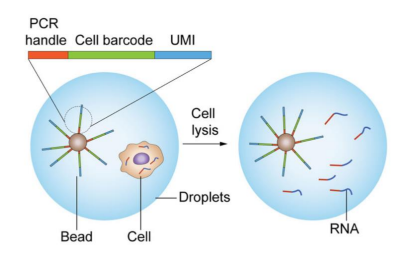


https://bauercore.fas.harvard.edu/10x-chromium-system

- Droplets of sequencing reagents
- Cell barcode + pooled sequencing



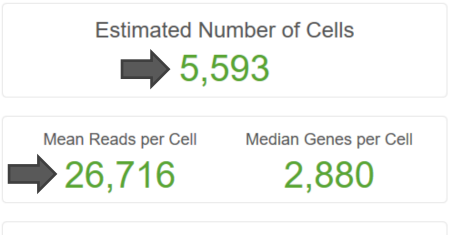
UMI and cell barcode



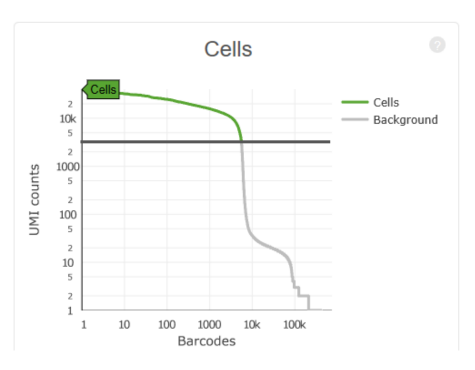
Hwang et al. Exp & Mol Med 50:96 (2018)

- All beads in each droplet have the same cell barcode
 - Reads with the same barcode came from the same cell
- Each PCR adapter contains different
 Unique Molecular Identifiers (UMI)
 - Reads with the same UMI came from the same original RNA molecule

Single-cell vs bulk data

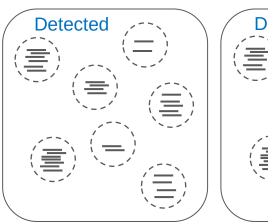


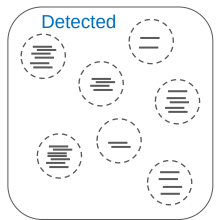




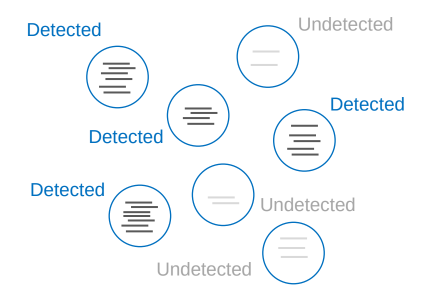
Detection limit of single-cell data







Single-cell RNA-seq



Challenges in single-cell data analysis

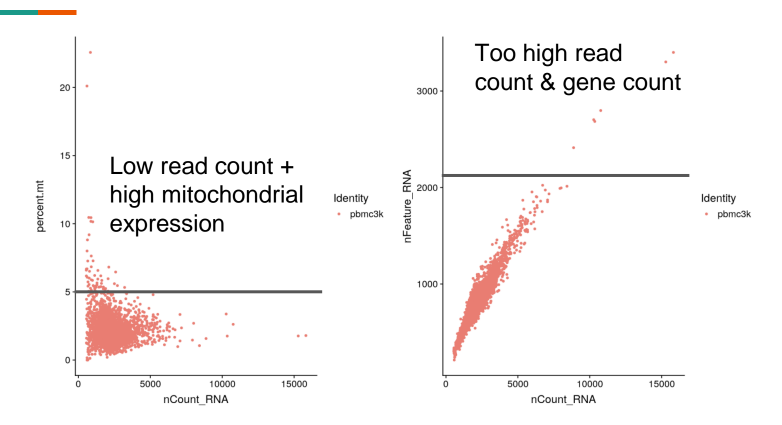
- Low read count per cell and gene
 - A lot of zeros in expression data
- Cells are biologically different
 - High variance across cells
- Cells are in continuous states of development
 - Not just control vs treatment
- Data is very large (256 GB of RAM for medium project)
 - 10,000 cells x 5,000 genes

Single-cell data preprocessing

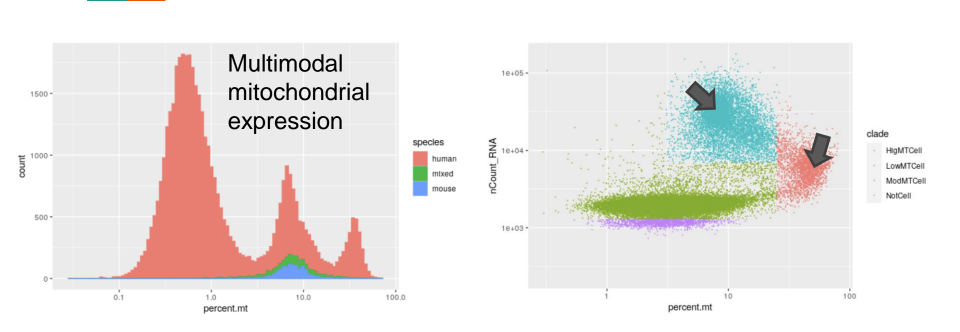
Key steps in single-cell data processing

