# 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.

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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#### Base R

Read the Base R documentation.

Library/Package	Piece of Code	What it's called	What it does
Base R	rowSums()	Row sums	Calculates sums for each row
Base R	colSums()	Column sums	Calculates sums for each column
Base R	t()	Transpose	Returns the transpose of a matrix or data frame
Base R	prcomp()	Principal Components Analysis	Executes a principal components analysis on specified matrix or data frame
Base R	<pre>&lt;- function(x) { <code> }</code></pre>	Function	Creates a function that would take the defined parameters as input and execute the commands within the curly braces

# Salmon and alevinQC

Read the command-line tool Salmon documentation.

Read the R package alevinQC documentation.

Software/package	Piece of Code	What it's called	What it does
Salmon	salmon alevin	Salmon Alevin	Runs the Alevin quantification from the command line
alevinQC	alevinQCReport()	Alevin QC Report	Produces a QC (quality check) report from the salmon alevin output

### SingleCellExperiment , txmimeta , and DropletUtils

Read the SingleCellExperiment package documentation (and e-book), and a vignette on its usage.

Note that some of the SingleCellExperiment functions link to documentation from other packages like SummarizedExperiment or ExperimentSubset.

In fact, SingleCellExperiment objects are based around existing Bioconductor functions in those packages, so the function usage is equivalent!

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

Read the DropletUtils package documentation.

Library/Package	Piece of Code	What it's called	What it does
SingleCellExperiment	SingleCellExperiment()	Single Cell Experiment	Creates a SingleCellExperiment object
SingleCellExperiment	colData()	Column Data	Extracts and stores cell-level metadata that describes features of the SingleCellExperiment object
SingleCellExperiment	rowData()	Row Data	Extracts and stores gene-level metadata that describes features of the SingleCellExperiment object
SingleCellExperiment	assay()	Assay	Extracts and stores a given assay from a SingleCellExperiment object
SingleCellExperiment	assayNames()	Assay names	Returns a vector of the names of all assays in a SingleCellExperiment object
SingleCellExperiment	logcounts()	Log counts	Extracts and stores log-transformed single-cell experiment count data as an assay of the SingleCellExperiment object
SingleCellExperiment	counts()	Counts	Extracts and stores raw single-cell experiment count data as an assay of the SingleCellExperiment object
SingleCellExperiment	reducedDim()	Reduced dim	Extracts or stores a given reduced dimension from a SingleCellExperiment object
SingleCellExperiment	reducedDimNames()	Reduced dim names	Returns a vector of the names of all reduced dimensions in a SingleCellExperiment object
S4Vectors	DataFrame()	Data frame	Not to be confused with data.frame() from Base R. This is a slightly different data frame-like object needed for storing information in  SingleCellExperiment object's colData slot.

Library/Package	Piece of Code	What it's called	What it does
tximeta	tximeta()	Transcript Quantification Import with Automatic Metadata	Load a directory of results produced by Salmon/or alevin output, including the associated metadata
DropletUtils	read10xCounts()	Read 10x counts	Load data from a 10x Genomics experiment into R
DropletUtils	emptyDrops()	Empty drops	Use the overall gene expression patterns in the sample to identify empty droplets
DropletUtils	emptyDropsCellRanger()	Empty drops Cell Ranger	Use an approach analogous to Cell Ranger's algorithm to identify empty droplets

### scran and scater

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

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

Library/Package	Piece of Code	What it's called	What it does
scran	quickCluster()	Quick Clustering	Groups similar cells into clusters which are stored in the SingleCellExperiment object and are used for the calculation of size factors by scran::computeSumFactors
scran	<pre>computeSumFactors()</pre>	Compute Sum Factors	Returns a numeric vector of computed sum factors for each cell cluster stored in the SingleCellExperiment object. The cluster-based size factors are deconvolved into cell-based size factors that are stored in the SingleCellExperiment object and used by the scran::normalize function for the normalization of each cell's gene expression profile
scran	getTopHVGs()	Get top highly variable genes	Identify variable genes in a SingleCellExperiment object, based on variance
scran	modelGeneVar()	model per gene variance	Model the per gene variance of a SingleCellExperiment object
scran	findMarkers()	Find marker genes	Find candidate marker genes for clusters of cells
scater	logNormCounts()	Normalize log counts	Returns the SingleCellExperiment object with normalized expression values for each cell, using the size factors stored in the object
scater	addPerCellQC()	Add per cell quality control	For a SingleCellExperiment object, calculate and add quality control per cell and store in colData
scater	addPerFeatureQC()	Add per feature quality control	For a SingleCellExperiment object, calculate and add quality control per feature (genes usually) and store in rowData
scater	calculatePCA()	Calculate PCA	Calculates principal components analysis on a SingleCellExperiment object, returning a PCA matrix
scater	runPCA()	Run PCA	Calculates principal components analysis on a SingleCellExperiment object, returning an SCE object with a PCA reduced dimension

Library/Package	Piece of Code	What it's called	What it does
scater	calculateUMAP()	Calculate UMAP	Calculates uniform manifold approximate projection on a SingleCellExperiment object, returning a UMAP matrix
scater	runUMAP()	Run UMAP	Calculates uniform manifold approximate projection on a SingleCellExperiment object, returning an SCE object with a UMAP reduced dimension
scater	calculateTSNE()	Calculate t- SNE	Calculates t-stochastic neighbor embedding on a SingleCellExperiment object, returning an SCE object with a TSNE reduced dimension
scater	runTSNE()	Calculate UMAP	Calculates t-stochastic neighbor embedding on a SingleCellExperiment object, returning a t-SNE matrix
scater	<pre>plotReducedDim()</pre>	Plot reduced dimensions	Plot a given reduced dimension slot from a SingleCellExperiment object by its name
scater	plotPCA()	Plot PCA	Plot the "PCA"-named reduced dimension slot from a SingleCellExperiment object
scater	plotUMAP()	Plot UMAP	Plot the "UMAP"-named reduced dimension slot from a SingleCellExperiment object

### purrr, stringr, and tibble

Read the purrr package documentation.

Read the stringr package documentation.

Read the tibble package documentation.

Library/Package	Piece of Code	What it's called	What it does
purrr	map()	map	Apply a function across each element of list; return a list
purrr	map_df()	map df	Apply a function across each element of list; return a data frame
purrr	imap()	imap	Apply a function across each element of list and its index/names
stringr	str_remove()	String remove	Remove matched string patterns
tibble	as_tibble()	As tibble	Coerce data.frame or matrix to a tibble

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))
}
```

# SingleR

Read the SingleR package documentation, and an e-book on its usage.

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