### Introduction to Single-Cell RNA-seq

Thanks to Dennis Wylie for some slides

### Why single-cell RNA-seq?

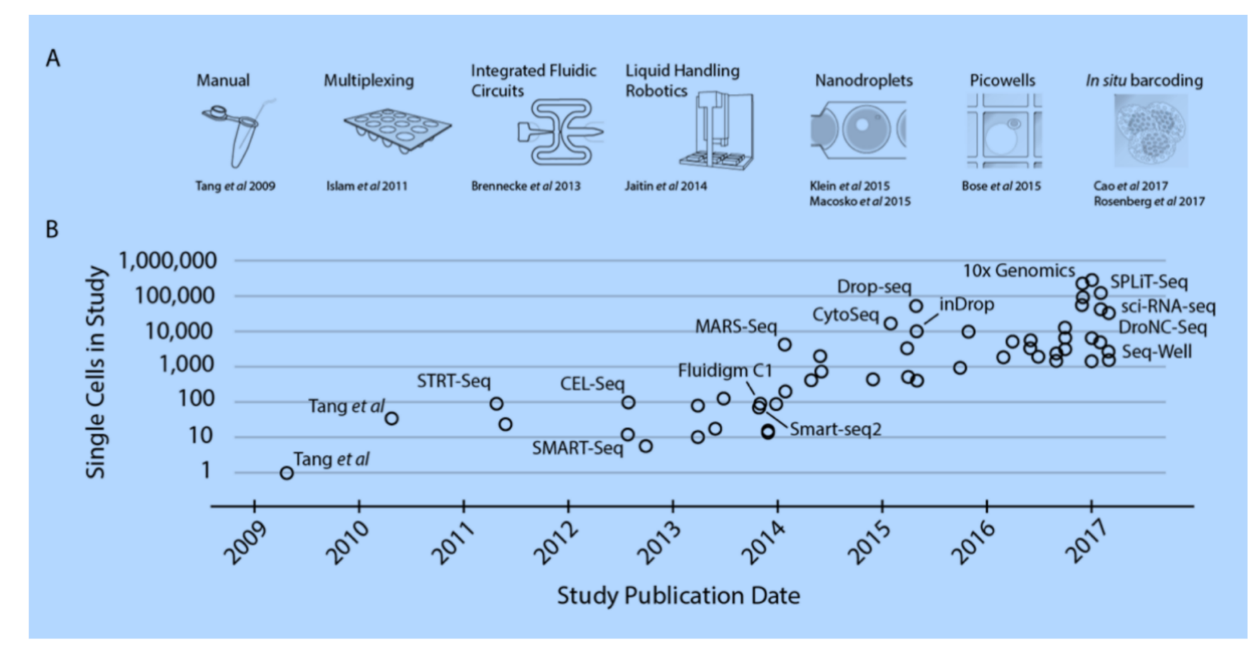
- Allows profiling of gene expression in individual cells.
  - To look at heterogeneity across cell type subpopulations

### Unique Challenges with single-cell RNA-seq

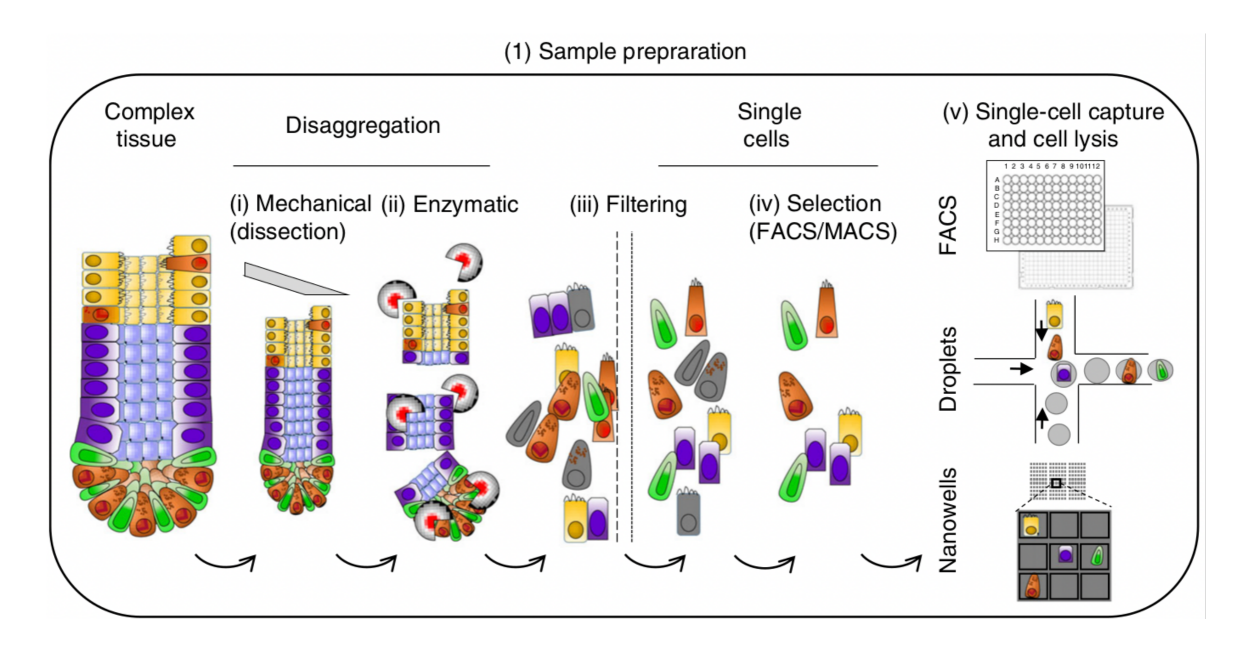
- Gene dropouts
  - Due to low amounts of RNA per cell.
  - Some cells are easier to capture than others.
- Large, but sparse gene expression matrix
  - Expression values for all genes across 10^2 to 10^5 cells.
  - Many zeros