- Quality filter
 - Low read count & gene count = non-cells
 - Very high read count & gene count = multi-cells
 - High mitochondrial expression = dead cells
- Within-sample normalization
 - Dealing with missing expression values
- Multi-sample integration
 - Single-cell data have strong batch effects

Basic quality filters

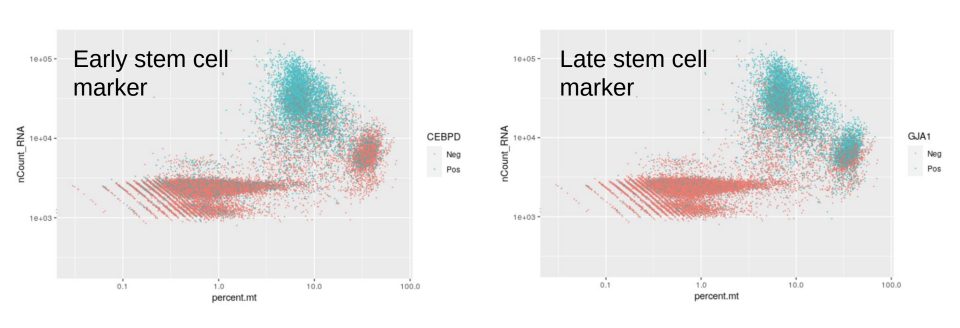


An exception

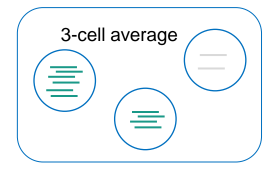


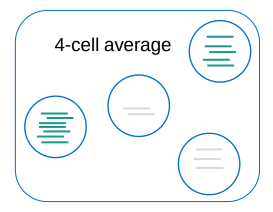
High mitochondrial activity in stem cells and some cell types

