# **Machine Learning 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.

#### **AnnotationDbi**

Read the AnnotationDbi package vignette here.

Library/Package	Piece of Code	What it's called	What it does
AnnotationDbi	keytypes()	Keytypes	Returns a character vector of column names/keytypes (eg., type of gene identifiers) available in an AnnotationDbi package.
AnnotationDbi	mapIDs()	Mapped IDs	Extracts the mapped ids for a set of keys (e.g., gene identifiers) of a specific keytype

# Base R

Read the Base R package documentation <u>here</u>.

Library/Package	Piece of Code	What it's called	What it does
Base R	round()	Round	Rounds the values in the object provided in the first argument to the number of decimal places specified in the second argument
Base R	<u>identical()</u>	Identical	Checks if two objects are exactly equal
Base R	prcomp()	Principal Components Analysis	Executes a principal components analysis on specified matrix or data frame
Base R	rowSums()	Row Sums	Returns the sums of the rows in a numeric array, matrix, or data.frame
Base R	rowMeans()	Row Means	Returns the means of the rows in a numeric array, matrix, or data.frame
Base R	<pre>quantile()</pre>	Sample Quantiles	Returns the sample quantiles for a given numeric vector of data and numeric vector of probabilities
Base R	cor()	Correlation	Computes correlation between columns using a specified correlation method, and returns a correlation matrix
Base R	as.dist()	Distance matrix compution	Returns a special object of class dist, a distance matrix used by the hclust() function
Base R	hclust()	Hierarchical Clustering	Performs hierarchical clustering analysis on a set of dissimilarities and methods
Base R	table()	Create Table	Creates a contingency table of counts for each combination of factor levels
Base R	<u>duplicated()</u>	Duplicated	Returns a logical vector, where TRUE represents elements of the object that are duplicates
Base R	any()	Any	Checks to see if at least one of the elements are TRUE when given a logical vector
Base R	cbind()	Cbind	Combines vectors, matrices, or data.frames by columns
Base R	<pre>pairwise.wilcox.test()</pre>	Pairwise Wilcoxon Rank Sum Tests	Calculates the pairwise comparisons between group levels

### **PLIER**

Read the PLIER package documentation <u>here</u>.

A PLIER package vignette can be found <a href="here">here</a> and can also serve as documentation for the commands in the table below.

Library/Package	Piece of Code	What it's called	What it does	
PLIER	combinePaths()	Combine Pathways	Combines the pathway data obtained from PLIER and returns the result as a matrix	
PLIER	commonRows()	Common Rows	Determines the rows (genes) that are common to the specified data matrices and returns them as a character vector	
PLIER	rowNorm()	Row Normalize	Normalizes each row (gene) by z-scoring the expression values	
PLIER	num.pc()	Number of Principal Components	Returns the number of significant principal components	
PLIER	PLIER()	Main PLIER Function	Main function of the Pathway-Level Information ExtractoR.	
PLIER	plotU()	Plot U Matrix	Plots the U matrix obtained from the PLIER function results, allowing insight into the pathways or cell types captured by the latent variables	

#### ComplexHeatmap

Read the ComplexHeatmap package documentation <u>here</u>.

Library/Package	Piece of Code	What it's called	What it does	
ComplexHeatmap	Heatmap()	Complex Heatmap	Constructs a heatmap whose graphics and features can be defined	
ComplexHeatmap	<pre>HeatmapAnnotation()</pre>	Heatmap Annotation Constructor	Creates an annotation object to be used in conjunction with a Heatmap	

## ggplot2

Read the ggplot2 package documentation here.

A vignette on the usage of the ggplot2 package can be found here.

Library/Package	Piece of Code	What it's called	What it does
ggplot2	<pre>geom_jitter()</pre>	Jittered Points	Adds a small amount of random variation at each point's location on a plot
ggplot2	labs()	Labels Sets the axis, legend, and plot labels if specified	
ggplot2	theme()	Theme	Sets the specified non-data elements of a plot (i.e. plot title, legend spacing, text size, etc.)

# tidyr

Read the tidyr package documentation here.

A vignette on the usage of the tidyr package can be found here. I Library/Package | Piece of Code | What it's called | What it does | I-----

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# data.table , purrr , dplyr , matrixStats , umap , ConsensusClusterPlus , plotly , tidyr , ggsignif

Documentation for each of these packages can be accessed by clicking the package name in the table below.

Library/Package	Piece of Code	What it's called	What it does
data.table	<pre>fread()</pre>	F read	Reads in data faster than base R
purrr	discard()	Discard	Discards the given elements
dplyr	pull()	Pull	Pulls a single variable out of a given table of data
matrixStats	rowSds()	Row Standard Deviations	Returns the standard deviation estimates for each row in a matrix
<u>matrixStats</u>	rowVars()	Row Variances	Returns the variance estimates for each row in a matrix
<u>umap</u>	umap(_)	Uniform Manifold Approximation and Projection (UMAP)	Computes a manifold approximation and projection on a given matrix or data.frame
ConsensusClusterPlus	ConsensusClusterPlus()	Consensus Clustering	Finds the consensus across multiple runs of the clustering algorithm
plotly	plot_ly()	Plotly Visualization	Initiates a plotly visualization with given R objects
ggsignif	<pre>geom_signif()</pre>	Create Significance Layer	Adds significance information to the plot. It can be used to run statistical tests and display the significance information from those tests. We use it differently, in a way that gives us more control, in the notebook.