### Single-cell RNA-seq Technology Improvements

#### https://arxiv.org/abs/1704.01379

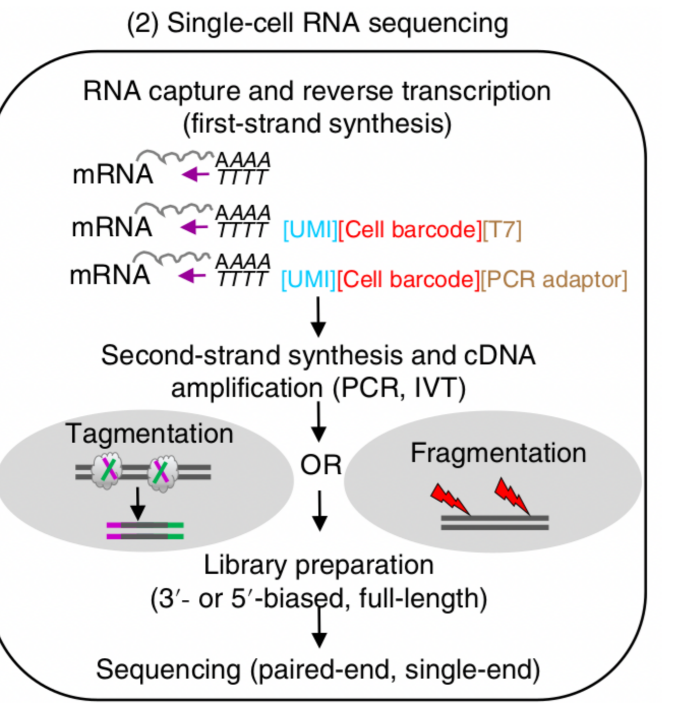


### Single-cell RNA-seq Sample Prep



Taken from Lafzi et al. (2018)

## Single-cell RNA-Seq Library Prep



<sup>•</sup>UMI (Unique molecular index)

- •Random 4-20 bp sequences attached to each RNA fragment/template to uniquely identify that RNA fragment/template.
- •One per fragment.
- •For detection of PCR duplicates.

#### •Cell barcode

- •A cell-specific sequence attached to RNA fragments.
- •One per cell
- •For differentiating by cell.

#### •Sample barcode/index

- •One per sample
- •Allows pooling multiple samples on the same sequencing run.

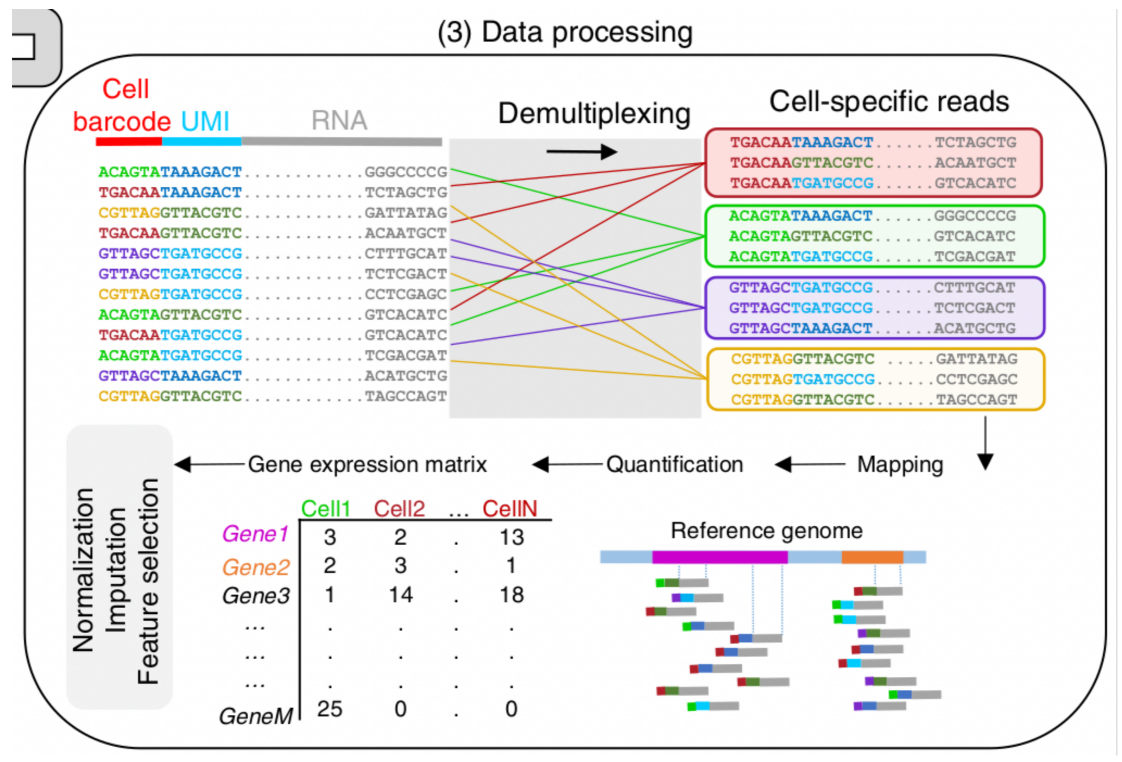
Taken from Lafzi et al. (2018)

