### **Advanced scRNA-seq Cheatsheet**

The tables below consist of valuable functions or commands that will help you through this module.

Each table represents a different library/tool and its corresponding commands.

#### You may also be interested in the following additional cheatsheets:

- Download the PDF for the Introduction to R and Tidyverse cheatsheet
- Download the PDF for the <u>Introduction to Single-Cell RNA sequencing cheatsheet</u>
- Download the PDF for the SCE and Seurat object comparison cheatsheet

Please note that these tables are not intended to tell you all the information you need to know about each command.

The hyperlinks found in each piece of code will take you to the documentation for further information on the usage of each command. Please be aware that the documentation will generally provide information about the given function's most current version (or a recent version, depending on how often the documentation site is updated). This will usually (but not always!) match what you have installed on your machine. If you have a different version of R or other R packages, the documentation may differ from what you have installed.

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#### scater

Read the <u>scater</u> <u>package documentation</u>, and a <u>vignette on its usage</u>.

| Library/Package | Piece of Code    | What it's called              | What it does  |
|-----------------|------------------|-------------------------------|---|
| scater          | plotReducedDim() | Plot<br>reduced<br>dimensions | Plot a given reduced dimension slot from a SingleCellExperiment object by its name                          |
| scater          | plotUMAP()       | Plot UMAP                     | Plot the "UMAP"-named reduced dimension slot from a SingleCellExperiment object                             |
| scater          | plotExpression() | Plot<br>expression            | Plot expression values for all cells in a SingleCellExperiment object, using the logcounts assay by default |

## miQC

Read the <u>miQC</u> <u>package documentation</u>, and a <u>vignette on its usage</u>.

| Library/Package | Piece of Code              | What<br>it's<br>called | What it does  |
|-----------------|----------------------------|------------------------|---|
| miQC            | <pre>mixtureModel()</pre>  | Mixture<br>model       | Fit a miQC mixture model to a SingleCellExperiment object for use in filtering  |
| miQC            | filterCells()              | Filter<br>cells        | Filter cells from a  SingleCellExperiment object based on a miQC model, returning a filtered SingleCellExperiment object                        |
| miQC            | plotMetrics()              | Plot<br>metrics        | Plot percent of mitochondrial reads against the number of unique genes found for each cell  |
| miQC            | <pre>plotModel()</pre>     | Plot<br>model          | <pre>miQC::plotMetics() with the miQC fitted model overlaid</pre>   |
| miQC            | <pre>plotFiltering()</pre> | Plot<br>filtering      | Plot percent of mitochondrial reads against the number of unique genes found, coloring points based on whether they will be filtered out or not |

### batchelor and harmony

Read the <u>batchelor</u> <u>package documentation</u>, and a <u>vignette on its usage</u>.

Read the <u>harmony</u> <u>package documentation</u>, and a <u>vignette on its usage</u>.

| Library/Package | Piece of Code              | What it's called                         | What it does  |
|-----------------|----------------------------|--|---|
| batchelor       | MultiBatchPCA()            | Multi-batch<br>PCA                       | Perform PCA across<br>multiple gene expression<br>matrices, weighted by batch<br>size   |
| batchelor       | <pre>fastMNN()</pre>       | Fast mutual nearest neighbors correction | Perform integration on an SCE object with mutual nearest neighbors using the fastmnn algorithm, returning an SCE object with batch-corrected principal components |
| harmony         | <pre>HarmonyMatrix()</pre> | Perform harmony integration on a matrix  | Perform integration with harmony on either a matrix of principle components or gene expression, returning a matrix of batch-corrected principal components        |

### SingleR

Read the <u>SingleR</u> <u>package documentation</u>, and an <u>e-book on its usage</u>.

| Library/Package | Piece of Code             | What<br>it's<br>called               | What it does   |
|-----------------|---------------------------|--------------------------------------|--|
| SingleR         | <pre>trainSingleR()</pre> | Train the<br>SingleR<br>classifier   | Build a SingleR classifier model object from an annotated reference dataset  |
| SingleR         | classifySingleR()         | Classify<br>cells<br>with<br>SingleR | Use a SingleR model object to assign cell types to the cells in an SCE object  |
| SingleR         | SingleR()                 | Annotate<br>scRNA-<br>seq data       | Combines  trainSingleR() and classifySingleR() to assign cell types to an SCE object from an annotated reference dataset |

### pheatmap and EnhancedVolcano

Read the <a href="pheatmap">pheatmap</a> <a href="package documentation">package documentation</a>.

