

Package ‘PathoStat’

September 18, 2025

Type Package

Title PathoStat Statistical Microbiome Analysis Package

Version 1.34.0

Date 2020-03-27

Author Solaiappan Manimaran <manimaran_1975@hotmail.com>, Matthew Bendall <bendall@gwmail.gwu.edu>, Sandro Valenzuela Diaz <sandrolvalenzuelad@gmail.com>, Eduardo Castro <castronallar@gmail.com>, Tyler Faits <tfaits@gmail.com>, Yue Zhao <jasonzhao0307@gmail.com>, Anthony Nicholas Federico <anfed@bu.edu>, W. Evan Johnson <wej@bu.edu>

Maintainer Solaiappan Manimaran <manimaran_1975@hotmail.com>, Yue Zhao <jasonzhao0307@gmail.com>

Description The purpose of this package is to perform Statistical Microbiome Analysis on metagenomics results from sequencing data samples. In particular, it supports analyses on the PathoScope generated report files. PathoStat provides various functionalities including Relative Abundance charts, Diversity estimates and plots, tests of Differential Abundance, Time Series visualization, and Core OTU analysis.

URL <https://github.com/mani2012/PathoStat>

BugReports <https://github.com/mani2012/PathoStat/issues>

License GPL (>= 2)

Depends R (>= 3.5)

Imports limma, corpcor, matrixStats, reshape2, scales, ggplot2, rentrez, DT, tidyr, plyr, dplyr, phyloseq, shiny, stats, methods, XML, graphics, utils, BiocStyle, edgeR, DESeq2, ComplexHeatmap, plotly, webshot, vegan, shinyjs, glmnet, gmodels, ROCR, RColorBrewer, knitr, devtools, ape

Collate 'pathoStat.R' 'utils.R' 'taxonomy.R' 'biomarker.R' 'allClasses.R' 'visualization.R' 'differentialAnalysis.R'

biocViews Microbiome, Metagenomics, GraphAndNetwork, Microarray,
PatternLogic, PrincipalComponent, Sequencing, Software,
Visualization, RNASeq, ImmunoOncology

RoxygenNote 6.1.1

Encoding UTF-8

Suggests rmarkdown, testthat

VignetteBuilder knitr

git_url <https://git.bioconductor.org/packages/PathoStat>

git_branch RELEASE_3_21

git_last_commit 1a2cb3a

git_last_commit_date 2025-04-15

Repository Bioconductor 3.21

Date/Publication 2025-09-17

Contents

Bootstrap_LOOCV_LR_AUC	3
Chisq_Test_Pam	4
findRAfromCount	4
findTaxonMat	5
findTaxonomy	6
findTaxonomy300	6
Fisher_Test_Pam	7
formatTaxTable	8
getShinyInput	8
getShinyInputCombat	9
getShinyInputOrig	9
getSignatureFromMultipleGlmnet	10
GET_PAM	10
grepTid	11
loadPathoscopeReports	12
loadPstat	13
log2CPM	13
LOOAUC_simple_multiple_noplot_one_df	14
LOOAUC_simple_multiple_one_df	14
PathoStat-class	15
percent	16
phyloseq_to_edgeR	16
plotPCAPlotly	17
plotPCoAPlotly	18
pstat_data	19
readPathoscopeData	19
runPathoStat	20
savePstat	21

Bootstrap_LOOCV_LR_AUC

3

setShinyInput

21

setShinyInputCombat

22

setShinyInputOrig

22

summarizeTable

23

TranslateIdToTaxLevel

23

Wilcox_Test_df

24

Index

25

Bootstrap_LOOCV_LR_AUC

Do bootstrap and LOOCV

Description

Do bootstrap and LOOCV

Usage

Bootstrap_LOOCV_LR_AUC(df, targetVec, nboot = 50)

Arguments

- df

Row is sample, column is feature. Required
- targetVec

y vector. Required
- nboot

number of BOOTSTRAP

Value

bootstrap loocv result dataframe

Examples

```
data('iris')
Bootstrap_LOOCV_LR_AUC(iris[,1:4],
c(rep(1,100), rep(0,50)), nboot = 3)
```

Chisq_Test_Pam	<i>Given PAM and disease/control annotation, do Chi-square test for each row of PAM</i>
----------------	---

