Differential Expression Analysis for Single-cell RNA-seq

The Data Lab

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

- Bulk tissue collection is a mixture of cell types, so gene expression differences between samples may be dependent on changes in cell type composition
- Additionally, changes in gene expression may not be due to the cell type we are interested in
- With single-cell, we can narrow in on a specific population of interest and identify differentially expressed genes in that specific population across a set of samples
- Focusing on a single population of cells minimizes the effect changes in cell type composition has on differentially expressed genes

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:

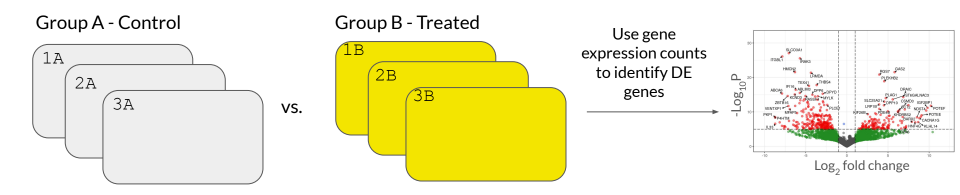
- Try to perform differential expression on a subpopulation/cell type that is not found among all samples of interest
- Perform differential expression without replicates! You should have at least 3 samples for each group being compared.

Differential expression analysis starts with good experimental design

- 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 to 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
 - Marker genes Identifying genes that are specific markers of a single group of cells within a single sample (e.g. CD4 is expressed in T-cells)
 - Differential expression Identifying genes that are differentially expressed in a group of cells between two conditions (e.g. genes expressed in stimulated T-cells vs. unstimulated T-cells)

The goal of differential expression is to identify genes that are differentially expressed between two groups

Single-cell counts matrices
≥3 samples per group



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

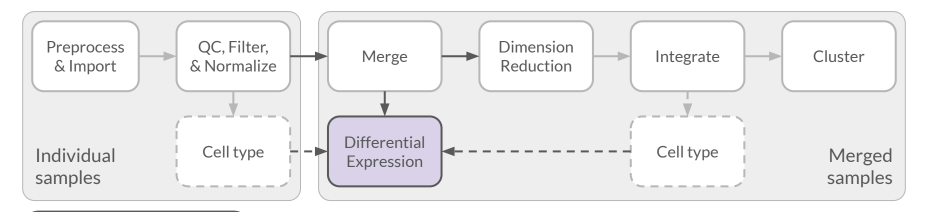
There are plenty of tools that exist for DE analysis in bulk RNA-seq (DESeq2, EdgeR, limma, etc.), so why can't we just use the same tools?

- 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

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

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



Differential Expression Pseudo-bulk counts aggregateAcrossCells() Calculate differential expression statistics DESeq() Visualize differentially expressed genes EnhancedVolcano() plotExpression()

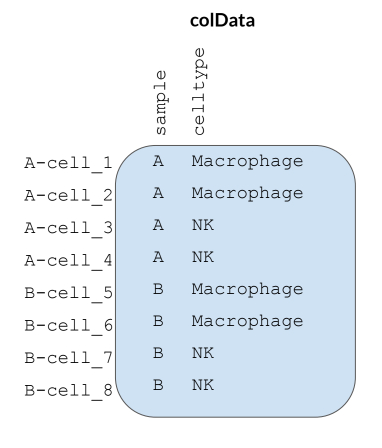
Pseudo-bulk

- Collapse gene expression counts by sample and cell type rather than individual cells
- Treats each sample rather than each cell as a replicate
- Differential expression statistics
 - Set up differential expression between two groups of samples
 - Use tools used for bulk RNA-seq analysis, like DESeq2, to identify differentially expressed genes between groups

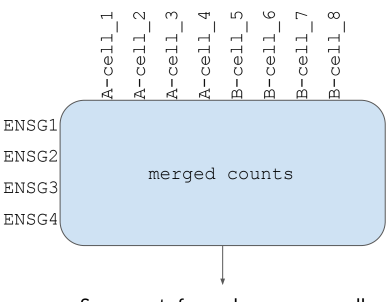
Visualize

- Look at all differentially expressed genes using a volcano plot
- Plot expression of individual genes across samples and subpopulations to validate results

Creating a pseudo-bulk SCE object



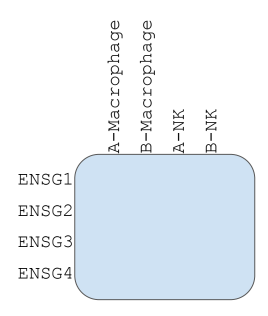
counts



Sum counts for each gene across all cells in desired group to use for differential expression (e.g., sample and treatment)

Creating a pseudo-bulk SCE object

Pseudo-bulked counts



- 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 by gene counts matrix rather than cell by 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

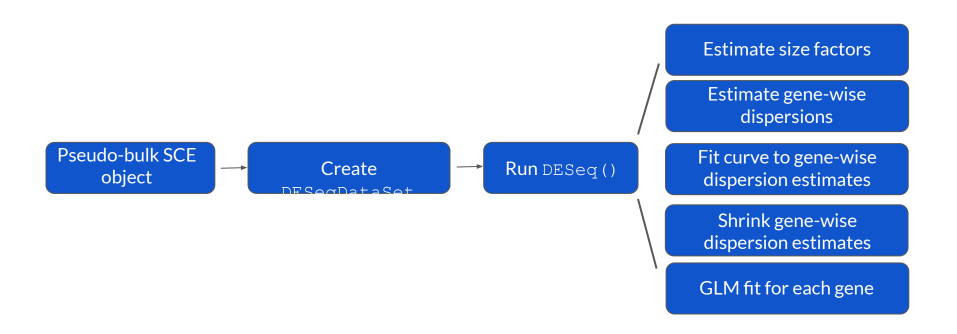
Using DESeq2 for differential expression analysis

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 log2-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