Stem cell markers in high-mitochondrial cells



Normalization with pooling





Low expression

Gene A Gene B

Gene C

Medium expression

Gene D

Gene E

Gene F

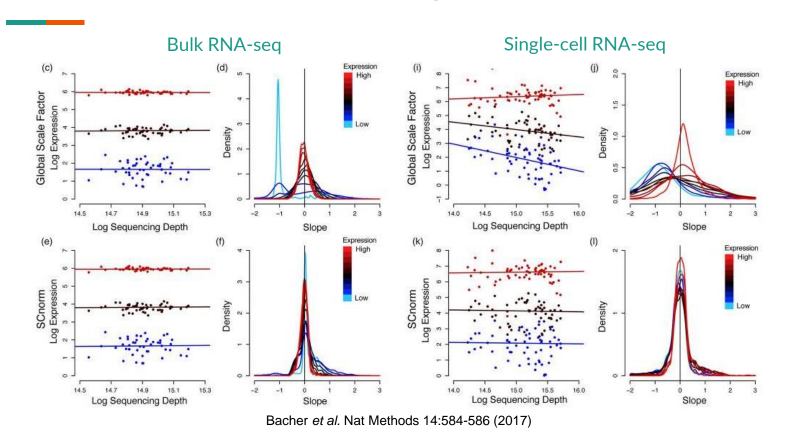
Gene G

High expression

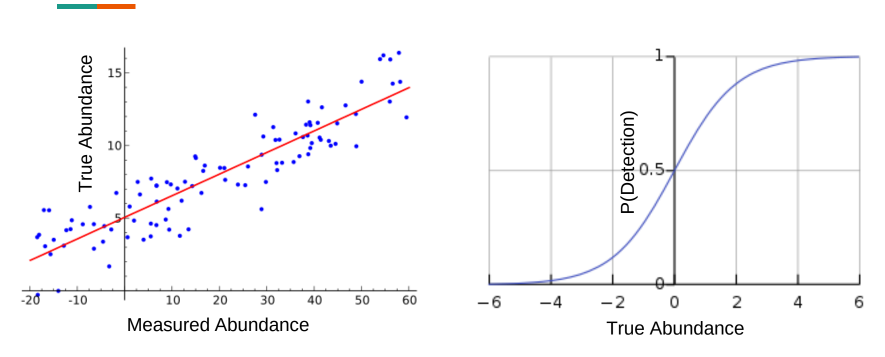
Gene H

Gene I

Expression-dependent scaling factor



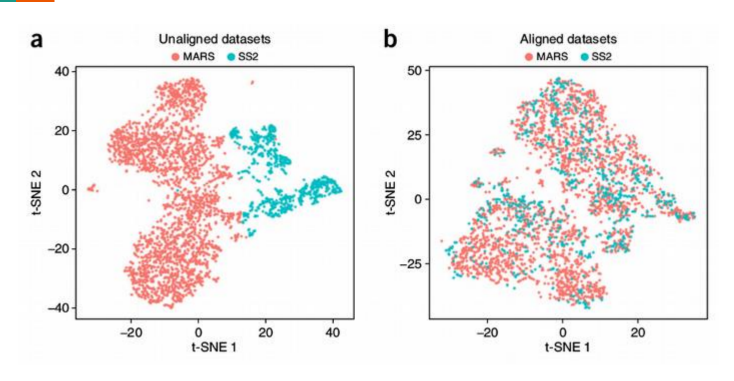
Modeling of detection probability



Measured abundance = $f(true abundance) \times P(detection | true abundance)$

Single-cell data integration

High bias across datasets



https://towardsdatascience.com/how-to-batch-correct-single-cell-7bad210c7ae1

Linear effect removal (Combat-Seq)

Negative binomial regression models

Gene-wise model: for a certain gene g, count in sample j from batch i $y_{gij} \sim NB(\mu_{gij}, \phi_{gi})$

parameter γ_{ai}

Explicit addition of batch
$$\log \mu_{gij} = \alpha_g + X_j \beta_g + \gamma_{gi} + \log N_j$$

parameter $\gamma_{g,j}$ $Var(y_{gij}) = \mu_{gij} + \phi_{gi} \mu_{gij}^2$

Decompose scaled counts into 3 components
$$\begin{bmatrix} & \alpha_g \\ & X_j\beta_g \\ & & \\ & \gamma_{gi} \end{bmatrix}$$

$$lpha_g$$
 Average level for gene g (in "negative" samples) $X_j eta_g$ Biological condition of sample j

Mean batch effect

 N_i = total read count for sample j

Dispersion batch effect

Estimate batch effect parameters

Estimate parameters using established methods in edgeR

Calculate "batch-free" distributions

We assume the adjusted data also follow a negative binomial distribution: $y_{qj}^* \sim NB(\mu_{qj}^*, \phi_q^*)$

Subtract batch parameter $\gamma_{g,i}$ \Longrightarrow $\log \mu_{gj}^* = \log \hat{\mu}_{gij} - \hat{\gamma}_{gi}$ $\phi_g^* = \frac{1}{N_{batch}} \sum_i \hat{\phi}_{gi}$

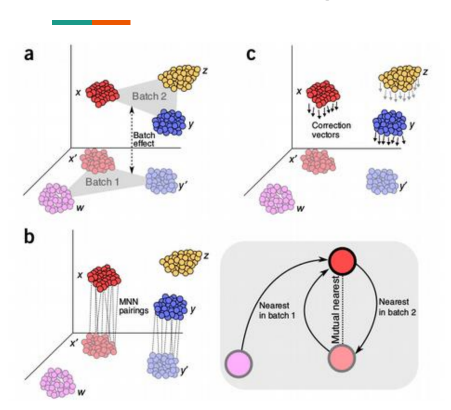


$$\log \mu_{gj}^* = \log \hat{\mu}_{gij} - \hat{\gamma}_g$$

$$\phi_g^* = \frac{1}{N_{batch}} \sum_i \hat{\phi}_{gi}$$

Zhang, Y. et al. NAR Genom and Bioinfo 2:lgaa078 (2020)

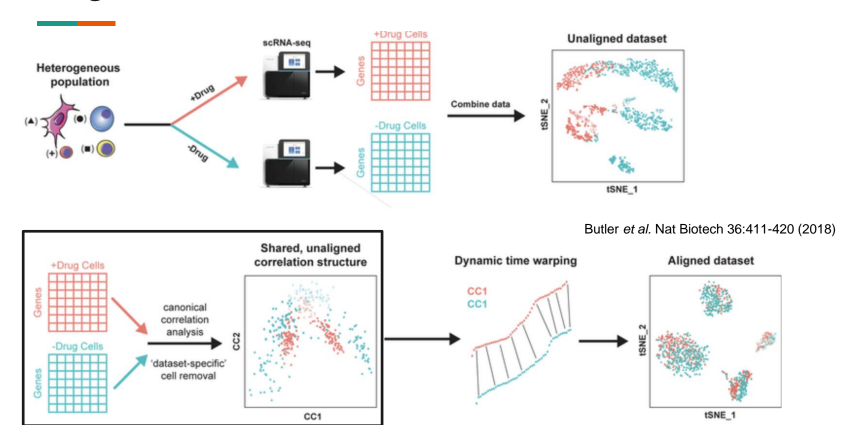
Mutual nearest neighbor (MNN)