### **Coverage Recommendations**

- How many cells per sample?
  - •~10,000 cells per 'typical' sample.
- How many reads per cell?
  - 30,000-50,000 reads per cell for 'typical' samples.
- This would take up >1 lane/sample

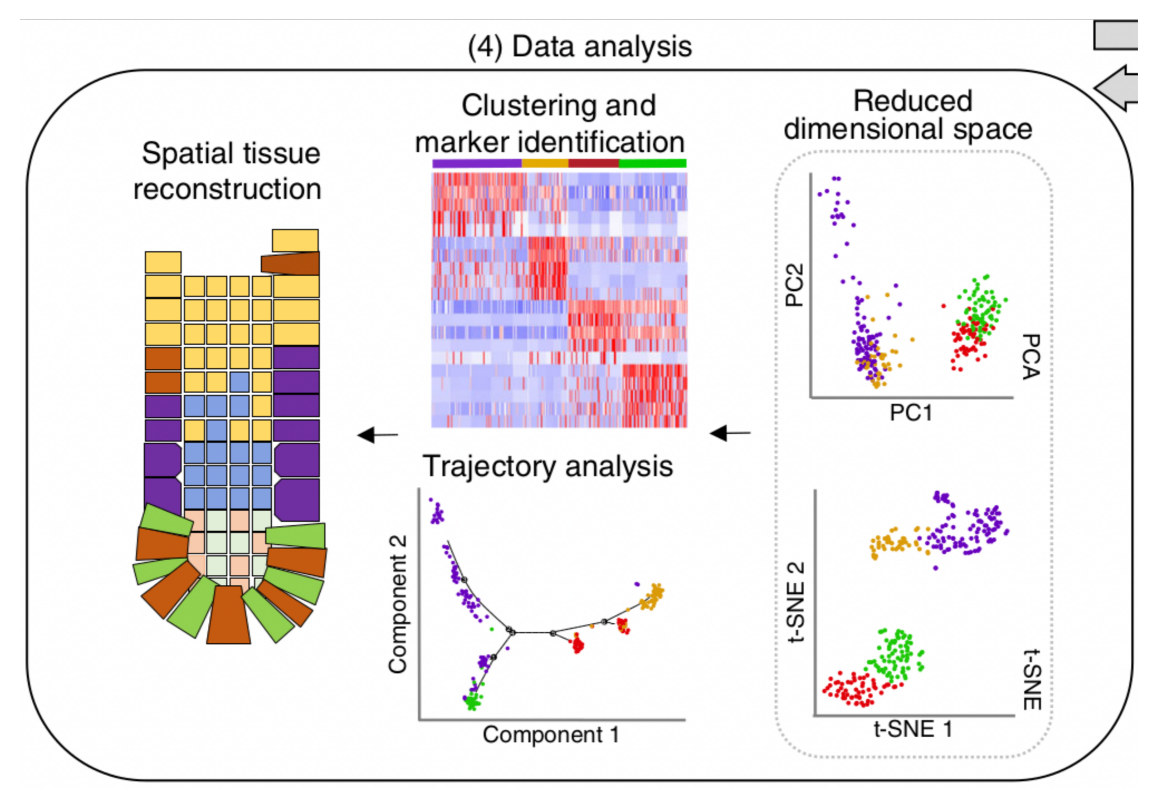
# Coverage decisions should be made based on the purpose/questions of the study.

### Single-cell RNA-Seq Data Analysis



Taken from Lafzi et al. (2018)

### Single-cell RNA-Seq Data Analysis



Taken from Lafzi et al. (2018)

### Single-cell RNA-seq Analysis Steps

Many, many tools available: http://www.scRNA-tools.org

- QC Assessment (FastQC)
- Alignment to reference (STAR within Cell Ranger)
- Quantification (within Cell Ranger, UMI-tools)

### Single-cell RNA-seq Analysis Steps

- Imputation (OPTIONAL) (SAVER)
- Normalization (scran or sctransform)
- **Dimensionality reduction** (tSNE, PCA, etc)
- Clustering (hierarchical, k-means, Seurat, etc)
- **Differential expression analysis** (deseq2, edgeR, limma, MAST)

# Cell Ranger

- Cell ranger is a set of analysis pipelines that process Chromium (10x) single-cell RNA-Seq data.
- 1. Assess quality
- 2. Aligns reads (using star)
- 3. UMI, cell barcode error correction and demultiplexing
- **'4.** Generates a gene expression matrix after 1 and 2.