Description

Given PAM and disease/control annotation, do Chi-square test for each row of PAM

Usage

```
Chisq_Test_Pam(pam, label.vec.num, pvalue.cutoff = 0.05)
```

Arguments

pam	Input data object that contains the data to be tested. Required
label.vec.num	The target binary condition. Required
pvalue.cutoff	choose p-value cut-off

Value

df.output object

Examples

```
tmp <- matrix(rbinom(12,1,0.5), nrow = 3)
rownames(tmp) <- c("a", "b", "c")
Chisq_Test_Pam(tmp, c(1,1,0,0))
```

findRAfromCount	<i>Return the Relative Abundance (RA) data for the given count OTU table</i>
-----------------	--

Description

Return the Relative Abundance (RA) data for the given count OTU table

Usage

```
findRAfromCount(count_otu)
```

Arguments

count_otu	Count OTU table
-----------	-----------------

Value

ra_otu Relative Abundance (RA) OTU table

Examples

```
data_dir <- system.file("data", package = "PathoStat")
infileName <- "pstat_data.rda"
pstat_test <- loadPstat(data_dir, infileName)
ra_otu <- findRAfromCount(phyloseq::otu_table(pstat_test))
```

findTaxonMat	<i>Find the Taxonomy Information Matrix</i>
--------------	---

Description

Find the Taxonomy Information Matrix

Usage

```
findTaxonMat(names, taxonLevels)
```

Arguments

names	Row names of the taxonomy matrix
taxonLevels	Taxon Levels of all tids

Value

taxmat Taxonomy Information Matrix

Examples

```
example_data_dir <- system.file("example/data", package = "PathoStat")
pathoreport_file_suffix <- "-sam-report.tsv"
datlist <- readPathoscopeData(example_data_dir, pathoreport_file_suffix,
input.files.name.vec = as.character(1:6))
dat <- datlist$data
ids <- rownames(dat)
tids <- unlist(lapply(ids, FUN = grepTid))
# taxonLevels <- findTaxonomy(tids[1:5])
# taxmat <- findTaxonMat(ids[1:5], taxonLevels)
```

findTaxonomy*Find the taxonomy for unlimited tids*

Description

Find the taxonomy for unlimited tids

Usage

```
findTaxonomy(tids)
```

Arguments

tids Given taxonomy ids

Value

taxondata Data with the taxonomy information

Examples

```
example_data_dir <- system.file("example/data", package = "PathoStat")
pathoreport_file_suffix <- "-sam-report.tsv"
datlist <- readPathoscopeData(example_data_dir, pathoreport_file_suffix,
input.files.name.vec = as.character(1:6))
dat <- datlist$data
ids <- rownames(dat)
tids <- unlist(lapply(ids, FUN = grepTid))
# taxonLevels <- findTaxonomy(tids[1:5])
```

findTaxonomy300*Find the taxonomy for maximum 300 tids*

Description

Find the taxonomy for maximum 300 tids

Usage

```
findTaxonomy300(tids)
```

Arguments

tids Given taxonomy ids

Value

taxondata Data with the taxonomy information

Examples

```
example_data_dir <- system.file("example/data", package = "PathoStat")
pathoreport_file_suffix <- "-sam-report.tsv"
datlist <- readPathoscopeData(example_data_dir,
pathoreport_file_suffix, input.files.name.vec = as.character(1:6))
dat <- datlist$data
ids <- rownames(dat)
tids <- unlist(lapply(ids, FUN = grepTid))
# taxonLevels <- findTaxonomy300(tids[1:5])
```

Fisher_Test_Pam	<i>Given PAM and disease/control annotation, do Chi-square test for each row of PAM</i>
-----------------	---

Description

Given PAM and disease/control annotation, do Chi-square test for each row of PAM

Usage

```
Fisher_Test_Pam(pam, label.vec.num, pvalue.cutoff = 0.05)
```

Arguments

pam	Input data object that contains the data to be tested. Required
label.vec.num	The target binary condition. Required
pvalue.cutoff	choose p-value cut-off

Value

df.output object

Examples

```
tmp <- matrix(rbinom(12,1,0.5), nrow = 3)
rownames(tmp) <- c("a", "b", "c")
Fisher_Test_Pam(tmp, c(1,1,0,0))
```

formatTaxTable	<i>Format taxonomy table for rendering</i>
----------------	--

Description

Format taxonomy table for rendering

Usage

```
formatTaxTable(ttable)
```

Arguments

ttable	Taxonomy table
--------	----------------

Value

Formatted table suitable for rendering with. DT::renderDataTable

getShinyInput	<i>Getter function to get the shinyInput option</i>
---------------	---