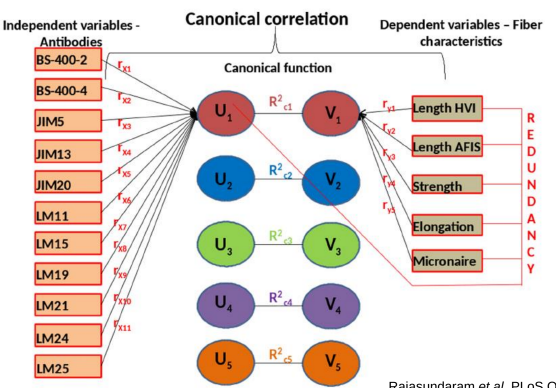
- Map clusters of cells together rather than individual cells
- Similar to reciprocal best hits from BLAST for identifying orthologs
- Apply the average mapping vector to unique cell types

Haghverdi et al., Nat. Biot. 36, 2018

Integration via canonical correlation



Canonical correlation analysis (CCA)



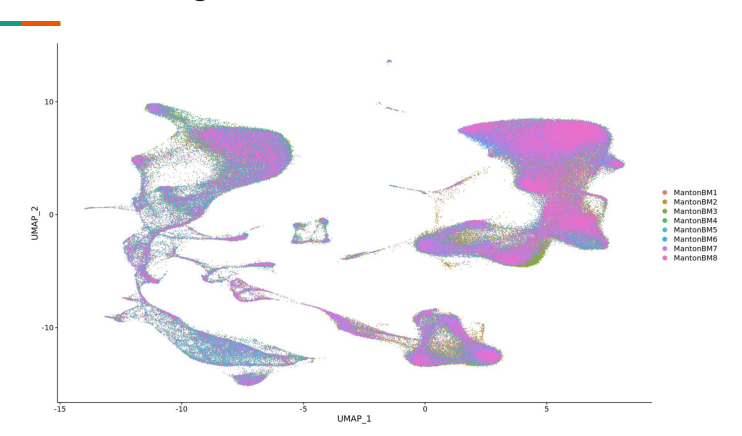
- Same samples with two different systems of observations
- Identify correlation structure observation systems (features)

Rajasundaram et al. PLoS ONE (2014)

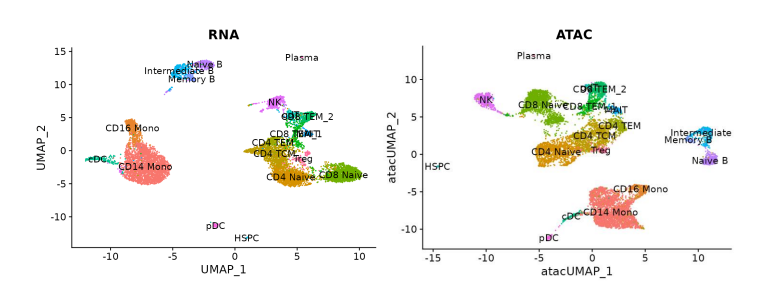
CCA for single-cell data

- Two single-cell experiments contain different cells (samples) but the same genes (features)
- CCA for single-cell data treats cells as feature describing the genes
 - Identify correlation structure between cells
 - **Expectation**: gene expressions of the same cell type will be most correlated
- Follow up by MNN-styled technique

Good data integration



Integrating RNA-seq with ATAC-seq

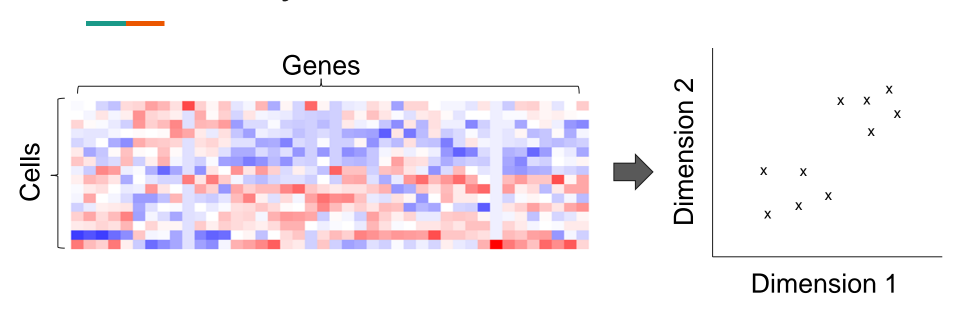


- ATAC-seq = open chromatin ~ gene expression level
- Transfer cell type label

Visualizing single-cell data

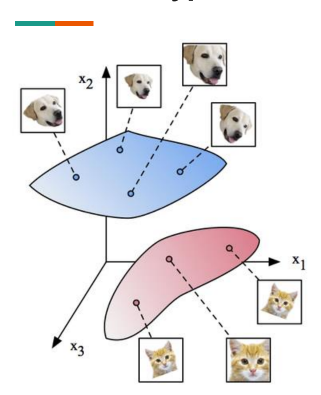
with dimensionality reduction techniques