5. Will also do further downstream analysis (normalization, clustering, DE analysis).

6. Analyses provided in a nice interactive report.

# Cell Ranger Web QC Page



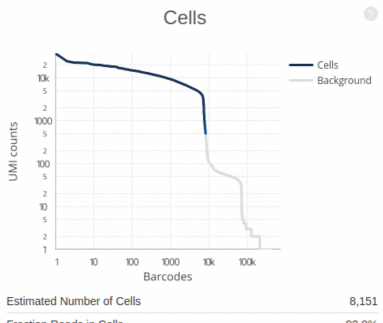
#### Cell Ranger · crtest ·

SUMMARY ANALYSIS

Estimated Number of Cells 8,151 Mean Reads per Cell Median Genes per Cell 28,670 2,405

Sequencing	
Number of Reads	233,692,380
Valid Barcodes	97.3%
Sequencing Saturation	58.4%
Q30 Bases in Barcode	95.4%
Q30 Bases in RNA Read	79.7%
Q30 Bases in UMI	95.9%

85.3%
77.4%
3.3%
15.2%
58.9%
56.5%
1.5%

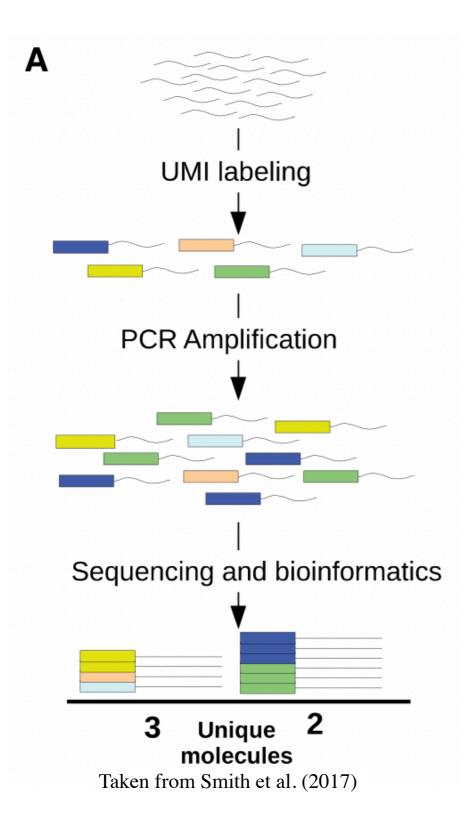


Estimated Number of Cells	8,151
Fraction Reads in Cells	92.9%
Mean Reads per Cell	28,670
Median Genes per Cell	2,405
Total Genes Detected	23,394
Median UMI Counts per Cell	5,484

#### Sample

Name	crtest	
Description		
Transcriptome	mm10	
Chemistry	Single Cell 3' v2	
Cell Ranger Version	3.0.0	

### Single-cell RNA-Seq Data Analysis UMI demultiplexing and error correction

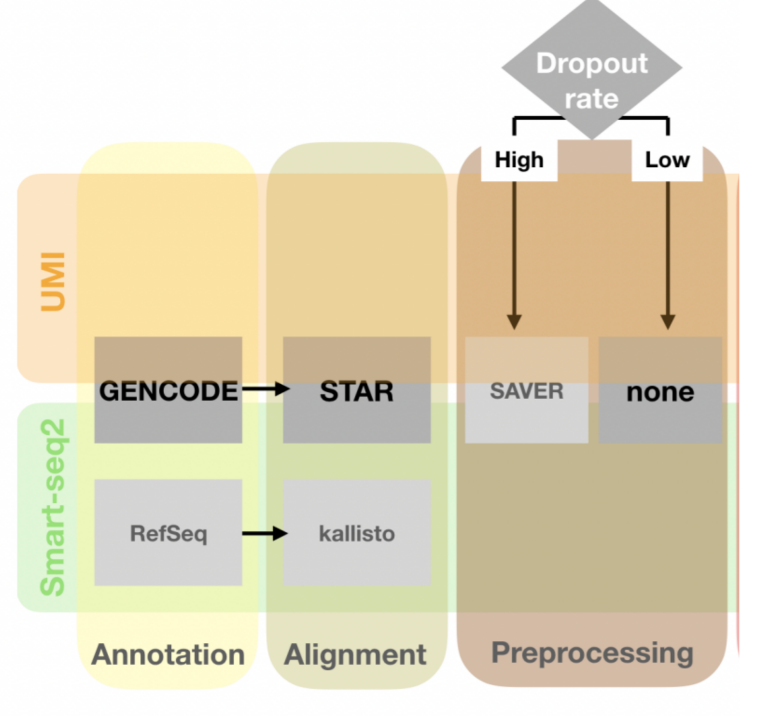


- UMIs can have sequencing errors.
- Some sequences tend to have more than others.
- UMI error correction/filtering
  - No homopolymers
  - No N's
  - No bases with quality lower than 10
  - If a UMI is 1 base pair substitution away from a higher-count UMI, it's corrected to the higher count UMI if they share a cell barcode.

### Imputation

- Method to deal with dropouts (genes with zero counts) by borrowing information from other cells.
- For a dropout gene X in cell Y,
  - Impute expression based on expression of gene X in other similar cells.

