

Package ‘PathoStat’

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Type Package

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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 (>= 3.3.1)

Imports MCMCpack, limma, corpcor, rmarkdown, knitr, pander, matrixStats, reshape2, scales, ggplot2, rentrez, BatchQC, DT, gtools, tidyr, plyr, dplyr, ape, phyloseq, shiny, grDevices, stats, methods, XML, graphics, utils, alluvial, BiocStyle, edgeR, preprocessCore, DESeq2

Collate 'pathoStat.R' 'utils.R' 'taxonomy.R' 'confRegion.R' 'allClasses.R' 'coreOTUModule.R' 'normalization.R'

Suggests testthat

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

SystemRequirements pandoc (<http://pandoc.org/installing.html>) for generating reports from markdown files.

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NeedsCompilation no

R topics documented:

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coreOTU

Compute Core OTUs for the given data matrix

Description

Compute Core OTUs for the given data matrix

Usage

```
coreOTU(zcounts, otuthreshold = 0.05, prevalence = 0.4)
```

Arguments

zcounts	Standardized counts
otuthreshold	Abundance cutoff threshold for the OTU to be picked
prevalence	Prevalence of the OTU at threshold cutoff among samples

Value

list containing core OTUs

Examples

```
example_data_dir <- system.file("example/data", package = "PathoStat")
pathoreport_file_suffix <- "-sam-report.tsv"
datlist <- readPathoscopeData(example_data_dir, pathoreport_file_suffix)
countdat <- datlist$countdata
coreotus <- coreOTU(countdat)
```

 coreOTUModule

Server function for Core OTU Module

Description

This function provides the server logic for the Core OTU tab. This function is not called directly; instead, it should be invoked within the Shiny app's server function using the `callModule` function. See <http://shiny.rstudio.com/articles/modules.html> for information about this design pattern.

Usage

```
coreOTUModule(input, output, session, pstat)
```

Arguments

input	Shiny server input object created by <code>callModule</code>
output	Shiny server output object created by <code>callModule</code>
session	Session created by <code>callModule</code>
pstat	PathoStat object (third argument to <code>callModule</code>).

Details

The `callModule` function should be invoked with this function as the first argument. `callModule` is responsible for creating the namespaced `input`, `output`, and `session` arguments. The second argument to `callModule` is the ID to be used for the namespace and *must* match the `id` argument provided to `coreOTUModuleUI`. The third argument to `callModule` should be a `PathoStat` object from the app's server function, and is passed to this function as the `pstat` argument.

Value

None

See Also

[coreOTUModuleUI](#) for the UI function, [callModule](#) to see how to invoke this function, or <http://shiny.rstudio.com/articles/modules.html> for more information about Shiny modules.

Examples

```
# This function is not called directly; instead, it should be invoked within
# the app's server function using the shiny::callModule function.
## Not run:
shinyServer(function(input, output, session) {
  shinyInput <- getShinyInput()
  pstat <- shinyInput$pstat
  callModule( coreOTUModule, "coreOTUModule", pstat )
})

## End(Not run)
```

coreOTUModuleUI

UI function for Core OTU Module

Description

This function creates the UI for the Core OTU tab. The tab panel can be included within a `tabsetPanel`, thus providing a simple way to add or remove this module from the Shiny app. The first argument, `id`, is the ID to be used for the namespace *and* must match the `id` argument provided to [coreOTUModule](#).

Usage

```
coreOTUModuleUI(id, label = "Core OTUs")
```

Arguments

<code>id</code>	Namespace for module
<code>label</code>	Tab label

Value

A `tabPanel` that can be included within a `tabsetPanel`.

See Also

[coreOTUModule](#) for the server function, [tabPanel](#) for the UI component returned by this function, or <http://shiny.rstudio.com/articles/modules.html> for more information about Shiny modules.