Dimensionality reduction

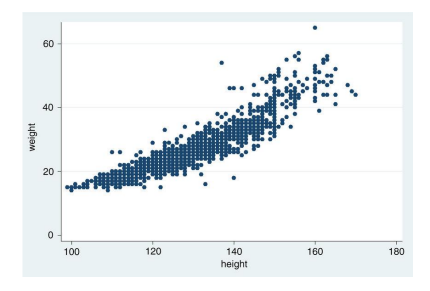


Collapse high-dimensional data on to 2D or 3D scatter plot that preserve some information in original dimension

Manifold hypothesis

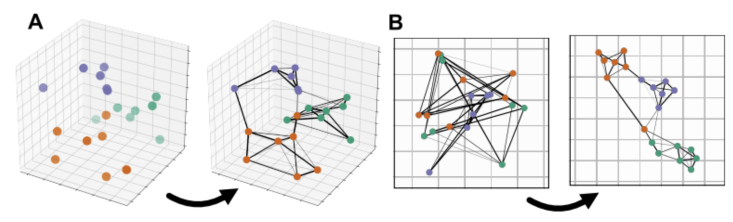


"Real-world, high-dimensional data lie on some low-dimensional manifolds"



Nordin, P. et al. Global Health Action 7:25351 (2014)

Dimensionality reduction algorithm sketch



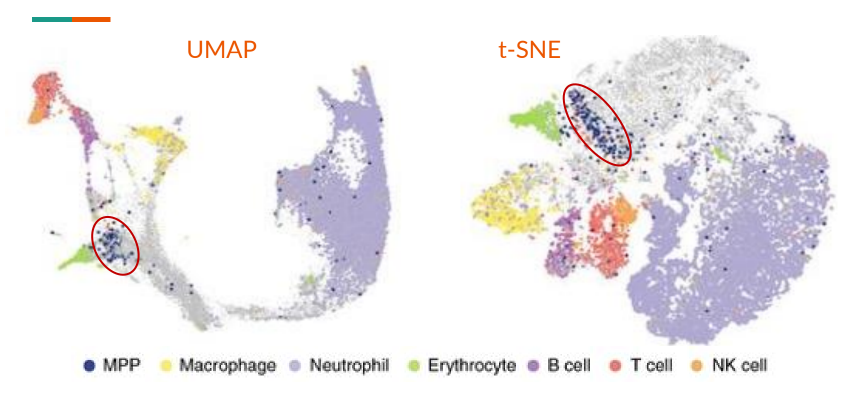
Step 1: Compute a graphical representation of the dataset

Sainburg, T. et al., Neural Comput 33(11):2881-2907 (2021)

Step 2 (non-parametric): Learn an embedding that preserves the structure of the graph

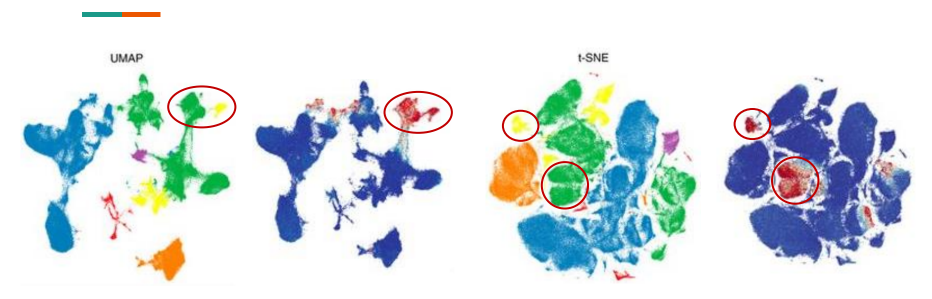
Similarity = correlation in gene expression across cells

t-SNE vs UMAP on single-cell data



Becht, E. et al. Nature Biotechnology 37:38-44 (2019)

t-SNE vs UMAP on single-cell data

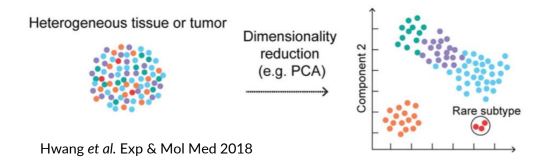


Becht, E. et al. Nature Biotechnology 37:38-44 (2019)

- Both are equally good at detecting individual cell types
- But UMAP is better at capturing transitions across cell types

