Bulk RNA-seq Cheatsheet

The tables below consist of valuable codes 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.

Library/Pacl	kage Piece of Code	What it's called	What it does
Base R	list.files()	List files	Produces a character vector of files or directories in the specified directory
Base R	names()	Names	Gets or sets the names of an object
Base R	colnames()	Column names	Gets or sets the column names of a matrix or data frame
Base R	all.equal()	All equal	Checks if two R objects are nearly equal
Base R	attr()	$egin{array}{c} { m Object} \ { m Attributes} \end{array}$	Gets or sets the attributes of an object
Base R	rowSums()	Row Sums	Returns the sum of the rows in a numeric matrix-like object (i.e., a matrix, data.frame, etc.)
Base R	relevel()	Relevel	Reorders the levels of a factor as specified
Base R	summary()	Object Summary	Returns a result summary of an object
Base R	as.data.frame()	Data Frame	Checks if an object is a data.frame, and transforms the object into one, if possible

		What it's	
Library/Package Piece of Code		called	What it does
DESeq2	rlog()	Regularized	Returns log2 transformed count
		Log	data
		Transformation	
DESeq2	vst()	Variance	Estimates the dispersion trend of
		Stabilizing	count data
		Transformation	
DESeq2	<pre>DESeqDataSetFromTximport()</pre>	DESeqDataSet	Creates a DESeqDataSet object
		constructor	
DESeq2	DESeq()	Differential	Estimates size factors, estimates
		Expression	dispersion, and performs negative
		Analysis	binomial fitting and Wald statistics
		Based on the	as steps in the default differential
		Negative	expression analysis
		Binomial	
		Distribution	
DESeq2	plotPCA()	PCA plot	Produces a principal component analysis plot for transformed data, while checking for batch effects

		What it's	
Library/Package Piece of Code		called	What it does
DESeq2	counts()	Counts	Saves the count data as a matrix
DESeq2	results()	Results	Returns the results table from a DESeq2 analysis
DESeq2	assay()	Assay	Returns count values in a matrix
		7771	
Library/Package Piece of Code		What it's called	What it does
fastp	fastp	FASTQ preprocessor	Preprocesses FASTQ files through adapter trimming, quality filtering, length filtering, and a number of additional option
FastQC	fastqc	FASTQC (Quality Control)	Performs quality control checks on raw sequence data and outputs a QC(quality control) report
		What it's	
Library/Packs	age Piece of Code	called	What it does
ggplot2	ggsave()	GG Save	Saves the last plot in working directory
ggplot2	<pre>last_plot()</pre>	Last plot	Returns the last plot produced
ggplot2	<pre>geom_point()</pre>	Geom point	Creates a scatterplot (when added to the ggplot() function)
ggplot2	<pre>xlab(); ylab()</pre>	X Axis Labels; Y Axis Labels	Modifies the labels on the x axis and on the y axis, respectively
ggplot2	<pre>coord_fixed()</pre>	Cartesian Coordinates with Fixed Aspect Ratio	Coerces the coordinates on the plot to represent a fixed specified ratio
Library/Package Piece of Code		What it's called	What it does
stringr	word()	Word	Extracts words from a character
readr tximport	<pre>write_rds() tximport()</pre>	Write RDS Tximport	vector Writes data to a .rds output file Imports transcript-level estimates and summarizes abundances, counts, and transcript lengths to the gene-level by default
dplyr	pull()	Pull	Extracts a variable as a vector
pheatmap	<pre>pheatmap()</pre>	Pheatmap	Plots clustered heatmaps

Tool	Piece of Code	What it's called	What it does
Salmon	salmon index	Salmon index	Builds a transcriptome index which is required for Salmon quantification (from the command line)
Salmon	salmon quant	Salmon quantification	Runs Salmon's quantification of transcript expression (from the command line)

Useful command line commands:

```
DODLO MacDock-Pro:~ ccdl$ pwd
/Users/ccdl Directory we start in
Applications Desktop Documents Downloads Library
                                                                    Movies
                                                                                  Music
CCDLs-MacBook-Pro:Desktop ccd1$ 1s
2019-chicago-training
                                                Screen Shot 2019-07-18 at 1.33.02 PM.png
                                               Screen Shot 2019-07-18 at 1.34.01 PM.png
Forms
Repos
                                               Screen Shot 2019-07ext9 at 9.28.52 PM.png
Screen Shot 2019-07-16 at 2.14.11 PM.png
                                               Screen Shot 2019-07-23 at 2.36.20 PM.png
CCDLs-MacBook-Pro:Desktop ccd1$

    Our new working directory

Hello world!
CCDLs-MBP: Desktop ccdl$ echo Hello bash!  echo command prints the string that follows
Hello bash!
[CCDLs-MBP:Desktop ccd1$ mkdir internet_file | Internet_file | mkdir command creates a directory in the current directory
CCDLs-MBP:Desktop ccdl$ cd internet_file/
[CCDLs-MBP:internet_file ccdl$ pwd

    wget downloads a file from the internet

                                         and saves it in the current directory
/Users/ccdl/Desktop/internet_file
CCDLs-MBP:internet_file ccdl$ wget -0 ccdl_webpage https://www.ccdatalab.org/
--2019-07-24 08:47:27-- https://www.ccdatalab.org/
Resolving www.ccdatalab.org (www.ccdatalab.org)... 198.185.159.144, 198.49.23.144, 198.49.23.145, ...
Connecting to www.ccdatalab.org (www.ccdatalab.org)|198.185.159.144|:443... connected.
[HTTP request sent, awaiting response... 200 OK
Length: 88867 (87K) [text/html]
Saving to: 'ccdl_webpage'
ccdl_webpage
                  2019-07-24 08:47:27 (901 KB/s) - 'ccdl_webpage' saved [88867/88867]
CCDLs-MBP:internet_file ccdl$ ls
                              ---- the command is shows us the file we downloaded via URL in the directory we created
ccdl_webpage
```