Read the **EnhancedVolcano** package documentation, and vignette on its usage.

| Library/Package | Piece of Code         | What it's called    | What it does  |
|-----------------|-----------------------|---------------------|---|
| pheatmap        | <pre>pheatmap()</pre> | Pretty<br>heatmap   | Plot a (pretty!) clustered heatmap  |
| EnhancedVolcano | EnhancedVolcano()     | Enhanced<br>volcano | Plot a volcano plot to visualize differential expression analysis results |

## DESeq2 and pseudo-bulking functions

Read the DESeq2 package documentation, and a vignette on its usage.

| Library/Package | Piece of Code                    | What it's called                               | What it does   |
|-----------------|----------------------------------|--|--|
| scuttle         | aggregateAcrossCells()           | Aggregate<br>data across<br>groups of<br>cells | Sum counts for each combination of features across groups of cells, commonly used to <i>pseudo-bulk</i> SCE counts |
| DESeq2          | DESeqDataSet()                   | DESeq<br>Dataset                               | Establish a DESeq object from a pseudo-bulked SingleCellExperiment object or a bulk SummarizedExperiment object    |
| DESeq2          | <pre>estimateSizeFactors()</pre> | Estimate size factors                          | Estimate size factors which are used to normalize counts for differential expression analysis                      |
| DESeq2          | rlog()                           | Apply a regularized log transformation         | Log2-transform counts in a  DESeq object for differential expression analysis                                      |
| DESeq2          | plotPCA()                        | Sample PCA<br>plot for<br>transformed<br>data  | Plot sample PCA from a log-<br>transformed DESeq object<br>to check for batch effects                              |
| DESeq2          | DESeq()                          | Perform differential expression analysis       | Perform differential expression: Estimate size factors, transform data, estimate dispersions, and perform testing. |
| DESeq2          | plotDispEsts()                   | Plot<br>dispersion<br>estimates                | Plot dispersion estimates from a fitted DESeq object to evaluate model fit   |
|                 |                                  |  |  |

| DESeq2 | results()      | Extract results from a DESeq analysis | Extract results from a fitted  DESeq object into a data frame                    |
|--------|----------------|---------------------------------------|--|
| DESeq2 | resultsNames() | Extract results names                 | Return coefficient names from a fitted DESeq object                              |
| DESeq2 | lfcShrink()    | Shrink log2<br>fold changes           | Add shrunken log2-fold changes to a results table produced by  DESeq2::results() |

### tidyverse functions

#### purrr functions

Read the <u>purr package documentation</u> and a <u>vignette on its usage</u>, and download the <u>purr package cheatsheet</u>.

| Library/Package | Piece of<br>Code  | What<br>it's<br>called | What it does  |
|-----------------|-------------------|------------------------|---|
| purrr           | <u>map()</u>      | map                    | Apply a function across each element of list; return a list                     |
| purrr           | <pre>imap()</pre> | imap                   | Apply a function across each element of list and its index/names; return a list |
| purrr           | <u>map2()</u>     | map2                   | Apply a function across each element of two lists at a time; return a list      |
| purrr           | reduce()          | Reduce                 | Reduce a list to a single value by applying a given function                    |

Note that purrr::map() functions can take advantage of R's new (as of version 4.1.0) anonymous function syntax:

```
# One-line syntax:
\(x) # function code goes here #

# Multi-line syntax:
\(x) {
    # function code goes  #
    # inside the curly braces #
}

# Example: Use an anonymous function with `purrr::map()`
# to get the colData's rownames for each SCE in `list_of_sce_objects`
purrr::map(
    list_of_sce_objects,
    \(x) rownames(colData(x))
)
```

#### ggplot2 functions

Read the <u>ggplot2</u> <u>package documentation</u> and an <u>overall reference for <u>ggplot2</u> <u>functions</u>, and download the <u>ggplot2</u> <u>package cheatsheet</u>.</u>

| Library/Package | Piece of Code                  | What<br>it's<br>called         | What it does   |
|-----------------|--------------------------------|--------------------------------|--|
| ggplot2         | geom_bar()                     | Barplot                        | Creates a barplot of counts for a given categorical variable when added as a layer to a ggplot() object  |
| ggplot2         | <pre>scale_fill_brewer()</pre> | Add<br>brewer<br>fill scale    | Apply a Brewer "fill" color palette to a categorical variable in a ggplot() object                       |
| ggplot2         | guides()                       | Guides                         | Function to customize legend ("guide") appearance  |
| ggplot2         | <pre>facet_grid()</pre>        | Facet<br>grid                  | Plot individual panels using specified variables to subset the data across rows and/or columns of a grid |
| ggplot2         | <u>vars(_)</u>                 | Vars                           | Helper function to specify variables to facet_grid() or facet_wrap()                                     |
| ggplot2         | theme_bw()                     | Black<br>and<br>white<br>theme | Display ggplot with gridlines but a white background   |
| ggplot2         | theme()                        | Theme                          | Customize elements of a ggplot plot theme  |
| ggplot2         | <pre>element_text()</pre>      | Element<br>text                | Customize textual elements of a ggplot theme   |

#### dplyr, tidyr, stringr, and tibble functions

Read the full documentation and download cheatsheets (where available) for these tidyverse packages at the following links:

- <u>dplyr</u> <u>documentation</u> and <u>dplyr</u> <u>cheatsheet</u>
- <u>tidyr</u> <u>documentation</u> and <u>tidyr</u> <u>cheatsheet</u>
- <u>stringr</u> <u>documentation</u> and <u>stringr</u> <u>cheatsheet</u>
- <u>tibble</u> <u>documentation</u>

| Library/Package | Piece of Code             | What<br>it's<br>called | What it does   |
|-----------------|---------------------------|------------------------|--|
| dplyr           | pull()                    | Pull                   | Extract a single column from a data frame into a stand-alone vector  |
| dplyr           | count()                   | Count                  | Count the number of observations in each group of a data frame   |
| dplyr           | <pre>left_join()</pre>    | Left join              | Joins two data frames together, retaining only rows present in the first ("left") argument to the function |
| dplyr           | <u>relocate()</u>         | Relocate               | Change column order in a data frame by relocating one or more columns                                      |
| dplyr           | <pre>case_when(_)</pre>   | Case<br>when           | Return a value based on a set of TRUE / FALSE comparisons; a vectorized if-else                            |
| tidyr           | <pre>pivot_longer()</pre> | Pivot<br>longer        | Convert a "wide" format data frame to a "long" format data frame   |
| tibble          | as_tibble()               | As tibble              | Convert an object to a tibble  |
| stringr         | str_detect()              | String<br>detect       | Returns TRUE / FALSE if a string contains a given substring  |
| stringr         | str_starts()              | String<br>starts       | Returns TRUE / FALSE if a string starts with a given substring   |

# Pathway analysis

msigdbr

Read the <u>msigdbr</u> <u>package documentation</u> and <u>its vignette</u>.

| Library/Package | Piece of Code     | What it's called                 | What it does   |
|-----------------|-------------------|----------------------------------|--|
| msigdbr         | msigdbr_species() | List msigdbr - supported species | Lists the species  msigdbr supports                            |
| msigdbr         | msigdbr()         | Retrieve<br>gene set             | Retrieves gene sets and member genes in long data frame format |

#### clusterProfiler and enrichplot

Read the <u>clusterProfiler</u> <u>package documentation (PDF)</u>.

| Library/Package | Piece of Code | What it's called                    | What it does   |
|-----------------|---------------|-------------------------------------|--|
| clusterProfiler | GSEA()        | Gene Set Enrichment Analysis (GSEA) | Performs a universal gene set<br>enrichment analysis on given<br>preranked (sorted) named<br>vector of statistics, where the<br>names in the vector are gene<br>identifiers of gene sets |
| enrichplot      | gseaplot()    | GSEA plot                           | Produces a plot displaying the distribution of gene set and enrichment score   |

### AUCell and GSEABase

Read the <u>AUCell</u> <u>package documentation</u> and <u>its vignette</u>.

| Library/Package | Piece of Code                  | What it's called       | What it does   |
|-----------------|--------------------------------|------------------------|--|
| GSEABase        | <pre>GeneSet()</pre>           | Gene set               | Constructs a gene set as a GeneSet object for use with AUCell  |
| GSEABase        | <pre>GeneSetCollection()</pre> | Gene set collection    | Constructs a collection of gene sets as a  GeneSetCollection object for use with  AUCell                                   |
| AUCell          | <u>AUCell_buildRankings()</u>  | Build cell<br>rankings | Builds a ranking of genes<br>for each cell that is used<br>to calculate the recovery<br>curve                              |
| AUCell          | AUCell_calcAUC()               | Calculate<br>AUC       | Calculates the area under the recovery curve (AUC) for each gene set in each cell  |
| AUCell          | AUCell_exploreThresholds()     | Explore<br>thresholds  | Calculates thresholds in AUC values that can be used to assign cells; optionally makes assignments and produces histograms |

### bluster

Read the <u>bluster</u> <u>package documentation</u> and vignettes on its usage:

- Flexible clustering for Bioconductor
- Assorted clustering diagnostics

| Library/Package | Piece of Code                   | What it's called                                   | What it does   |
|-----------------|---------------------------------|--|--|
| bluster         | clusterRows()                   | Cluster rows<br>of a matrix                        | Perform clustering using a variety of algorithms on a matrix-like object               |
| bluster         | <pre>approxSilhouette()</pre>   | Approximate silhouette width                       | Calculate an approximate silhouette width for each cell given a set of clusters        |
| bluster         | <pre>neighborPurity()</pre>     | Compute<br>neighborhood<br>purity                  | Calculate neighborhood purity for each cell given a set of clusters                    |
| bluster         | <pre>bootstrapStability()</pre> | Assess<br>cluster<br>stability by<br>bootstrapping | Generate cluster bootstrap replicates to estimate cluster robustness to sampling noise |