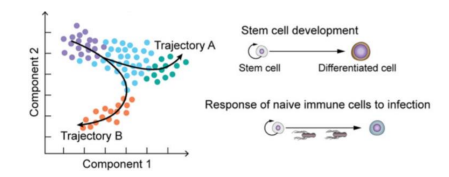
Single-cell analysis overview

Cell clustering and trajectory reconstryction

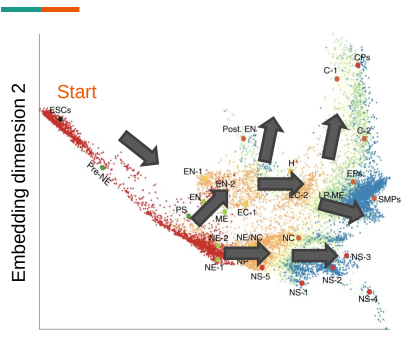


Clustering of cells with similar omics signatures reveal groups of different cell types and developmental stages

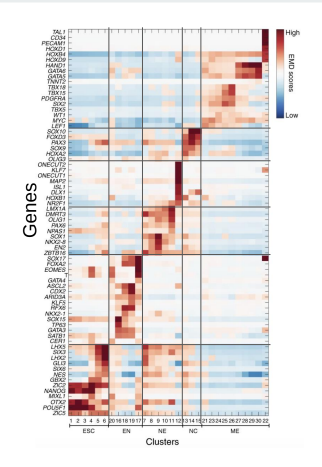
Trajectory modeling with random walk, diffusion, or Markov chain reconstruct the paths of cell development



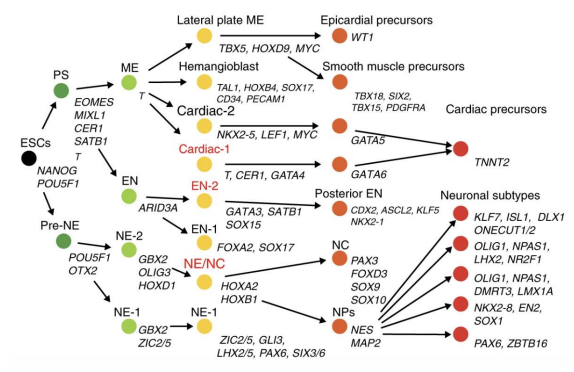
An end-to-end example



Embedding dimension 1

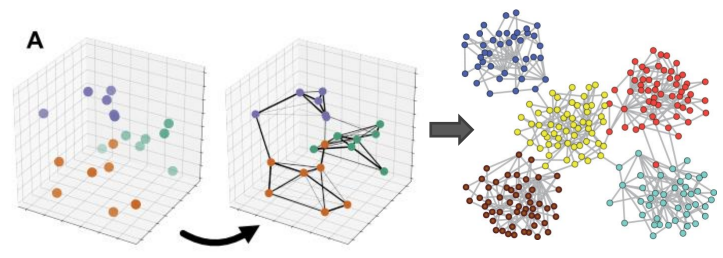


An end-to-end example



Source: Moon et al. Nature Biotechnology 37:1482-92 (2019)

Algorithm sketch for cell type clustering



Sainburg, T. et al., Neural Comput 33(11):2881-2907 (2021)

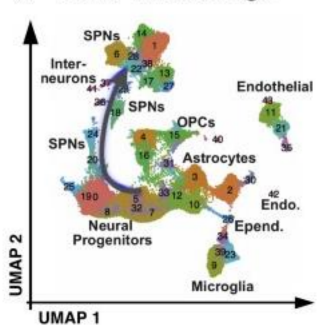
https://github.com/topics/graph-clustering

- Connect cells with similar gene expression profile
- Split network into modules with dense edges

Cell developmental trajectory

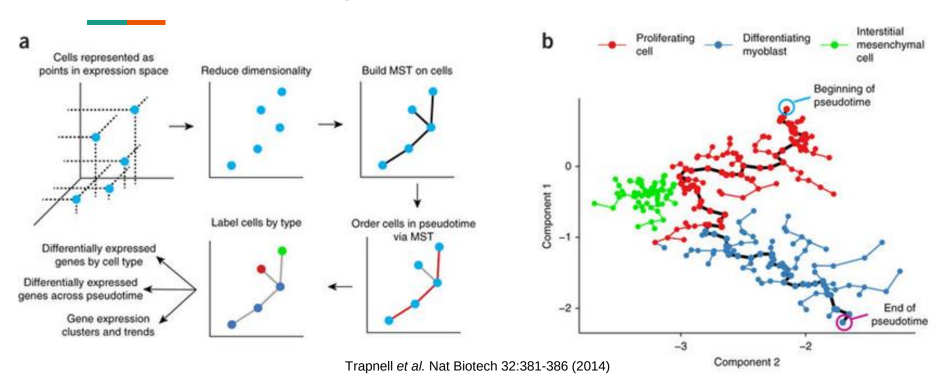
Cells can be arrange along developmental path