Description

Getter function to get the shinyInput option

Usage

```
getShinyInput()
```

Value

shinyInput option

Examples

```
getShinyInput()
```

<code>getShinyInputCombat</code>	<i>Getter function to get the shinyInputCombat option</i>
----------------------------------	---

Description

Getter function to get the shinyInputCombat option

Usage

```
getShinyInputCombat()
```

Value

shinyInputCombat option

Examples

```
getShinyInputCombat()
```

<code>getShinyInputOrig</code>	<i>Getter function to get the shinyInputOrig option</i>
--------------------------------	---

Description

Getter function to get the shinyInputOrig option

Usage

```
getShinyInputOrig()
```

Value

shinyInputOrig option

Examples

```
getShinyInputOrig()
```

```
getSignatureFromMultipleGlmnet
```

Use Lasso to do feature selection

Description

Use Lasso to do feature selection

Usage

```
getSignatureFromMultipleGlmnet(df.input, target.vec, nfolds = 10,
  logisticRegression = TRUE, nRun = 100, alpha = 1)
```

Arguments

df.input	Row is sample, column is feature. Required
target.vec	y vector. Required
nfolds	glmnet CV nfolds
logisticRegression	doing logistic regression or linear regression.
nRun	number of glmnet runs
alpha	same as in glmnet

Value

signature

Examples

```
data('iris')
getSignatureFromMultipleGlmnet(iris[,1:4],
  c(rep(1,100), rep(0,50)), nfolds = 3, nRun = 10)
```

```
GET_PAM
```

transform cpm counts to presence-absence matrix

Description

transform cpm counts to presence-absence matrix

Usage

```
GET_PAM(df)
```

Arguments

df Input data object that contains the data to be tested. Required

Value

df.output object

Examples

```
GET_PAM(data.frame(a = c(1,3,0), b = c(0,0.1,2)))
```

grepTid	<i>Greps the tid from the given identifier string</i>
---------	---

Description

Greps the tid from the given identifier string

Usage

```
grepTid(id)
```

Arguments

id Given identifier string

Value

tid string

Examples

```
grepTid("ti|700015|org|Coriobacterium_glomerans_PW2")
```

`loadPathoscopeReports` *Loads all data from a set of PathoID reports. For each column in the PathoID report, construct a matrix where the rows are genomes and the columns are samples. Returns a list where each element is named according to the PathoID column. For example, `ret[["Final.Best.Hit.Read.Numbers"]]` on the result of this function will get you the final count matrix. Also includes elements "total_reads" and "total_genomes" from the first line of the PathoID report.*

Description

Loads all data from a set of PathoID reports. For each column in the PathoID report, construct a matrix where the rows are genomes and the columns are samples. Returns a list where each element is named according to the PathoID column. For example, `ret[["Final.Best.Hit.Read.Numbers"]]` on the result of this function will get you the final count matrix. Also includes elements "total_reads" and "total_genomes" from the first line of the PathoID report.

Usage

```
loadPathoscopeReports(reportfiles, nrows = NULL)
```

Arguments

<code>reportfiles</code>	Paths to report files
<code>nrows</code>	Option to read first N rows of PathoScope reports

Value

Returns a list where each element is named according to the PathoID column. For example, `ret[["Final.Best.Hit.Read.Numbers"]]` on the result of this function will get you the final count matrix. Also includes elements "total_reads" and "total_genomes" from the first line of the PathoID report.

Examples

```
input_dir <- system.file("example/data", package = "PathoStat")
reportfiles <- list.files(input_dir, pattern = "*-sam-report.tsv",
full.names = TRUE)
```

loadPstat	<i>Load the R data(.rda) file with pathostat object</i>
-----------	---

Description

Load the R data(.rda) file with pathostat object

Usage

```
loadPstat(indir = ".", infileName = "pstat_data.rda")
```

Arguments

indir	Input Directory of the .rda file
infileName	File name of the .rda file

Value

pstat pathostat object (NULL if it does not exist)

Examples

```
data_dir <- system.file("data", package = "PathoStat")
infileName <- "pstat_data.rda"
pstat <- loadPstat(data_dir, infileName)
```

log2CPM	<i>Compute log2(counts per mil reads) and library size for each sample</i>
---------	--