### Single-cell RNA-seq Analyses Benchmarked



- STAR mapper works well for UMI based/ chromium (10x) scRNA-Seq data
- Imputation only if dropout rate is high

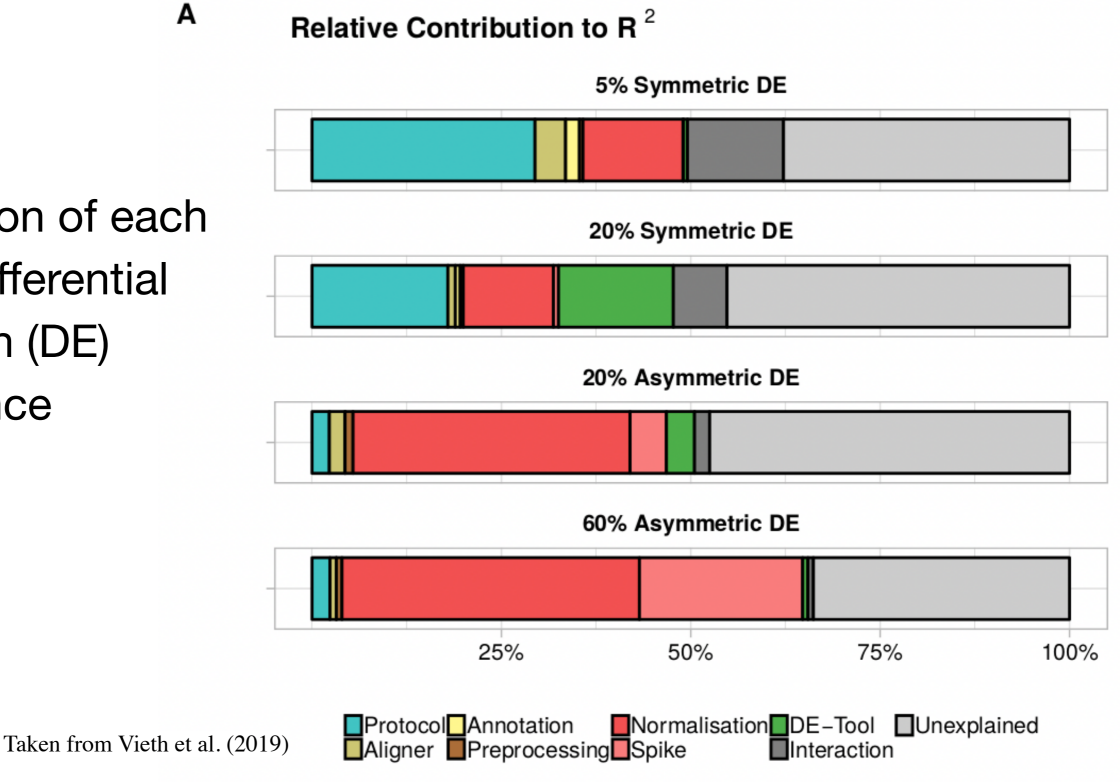
Taken from Vieth et al. (2019).

### Normalization is important!

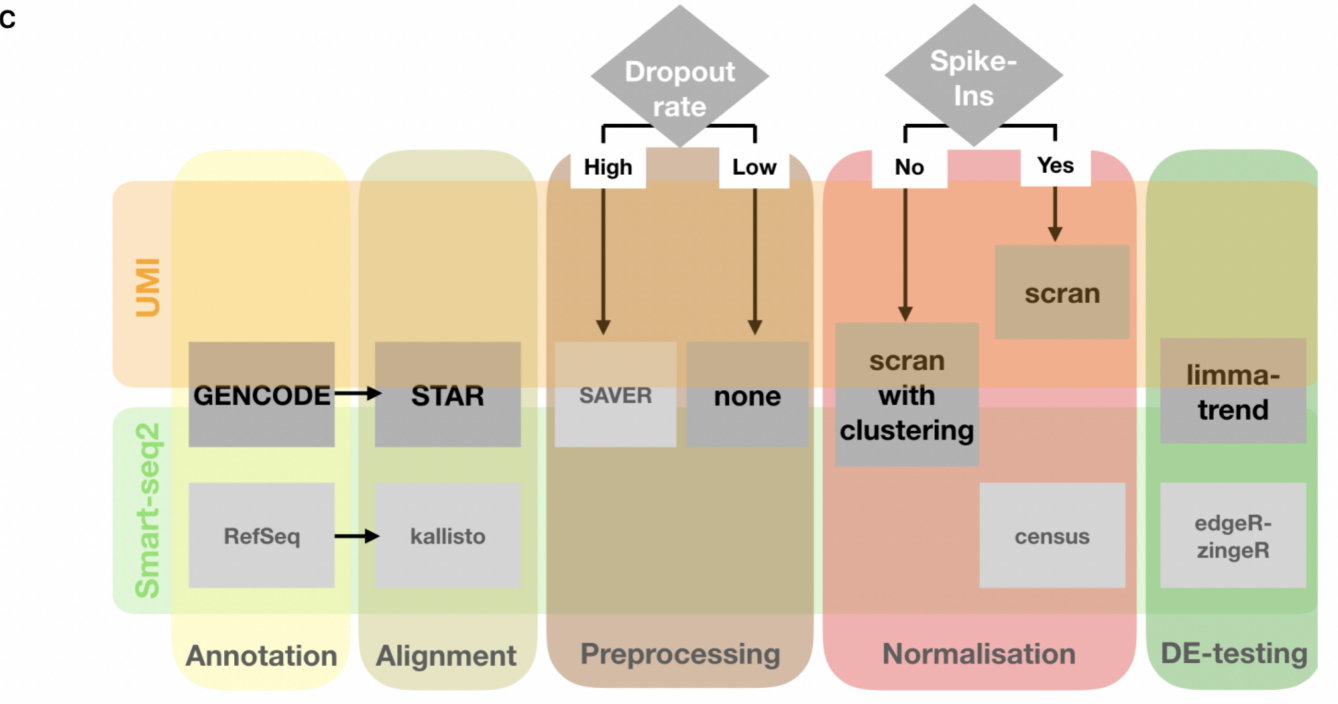
- Remove Technical variations without removing biological variation
  - dropout events, amplification bias, sequencing depth
  - batch effects
- Why is it different from normalization of bulk RNA-Seq?
  - "One main assumption in traditional DE-analysis is that differences in expression are symmetric. This implies that either a small fraction of genes is DE while the expression of the majority of genes remains constant or similar numbers of genes are up-and down-regulated so that the mean total mRNA content does not differ between groups. This assumption is no longer true when diverse cell types are considered." - Taken from Vieth et al. (2019).