(b) UMAP Embeddings



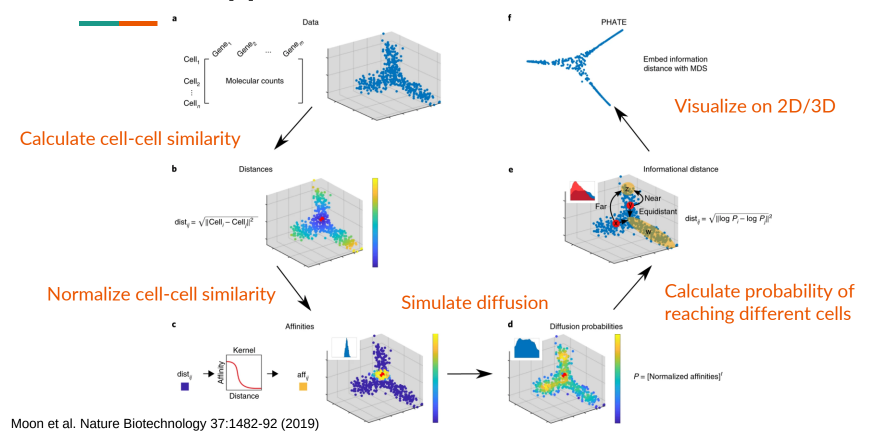
- Cells are in continuous developmental states
- Similar gene expression implies similar state
- Reconstruct pseudotime
- Identify important genes for development
 - Expression change along the trajectory
 - Expression switch at a particular time point

Minimum spanning tree

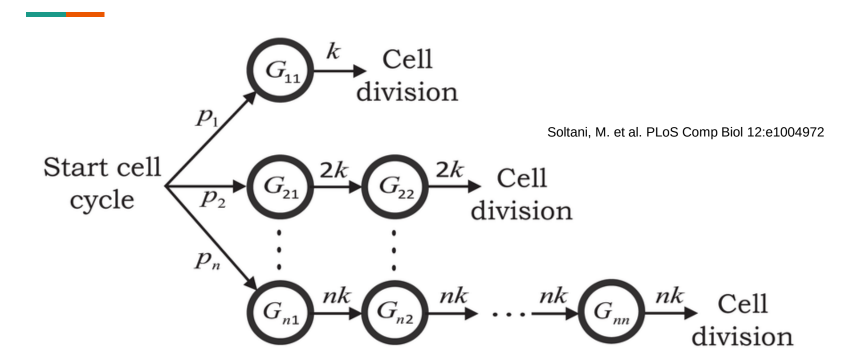


Trajectory along the most similar cells

Diffusion approach for cell-cell transitions

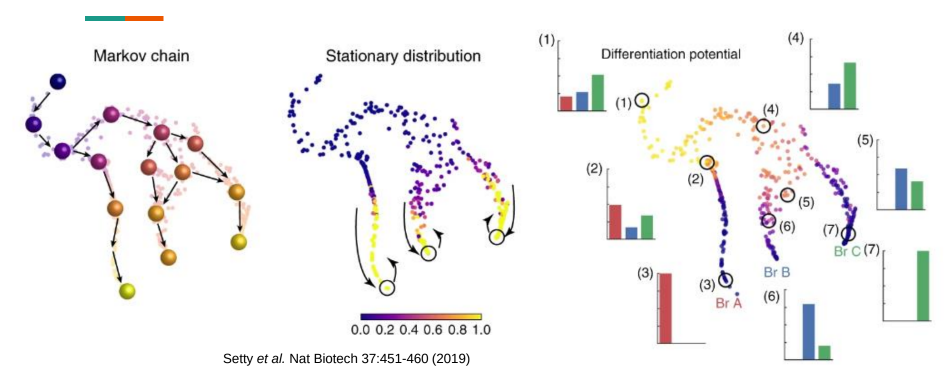


Markov chain model for cell development



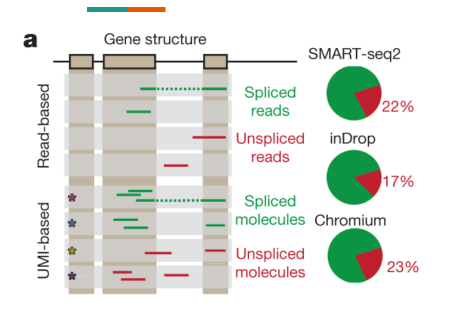
Cell state + state transition probability + dependency on previous state

