

# Differential Expression Analysis for Single-cell RNA-seq

The Data Lab

# Why differential expression (DE) analysis with single-cell?


Differential expression analysis can be used to identify differentially expressed genes in a specific cell type or cell subpopulation across samples (e.g., comparison of T-cells across treatment groups).

# The Do's and Don'ts of Differential Expression

## Do:

- Compare expression of genes in specific cell types across sample groups (e.g., between two treatment types)
- Identify differentially expressed genes in a subpopulation of cells across sample groups

## Don't:

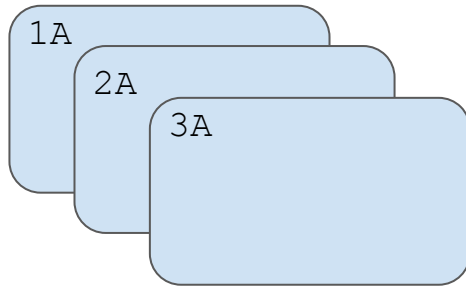
- Try to perform differential expression on a subpopulation/cell type that is not found among all samples of interest
  - Perform differential expression without replicates!
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# Differential expression analysis starts with good experimental design

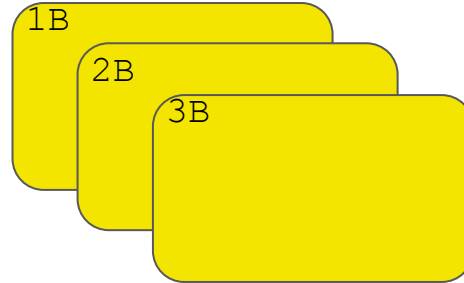
- What biological question are you trying to answer?
- The same rules of bulk RNA-seq DE analysis apply – you need biological replicates!
- Cells are NOT replicates
  - Cells from one sample are more similar to each other than cells from different samples
  - The goal is to highlight variation between samples not between cells in a sample
- Differential expression is not the same as identifying marker genes for clusters or cell types for a single sample

# Differential expression analysis starts with good experimental design

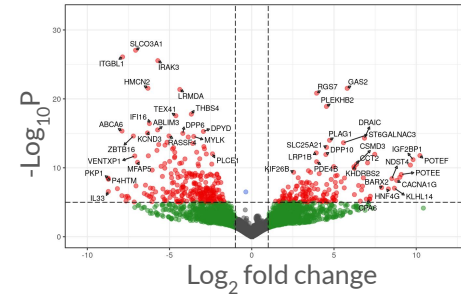
Single-cell counts matrices  
≥3 samples per group



vs.



Use gene  
expression counts  
to identify DE  
genes



# How do we perform DE analysis on single-cell data?

There are plenty of tools that exist for DE analysis in bulk RNA-seq (DESeq2, EdgeR, limma, etc.), so why can't we just use the same tools?

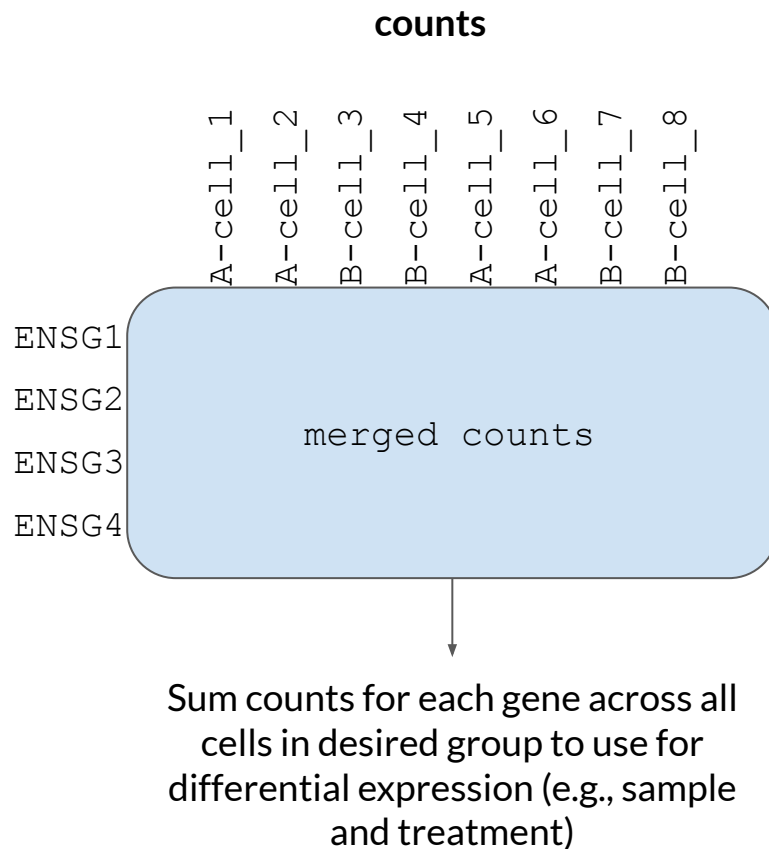
- High drop-out rate in single-cell, low gene expression counts
- Cells are treated independently, masking variation across the sample population
- Correlation of gene expression within cells from the same sample is unaccounted for

Potential solutions:

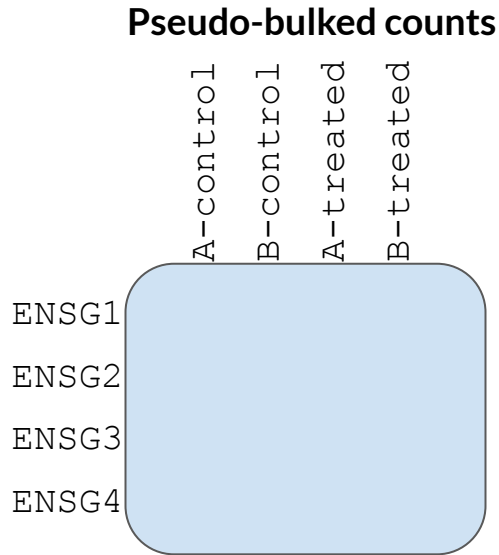
- **Calculate pseudo-bulk counts prior to using DE methods developed for bulk RNA-seq**
- Fitting a mixed-effects model to consider both drop-out and correlation between cells from the same sample (computationally intensive)
- Test for differences between distribution of a gene across a group of cells rather than between mean gene expression values

# Creating a pseudo-bulk SCE object

|          | sample | colData   |
|----------|--------|-----------|
|          |        | treatment |
| A-cell_1 | A      | Control   |
| A-cell_2 | A      | Control   |
| B-cell_3 | B      | Control   |
| B-cell_4 | B      | Control   |
| A-cell_5 | A      | Treated   |
| A-cell_6 | A      | Treated   |
| B-cell_7 | B      | Treated   |
| B-cell_8 | B      | Treated   |




# Creating a pseudo-bulk SCE object



- The resulting object will have one column for each group of cells and one row for each gene
- Similar to bulk RNA-seq we now have a sample x gene counts matrix rather than cell x gene matrix



# Why do we pseudo-bulk?

1. Produces larger and less sparse counts so we can use standard methods for normalization and DE
  2. Collapses gene expression counts by samples so samples rather than cells represent replicates
  3. Masks variance within a sample to emphasize variance across samples
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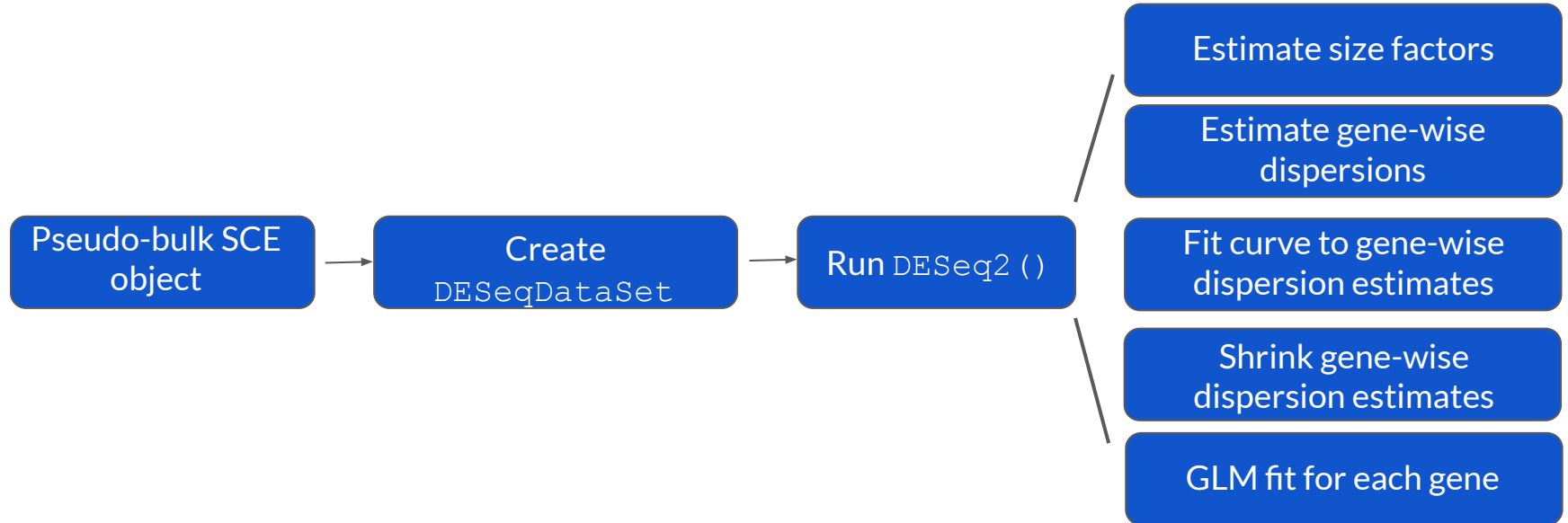
# Using DESeq2 for differential expression analysis

**Input:** Pseudo-bulk counts matrix based on the DE experiment design

- Create the pseudo-bulked counts matrix using the raw, uncorrected, unnormalized counts
- Looking at a specific cell type between treatment groups? Group by sample, cell type, and treatment

**Output:** Table of genes with associated log<sub>2</sub>-fold change, p-value, and adjusted p-value

# Using DESeq2 for differential expression analysis



# Some caveats of single-cell differential expression:

- Differential expression comes with some light circularity
  - When we pick cell types or groups of cells to perform DE, we typically have picked those groups based on expression of a subset of genes
  - This may mean we miss differences between samples, especially if those differences are large enough to change a cell label
- Use raw counts, not corrected gene expression data!
  - Correction/integration will transform the data so that between sample variation is not preserved, sometimes resulting in negative gene expression values
  - DESeq2 has been optimized for count data such that normalization and correction will affect the distribution in ways that may not be compatible with the model