### Normalization is important!

Contribution of each step on differential expression (DE) performance



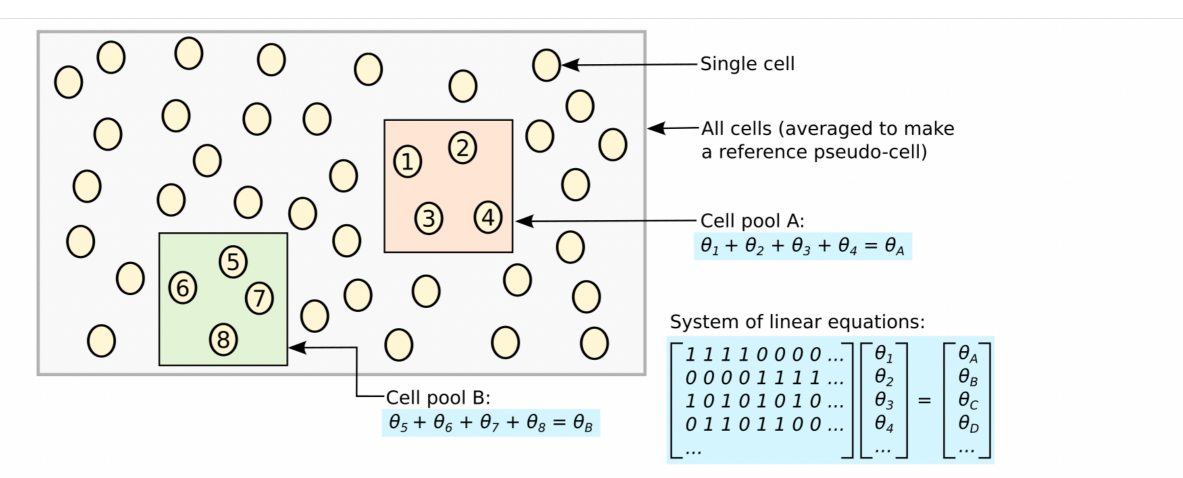
### Single-cell RNA-seq Analyses Benchmarked



Taken from Vieth et al. (2019).

### Normalization with SCRAN

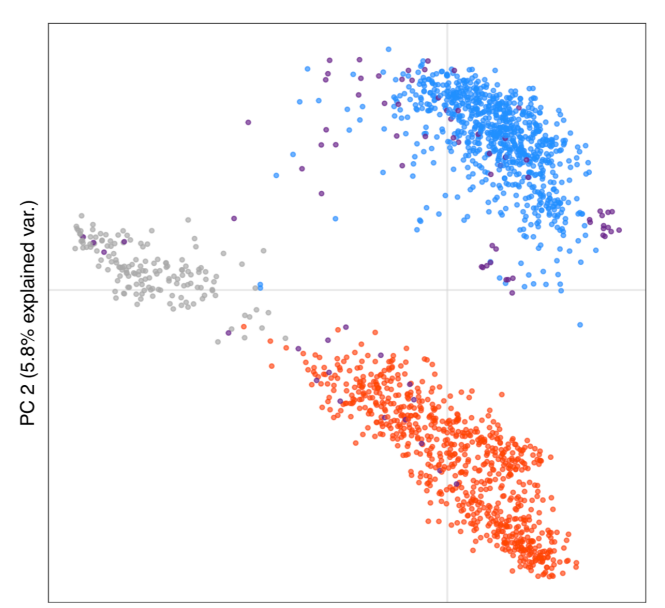
- Cluster cells into cell pools by similarity first.
- Perform normalization within each cluster/cell pool.



**Fig. 3** Schematic of the deconvolution method. All cells in the data set are averaged to make a reference pseudo-cell. Expression values for cells in pool A are summed together and normalized against the reference to yield a pool-based size factor  $\theta_A$ . This is equal to the sum of the cell-based factors  $\theta_j$  for cells j = 1-4 and can be used to formulate a linear equation. (For simplicity, the  $t_j$  term is assumed to be unity here.) Repeating this for multiple pools (e.g., pool B) leads to the construction of a linear system that can be solved to estimate  $\theta_j$  for each cell j