Examples

```
shiny::mainPanel(  
  shiny::tabsetPanel(  
    coreOTUModuleUI("coreOTUModule")  
  )  
)
```

coreOTUNormalize	<i>Compute Empirical Bayes OTU Normalized data</i>
------------------	--

Description

Compute Empirical Bayes OTU Normalized data

Usage

```
coreOTUNormalize(zcounts, wt = 0.25, otuthreshold = 0.05,  
  prevalence = 0.4)
```

Arguments

zcounts	counts data to be normalized
wt	Weight parameter indicating how much information to borrow across samples using Empirical Bayes
otuthreshold	Abundance cutoff threshold for the OTU to be picked
prevalence	Prevalence of the OTU at threshold cutoff among samples

Value

list containing Empirical Bayes coreOTU Normalized 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)  
countdat <- datlist$countdata  
coreotunormdat <- coreOTUNormalize(countdat)
```

coreOTUQuantile	<i>Compute coreOTU Quantile Normalized data</i>
-----------------	---

Description

Compute coreOTU Quantile Normalized data

Usage

```
coreOTUQuantile(zcounts, otuthreshold = 0.05, prevalence = 0.4)
```

Arguments

zcounts	counts data to be normalized
otuthreshold	Abundance cutoff threshold for the OTU to be picked
prevalence	Prevalence of the OTU at threshold cutoff among samples

Value

list containing coreOTU Quantile Normalized 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)
countdat <- datlist$countdata
coreotunormdat <- coreOTUQuantile(countdat)
```

createPathoStat	<i>Generates a PathoStat object from the PathoScope reports for further analysis using the interactive shiny app</i>
-----------------	--

Description

Generates a PathoStat object from the PathoScope reports for further analysis using the interactive shiny app

Usage

```
createPathoStat(input_dir = ".", sample_data_file = "sample_data.tsv",
  pathoreport_file_suffix = "-sam-report.tsv")
```

Arguments

input_dir	Directory where the tsv files from PathoScope are located
sample_data_file	Sample Data file with information about samples
pathoreport_file_suffix	PathoScope report files suffix

Value

pstat The pathostat object generated from the given tsv files

Examples

```
example_data_dir <- system.file("example/data", package = "PathoStat")
pstat <- createPathoStat(input_dir=example_data_dir,
  sample_data_file="sample_data.tsv")
```

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 <- loadPstat(data_dir, infileName)
ra_otu <- findRAfromCount(phyloseq::otu_table(pstat))
```

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

Description

Find the taxonomy for the given taxon id

Usage

```
findTaxonLevel(tid)
```

Arguments

tid Given taxon id

Value

taxonomy LineageEx

Examples

```
example_data_dir <- system.file("example/data", package = "PathoStat")
pathoreport_file_suffix <- "-sam-report.tsv"
datlist <- readPathoscopeData(example_data_dir, pathoreport_file_suffix)
dat <- datlist$data
ids <- rownames(dat)
tids <- unlist(lapply(ids, FUN = grepTid))
taxonLevel <- findTaxonomy(tids[1])
```

findTaxonMat

Find the Taxonomy Information Matrix

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)
dat <- datlist$data
ids <- rownames(dat)
tids <- unlist(lapply(ids, FUN = grepTid))
taxonLevels <- findTaxonomy(tids[1:5])
taxmat <- findTaxonMat(ids[1:5], taxonLevels)
```

findTaxonomy	<i>Find the taxonomy for each taxon ids</i>
--------------	---

Description

Find the taxonomy for each taxon ids

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)
dat <- datlist$data
ids <- rownames(dat)
tids <- unlist(lapply(ids, FUN = grepTid))
taxonLevels <- findTaxonomy(tids[1:5])
```

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()
```

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

Description

Getter function to get the shinyInputCombat option

Usage

```
getShinyInputCombat()
```

Value

shinyInputCombat option

Examples

```
getShinyInputCombat()
```

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

Description

Getter function to get the shinyInputOrig option

Usage

```
getShinyInputOrig()
```

Value

shinyInputOrig option

Examples

```
getShinyInputOrig()
```

get_core	<i>Select rows of OTU matrix that meet given detection and prevalence thresholds</i>
----------	--

Description

Select rows of OTU matrix that meet given detection and prevalence thresholds

Usage

```
get_core(pstat, detection, prevalence)
```

Arguments

pstat	PathoStat object
detection	An integer specifying the minimum value considered to be "detected"
prevalence	An integer specifying the minimum number of samples that must be detected

Value

Subsetted PathoStat object

get_coremat	<i>Create core OTU matrix containing number of OTUs detected at varying detection and prevalence thresholds.</i>
-------------	--

Description

Create core OTU matrix containing number of OTUs detected at varying detection and prevalence thresholds.