Description

Compute log2(counts per mil reads) and library size for each sample

Usage

```
log2CPM(qcounts, lib.size = NULL)
```

Arguments

qcounts	quantile normalized counts
lib.size	default is colsums(qcounts)

Value

list containing log2(quantile counts per mil reads) and library sizes

Examples

```
log2CPM(matrix(1:12, nrow = 3))
```

LOOAUC_simple_multiple_noplot_one_df
<i>LOOCV</i>

Description

LOOCV

Usage

```
LOOAUC_simple_multiple_noplot_one_df(df, targetVec)
```

Arguments

- df Row is sample, column is feature. Required
- targetVec y vector. Required

Value

mean auc

Examples

```
data('iris')
LOOAUC_simple_multiple_noplot_one_df(iris[,1:4],
c(rep(1,100), rep(0,50)))
```

LOOAUC_simple_multiple_one_df
<i>LOOCV with ROC curve</i>

Description

LOOCV with ROC curve

Usage

```
LOOAUC_simple_multiple_one_df(df, targetVec)
```

Arguments

- df Row is sample, column is feature. Required
- targetVec y vector. Required

Value

the ROC

Examples

```
data('iris')
LOOAUC_simple_multiple_one_df(iris[,1:4],
c(rep(1,100), rep(0,50)))
```

PathoStat-class

PathoStat class to store PathoStat input data including phyloseq object

Description

Contains all currently-supported BatchQC output data classes:

Details

slots:

average_count a single object of class otu_tableOrNULL

besthit_count a single object of class otu_tableOrNULL

highconf_count a single object of class otu_tableOrNULL

lowconf_count a single object of class otu_tableOrNULL

Examples

```
otumat = matrix(sample(1:100, 100, replace = TRUE), nrow = 10, ncol = 10)
rownames(otumat) <- paste0("OTU", 1:nrow(otumat))
colnames(otumat) <- paste0("Sample", 1:ncol(otumat))
taxmat = matrix(sample(letters, 70, replace = TRUE),
nrow = nrow(otumat), ncol = 7)
rownames(taxmat) <- rownames(otumat)
colnames(taxmat) <- c("Domain", "Phylum", "Class",
"Order", "Family", "Genus", "Species")
OTU = phyloseq::otu_table(otumat, taxa_are_rows = TRUE)
TAX = phyloseq::tax_table(taxmat)
physeq = phyloseq::phyloseq(OTU, TAX)
pathostat1(physeq)
```

percent	<i>Compute percentage</i>
---------	---------------------------

Description

Compute percentage

Usage

```
percent(x, digits = 2, format = "f")
```

Arguments

x	a number or a vector
digits	how many digit of percentage
format	numeric format, "f" for float

Value

the percentage

Examples

```
percent.vec <- percent(c(0.9, 0.98))
```

phyloseq_to_edgeR	<i>Convert phyloseq OTU count data into DGEList for edgeR package</i>
-------------------	---

Description

Further details.

Usage

```
phyloseq_to_edgeR(physeq, group, method = "RLE", ...)
```

Arguments

physeq	(Required).
group	(Required). A character vector or factor giving the experimental group/condition for each sample/library.
method	(Optional).
...	Additional arguments passed on to

Value

dispersion

Examples

```
data_dir_test <- system.file("data", package = "PathoStat")
pstat_test <- loadPstat(indir=data_dir_test,
  inFile="pstat_data_2_L1.rda")
phyloseq_to_edgeR(pstat_test, group="Sex")
```

plotPCAPlotly	<i>Plot PCA</i>
---------------	-----------------

Description

Plot PCA

Usage

```
plotPCAPlotly(df.input, condition.color.vec,
  condition.color.name = "condition", condition.shape.vec = NULL,
  condition.shape.name = "condition", columnTitle = "Title",
  pc.a = "PC1", pc.b = "PC2")
```

Arguments

- df.input Input data object that contains the data to be plotted. Required
- condition.color.vec color vector. Required
- condition.color.name color variable name. Required
- condition.shape.vec shape vector. Required
- condition.shape.name shape variable name. Required
- columnTitle Title to be displayed at top of heatmap.
- pc.a pc.1
- pc.b pc.2