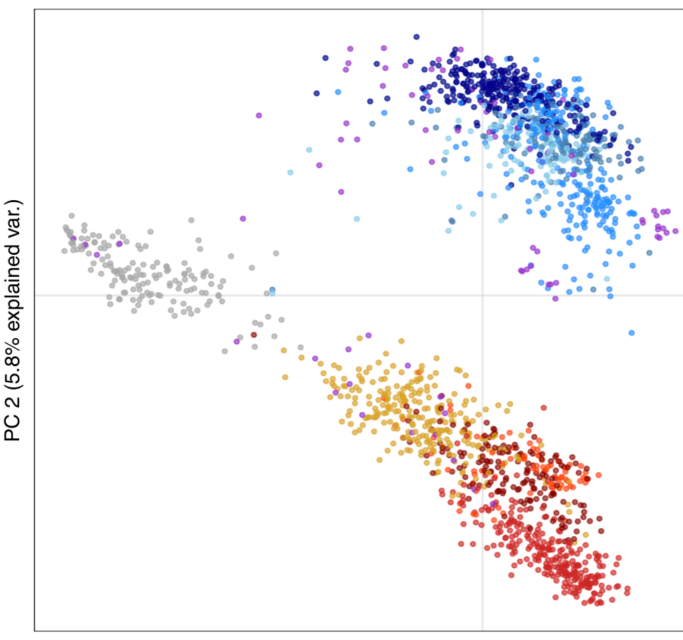
## **Dimensionality Reduction**

- Why?
  - Reduce the number of dimensions in a high dimensional data for visualization.
  - To prepare the dataset for subsequent clustering.



PC1 (12.9% explained var.)

### Dimensionality Reduction PCA

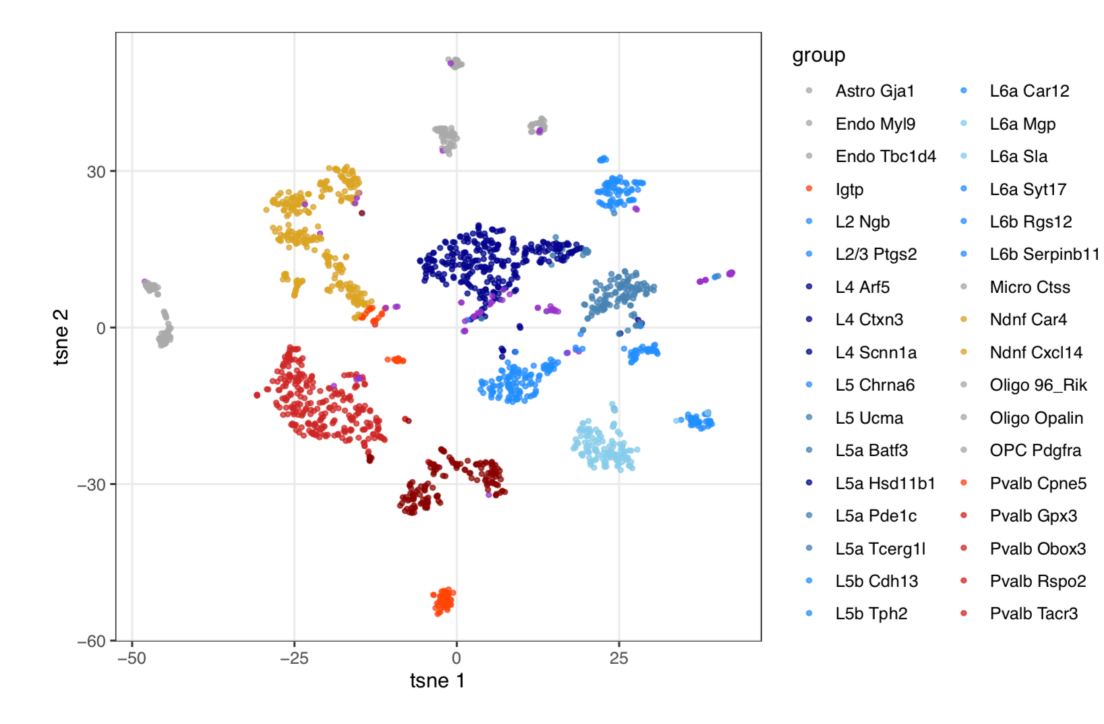


<ul> <li>As</li> </ul>	tro Gja1 🔹 🔹	L6a Car12	٠	Pvalb Tpbg
<ul> <li>En</li> </ul>	do Myl9 🛛 🔍	L6a Mgp	٠	Pvalb Wt1
• En	do Tbc1d4 🛛 🔍	L6a Sla	•	Smad3
• Igt	р •	L6a Syt17	٠	Sncg
• L2	Ngb •	L6b Rgs12	٠	Sst Cbln4
• L2	/3 Ptgs2	L6b Serpinb11	٠	Sst Cdk6
• L4	Arf5 •	Micro Ctss	•	Sst Chodl
• L4	Ctxn3 •	Ndnf Car4	٠	Sst Myh8
• L4	Scnn1a 🛛 🔸	Ndnf Cxcl14	٠	Sst Tacstd2
• L5	Chrna6 •	Oligo 96_Rik	٠	Sst Th
• L5	Ucma •	Oligo Opalin	٠	Unclassified
• L5	a Batf3	OPC Pdgfra	٠	Vip Chat
• L5	a Hsd11b1 🛛 🔸	Pvalb Cpne5	•	Vip Gpc3
• L5	a Pde1c 🔹 🔹	Pvalb Gpx3	•	Vip Mybpc1
• L5	a Tcerg1l 🛛 🔹	Pvalb Obox3	•	Vip Parm1
• L5	b Cdh13	Pvalb Rspo2	•	Vip Sncg
• L5	b Tph2 🛛 🔸	Pvalb Tacr3		

PC1 (12.9% explained var.)