Usage

```
get_coremat(pstat)
```

Arguments

pstat	PathoStat object
-------	------------------

Value

Data frame containing number of OTUs at varying detection and prevalence thresholds, with rows corresponding to number of samples and columns corresponding to detection thresholds. An additional column called "prev" contains the sample threshold for each row.

get_coremat_lineplot	<i>Create line plot from core OTU matrix</i>
----------------------	--

Description

Create line plot from core OTU matrix

Usage

```
get_coremat_lineplot(coremat)
```

Arguments

coremat	Core OTU matrix (data.frame)
---------	------------------------------

Value

Line plot with number of OTUs on the x-axis and detection threshold on the y-axis. Lines connect data points with the same number of samples.

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

```
tid <- grepTid("ti|367928|org|Bifidobacterium_adolescentis_ATCC_15703")
```

loadPathoscopeReports	<i>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.</i>
-----------------------	--

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

reportfiles	Paths to report files
nrows	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)
loadPathoscopeReports(reportfiles)
```

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

Description

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

Usage

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

Arguments

<code>indir</code>	Input Directory of the .rda file
<code>infileName</code>	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

```
example_data_dir <- system.file("example/data", package = "PathoStat")
pathoreport_file_suffix <- "-sam-report.tsv"
datlist <- readPathoscopeData(example_data_dir, pathoreport_file_suffix)
countdat <- datlist$countdata
lcpm <- log2CPM(countdat)
```

pathostat	<i>Build PathoStat-class object from its phyloseq component.</i>
-----------	--

Description

Build PathoStat-class object from its phyloseq component.

Usage

```
pathostat(physeq1)
```

Arguments

physeq1	phyloseq object
---------	-----------------

Value

pstat The pathostat object generated from the given phyloseq object

Examples

```
rich_dense_biom = system.file("extdata", "rich_dense_otu_table.biom",
  package="phyloseq")
phyob <- phyloseq::import_biom(rich_dense_biom)
pstat_biom <- pathostat(phyob)
```

PathoStat-class	<i>PathoStat class to store PathoStat input data including phyloseq object</i>
-----------------	--

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

plotConfRegion	<i>Compute the confidence region for the given proportions</i>
----------------	--

Description

Compute the confidence region for the given proportions

Usage

```
plotConfRegion(p1, p2, size = 100, uselogit = TRUE, n = 10000,
  seed = 1000, jit = FALSE)
```

Arguments

p1	Read counts for first taxon
p2	Read counts for second taxon
size	Total read counts in the sample
uselogit	Use logit transformation to compute confidence region
n	Total number of simulation points to generate
seed	Seed to use in random simulation
jit	jitter option (FALSE by default) for the plot

Value

Confidence region plot

Examples

```
p1 <- 20
p2 <- 25
size <- 200
plotConfRegion(p1, p2, size, uselogit=FALSE)
```

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")
```

Arguments

input_dir Directory where the tsv files from PathoScope are located

pathoreport_file_suffix PathoScope report files suffix

Value

List of final guess relative abundance and count data

Examples

```
example_data_dir <- system.file("example/data", package = "PathoStat")
readPathoscopeData(input_dir=example_data_dir)
```

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_file = "PathoStat_report.html",
             report_dir = ".", report_option_binary = "111111111",
             view_report = FALSE, interactive = TRUE)
```

Arguments

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

Value

outputfile 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(pstat_data)  
outfile <- savePstat(pstat)
```

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 *Setter function to set the shinyInputCombat option*

Description

Setter function to set the shinyInputCombat option

Usage

```
setShinyInputCombat(x)
```

Arguments

x shinyInputCombat option

Value

shinyInputCombat option

Examples

```
setShinyInputCombat(NULL)
```

setShinyInputOrig *Setter function to set the shinyInputOrig option*

Description

Setter function to set the shinyInputOrig option

Usage

```
setShinyInputOrig(x)
```

Arguments

x shinyInputOrig option

Value

shinyInputOrig option

Examples

```
setShinyInputOrig(NULL)
```

sizeNormalize	<i>Normalize the given data based on library size</i>
---------------	---

Description

Normalize the given data based on library size

Usage

```
sizeNormalize(zcounts)
```

Arguments

zcounts Input counts data matrix

Value

accounts Normalized counts data 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)
countdat <- datlist$countdata
accounts <- sizeNormalize(countdat)
```

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