Value

the plot

Examples

```
data('iris')
plotPCAPlotly(t(iris[,1:4]),
condition.color.vec = c(rep(1,100), rep(0,50)),
condition.shape.vec = c(rep(0,100), rep(1,50)))
```

plotPCoAPlotly

Plot PCoA

Description

Plot PCoA

Usage

```
plotPCoAPlotly(physeq.input, condition.color.vec,
condition.color.name = "condition", condition.shape.vec = NULL,
condition.shape.name = "condition", method = "bray",
columnTitle = "Title", pc.a = "Axis.1", pc.b = "Axis.2")
```

Arguments

physeq.input	Input data object that contains the data to be plotted. Required
condition.color.vec	color vector. Required
condition.color.name	color variable name. Required
condition.shape.vec	shape vector. Required
condition.shape.name	shape variable name. Required
method	which distance metric
columnTitle	Title to be displayed at top of heatmap.
pc.a	pc.1
pc.b	pc.2

Value

the plot

Examples

```
data_dir_test <- system.file("data", package = "PathoStat")
pstat_test <- loadPstat(indir=data_dir_test,
infileName="pstat_data_2_L1.rda")
plotPCoAPlotly(pstat_test, condition.color.vec = rbinom(33,1,0.5),
condition.shape.vec = rbinom(33,1,0.5))
```

pstat_data	<i>pathostat object generated from example pathoscope report files</i>
------------	--

Description

This example data consists of 33 samples from a diet study with 11 subjects taking 3 different diets in random order

Usage

```
pstat
```

Format

pathostat object extension of phyloseq-class experiment-level object:

otu_table OTU table with 41 taxa and 33 samples

sample_data Sample Data with 33 samples by 18 sample variables

tax_table Taxonomy Table with 41 taxa by 9 taxonomic ranks

sample_data Phylogenetic Tree with 41 tips and 40 internal nodes

Value

pathostat object

readPathoscopeData	<i>Reads the data from PathoScope reports and returns a list of final guess relative abundance and count data</i>
--------------------	---

Description

Reads the data from PathoScope reports and returns a list of final guess relative abundance and count data

Usage

```
readPathoscopeData(input_dir = ".",
  pathoreport_file_suffix = "-sam-report.tsv", use.input.files = FALSE,
  input.files.path.vec = NULL, input.files.name.vec = NULL)
```

Arguments

`input_dir` Directory where the tsv files from PathoScope are located
`pathoreport_file_suffix`
 PathoScope report files suffix
`use.input.files`
 whether input dir to pathoscope files or directly pathoscope files
`input.files.path.vec`
 vector of pathoscope file paths
`input.files.name.vec`
 vector of pathoscope file names

Value

List of final guess relative abundance and count data

Examples

```

example_data_dir <- system.file("example/data", package = "PathoStat")
pathoreport_file_suffix <- "-sam-report.tsv"
datlist <- readPathoscopeData(example_data_dir, pathoreport_file_suffix,
input.files.name.vec = as.character(1:6))
  
```

runPathoStat	<i>Statistical Microbiome Analysis on the pathostat input and generates a html report and produces interactive shiny app plots</i>
--------------	--

Description

Statistical Microbiome Analysis on the pathostat input and generates a html report and produces interactive shiny app plots

Usage

```

runPathoStat(pstat = NULL, report_dir = ".",
report_option_binary = "11111111", interactive = TRUE)
  
```

Arguments

`pstat` phyloseq extension pathostat object
`report_dir` Output report directory path
`report_option_binary`
 9 bits Binary String representing the plots to display and hide in the report
`interactive` when TRUE, opens the interactive shinyApp

Value

outfile The output file with all the statistical plots

Examples

```
runPathoStat(interactive = FALSE)
```

savePstat	<i>Save the pathostat object to R data(.rda) file</i>
-----------	---

Description

Save the pathostat object to R data(.rda) file

Usage

```
savePstat(pstat, outdir = ".", outfileName = "pstat_data.rda")
```

Arguments

pstat	pathostat object
outdir	Output Directory of the .rda file
outfileName	File name of the .rda file

Value

outfile .rda file

Examples

```
data_dir_test <- system.file("data", package = "PathoStat")
pstat_test <- loadPstat(indir=data_dir_test,
  infileName="pstat_data_2_L1.rda")
outfile <- savePstat(pstat_test)
```

setShinyInput	<i>Setter function to set the shinyInput option</i>
---------------	---