Image generated by Dennis Wylie

### **Dimensionality Reduction** tSNE (t-Distributed Stochastic Neighbor Embedding )



Pvalb Tpbg

Pvalb Wt1

Sst Cbln4

Sst Cdk6

Sst Chodl

Sst Myh8

Sst Th

Sst Tacstd2

Unclassified

Vip Chat

Vip Gpc3

Vip Mybpc1

Vip Parm1

Vip Sncg

Smad3

Sncg

•

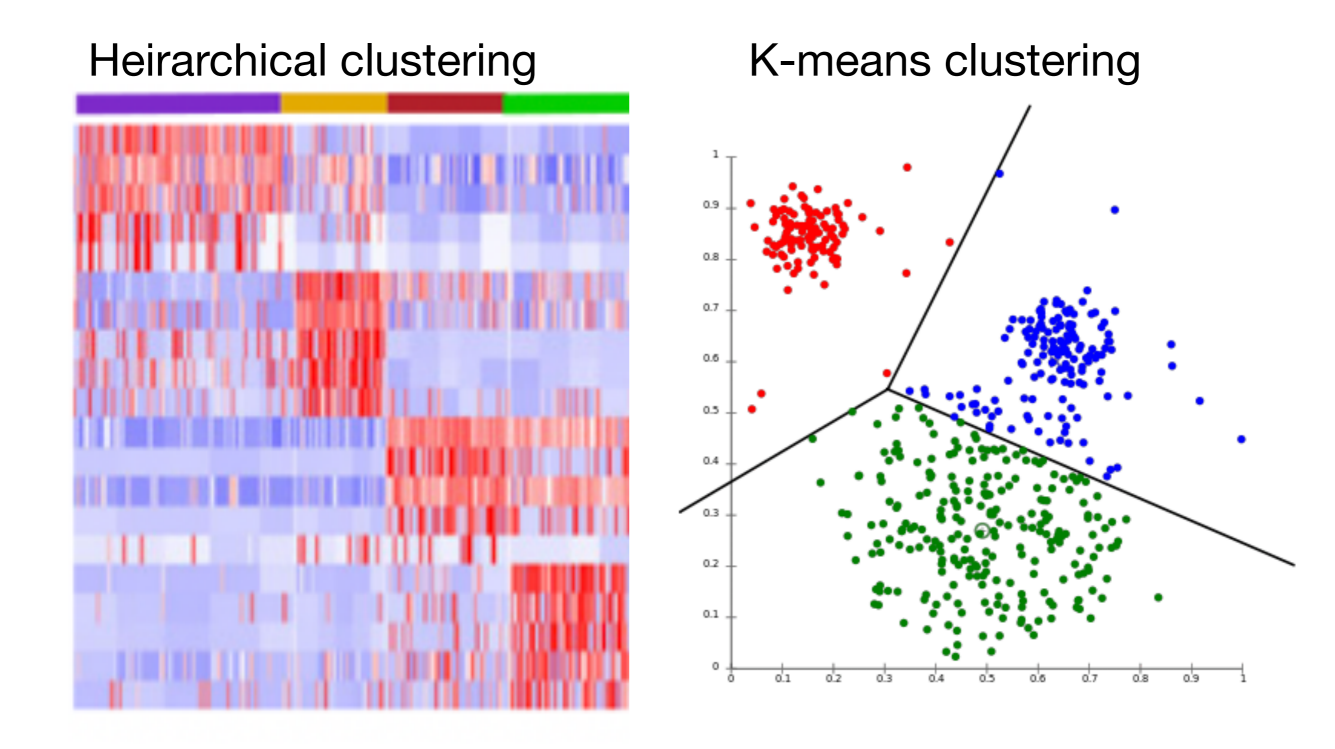
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# Clustering

### **Clustering to identify cell subpopulations**



### **Differential Expression Analysis**

#### Important to distinguish:

- 1. DE between predefined cell populations across different samples.
- 2. DE between clustering-defined subpopulations in 1 sample
  - applying standard statistical tests to clusters learned from same data set will result in very biased p values!
  - Use fold changes(effect sizes) to identify driver genes for each cluster compared to every other cluster.

### **Differential Expression Analysis**

- Bulk RNA-Seq DE analyses methods:
  - DESeq2
  - edgeR
- Specialized scRNA-Seq DE analysis methods
  - Single Cell Differential Expression (SCDE)
  - Model-based Analysis of Single-cell Transcriptomics (MAST)
- Soneson & Robinson (2018) evaluated 36 DE approaches:
  - "bulk RNA-seq analysis methods do not generally perform worse than those developed specifically for scRNA-seq."

# Conclusions

- Lots of tools available for scRNA-Seq. Tools are actively being developed, updated and benchmarked.
- **Mapping** using bulk RNA-Seq methods is just fine.
- When quantifying genes, UMIs should be taken into account and error corrected.
- Normalization is one of the most important steps in scRNA-seq data analysis and needs to be treated differently from bulk RNA-Seq datasets. Scran normalization works very well.
- **Clustering** using clustering methods to identify group for doing DE analysis.
- **DE analysis** using standard bulk RNA-Seq methods works fine.