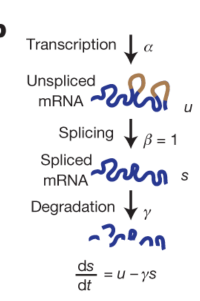
Estimating differentiation potential

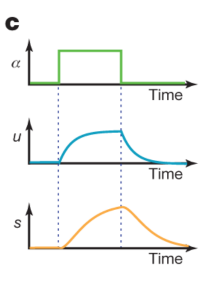


Differentiation potential = probability of reaching multiple final cell types

Dynamics of unspliced transcript



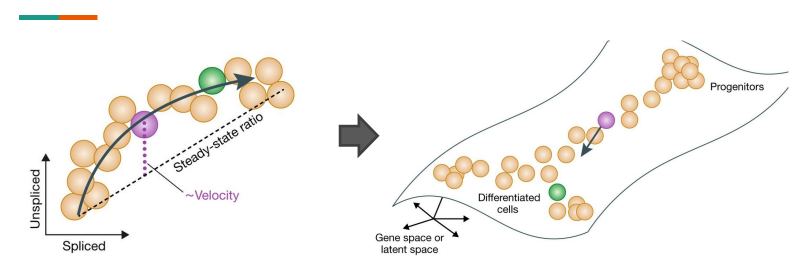




La Manno et al. Nature 2018

- When a gene is activated, level of unspliced transcripts rises first
- When a gene is repressed, level of unspliced transcripts drops first

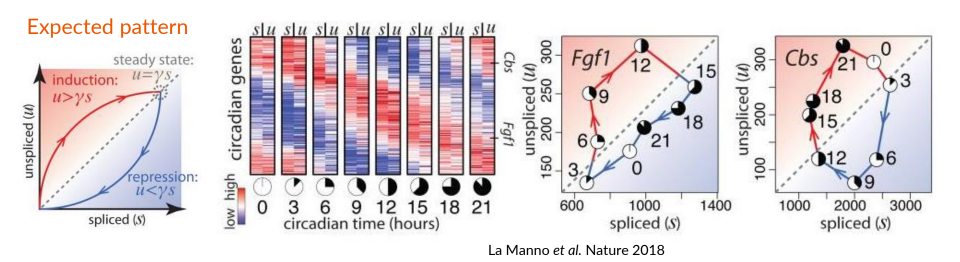
RNA velocity model



Bergen, V. et al. Mol Sys Biol 17:e10282 (2021)

- Ratio of spliced and unspliced isoforms tells gene activation state
- Compare to nearby cells to identify direction of activation or repression

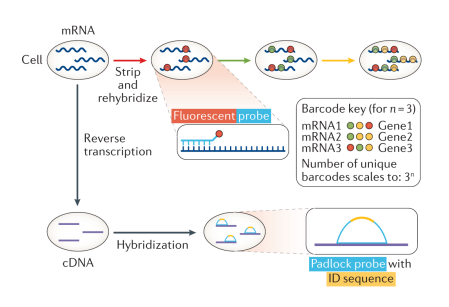
Proof of RNA velocity

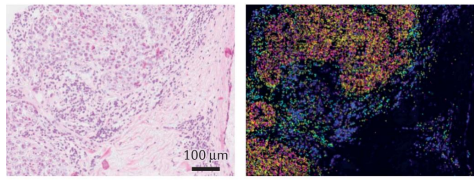


Circadian genes are genes whose expression cycle with the time of day

Spatial transcriptomics

Extended NanoString

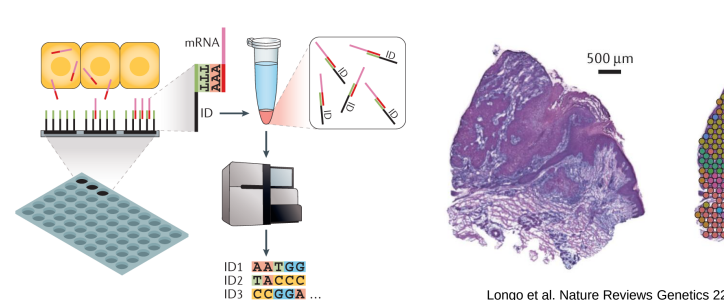


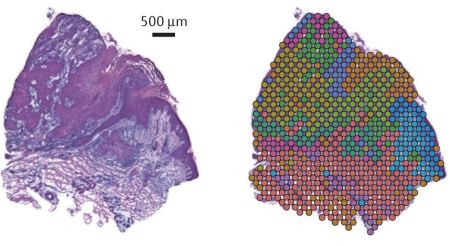


Longo et al. Nature Reviews Genetics 22:627-644 (2021)

In-situ fluorescence labeling of selected RNA transcripts

Spatial barcoding





Longo et al. Nature Reviews Genetics 22:627-644 (2021)

Spatial cell isolation + barcoding

Single-nucleus sequencing

- Isolate nuclei instead of whole-cells
 - Only capture RNA expression in nucleus
- Good for cells that are difficult to isolate: adipocyte, neuron, etc.
 - Also works well with preserved tissues

Summary

- Benefits of single-cell technology
- Difference between single-cell and bulk transcriptomics
- New analysis ideas
 - Visualization of high-dimensional data (preview)
 - Cell type clustering
 - Developmental trajectory reconstruction
- Spatial transcriptomics

Any question?

See you on Thursday September 29th