Description

Setter function to set the shinyInput option

Usage

```
setShinyInput(x)
```

Arguments

x	shinyInput option
---	-------------------

Value

shinyInput option

Examples

```
setShinyInput(NULL)
```

setShinyInputCombat	<i>Setter function to set the shinyInputCombat option</i>
---------------------	---

Description

Setter function to set the shinyInputCombat option

Usage

```
setShinyInputCombat(x)
```

Arguments

x	shinyInputCombat option
---	-------------------------

Value

shinyInputCombat option

Examples

```
setShinyInputCombat(NULL)
```

setShinyInputOrig	<i>Setter function to set the shinyInputOrig option</i>
-------------------	---

Description

Setter function to set the shinyInputOrig option

Usage

```
setShinyInputOrig(x)
```

Arguments

x	shinyInputOrig option
---	-----------------------

Value

shinyInputOrig option

Examples

```
setShinyInputOrig(NULL)
```

summarizeTable	<i>Summarize sample</i>
----------------	-------------------------

Description

Creates a table of summary metrics

Usage

```
summarizeTable(pstat)
```

Arguments

pstat Input pstat

Value

A data.frame object of summary metrics.

Examples

```
data_dir_test <- system.file("data", package = "PathoStat")
pstat_test <- loadPstat(indir=data_dir_test,
  infileName="pstat_data_2_L1.rda")
st.tmp <- summarizeTable(pstat_test)
```

TranslateIdToTaxLevel	<i>Find the taxonomy for the given taxon id name</i>
-----------------------	--

Description

Find the taxonomy for the given taxon id name

Usage

```
TranslateIdToTaxLevel(pstat, input.id.vec, tax.level)
```

Arguments

pstat	pathostat object
input.id.vec	names containing id
tax.level	target taxon level

Value

target taxon level names

Examples

```
data_dir_test <- system.file("data", package = "PathoStat")
pstat_test <- loadPstat(indir=data_dir_test,
  inFile="pstat_data_2_L1.rda")
names.new <- TranslateIdToTaxLevel(pstat_test,
  c("ti|862962|org|Bacteroides_fragilis_638R",
    "ti|697329|org|Ruminococcus_albus_7" ),
  "genus")
```

Wilcox_Test_df	<i>Mann-whitney test for a dataframe</i>
----------------	--

Description

Mann-whitney test for a dataframe

Usage

```
Wilcox_Test_df(df, label.vec.num, pvalue.cutoff = 0.05)
```

Arguments

df	Input data object that contains the data to be tested. Required
label.vec.num	The target binary condition. Required
pvalue.cutoff	choose p-value cut-off

Value

df.output object

Examples

```
data('iris')
Wilcox_Test_df(t(iris[,1:4]),
  c(rep(1,100), rep(0,50)))
```

Index

- * **datasets**
 - pstat_data, [19](#)
- Bootstrap_LOOCV_LR_AUC, [3](#)
- Chisq_Test_Pam, [4](#)
- findRAfromCount, [4](#)
- findTaxonMat, [5](#)
- findTaxonomy, [6](#)
- findTaxonomy300, [6](#)
- Fisher_Test_Pam, [7](#)
- formatTaxTable, [8](#)
- GET_PAM, [10](#)
- getShinyInput, [8](#)
- getShinyInputCombat, [9](#)
- getShinyInputOrig, [9](#)
- getSignatureFromMultipleGlmnet, [10](#)
- grepTid, [11](#)
- loadPathoscopeReports, [12](#)
- loadPstat, [13](#)
- log2CPM, [13](#)
- LOOAUC_simple_multiple_noplot_one_df, [14](#)
- LOOAUC_simple_multiple_one_df, [14](#)
- PathoStat-class, [15](#)
- pathostat1 (PathoStat-class), [15](#)
- percent, [16](#)
- phyloseq_to_edgeR, [16](#)
- plotPCAPlotly, [17](#)
- plotPCoAPlotly, [18](#)
- pstat (pstat_data), [19](#)
- pstat_data, [19](#)
- readPathoscopeData, [19](#)
- runPathoStat, [20](#)
- savePstat, [21](#)
- setShinyInput, [21](#)
- setShinyInputCombat, [22](#)
- setShinyInputOrig, [22](#)
- summarizeTable, [23](#)
- TranslateIdToTaxLevel, [23](#)
- Wilcox_Test_df, [24](#)