

Package ‘animalcules’

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Title Interactive microbiome analysis toolkit

Version 1.10.0

Description animalcules is an R package for utilizing up-to-date data analytics, visualization methods, and machine learning models to provide users an easy-to-use interactive microbiome analysis framework. It can be used as a standalone software package or users can explore their data with the accompanying interactive R Shiny application. Traditional microbiome analysis such as alpha/beta diversity and differential abundance analysis are enhanced, while new methods like biomarker identification are introduced by animalcules. Powerful interactive and dynamic figures generated by animalcules enable users to understand their data better and discover new insights.

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LazyData false

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alpha_div_boxplot *Alpha diversity boxplot*

Description

Alpha diversity boxplot

Usage

```
alpha_div_boxplot(  
  MAE,  
  tax_level,  
  condition,  
  alpha_metric = c("inverse_simpson", "gini_simpson", "shannon", "fisher", "coverage",  
  "unit")  
)
```

Arguments

MAE	A multi-assay experiment object
tax_level	The taxon level used for organisms
condition	Which condition to group samples
alpha_metric	Which alpha diversity metric to use

Value

A plotly object

Examples

```
data_dir = system.file('extdata/MAE.rds', package = 'animalcules')  
toy_data <- readRDS(data_dir)  
p <- alpha_div_boxplot(toy_data,  
  tax_level = 'genus',  
  condition = 'DISEASE',  
  alpha_metric = 'shannon')  
p
```

alpha_div_test *Get alpha diversity*

Description

Get alpha diversity

Usage

```
alpha_div_test(sam_table, alpha_stat)
```

Arguments

sam_table	A datafram with 2 cols, richness and condition
alpha_stat	Wilcoxon rank sum test or T-test for the test

Value

A dataframe

Examples

```
df_test <- data.frame(richness = seq_len(10),
condition = c(rep(1,5), rep(0,5)))
alpha_div_test(df_test,alpha_stat='Wilcoxon rank sum test')
```

counts_to_logcpm *Covert a counts table to a relative abundances table*

Description

Covert a counts table to a relative abundances table

Usage

```
counts_to_logcpm(counts_table)
```

Arguments

counts_table	A organism x sample data frame of counts
--------------	--

Value

A organism x sample data frame of logcpm counts

Examples

```
logcpm <- counts_to_logcpm(as.data.frame(matrix(seq_len(12),4)))
```

counts_to_relabu

Covert a counts table to a relative abundances table

Description

Covert a counts table to a relative abundances table

Usage

```
counts_to_relabu(counts_table)
```

Arguments

counts_table A organism x sample data frame of counts

Value

A organism x sample data frame of relative abundances

Examples

```
counts_to_relabu(matrix(seq_len(12),4))
```

df_char_to_factor

Factorize all categorical columns

Description

Factorize all categorical columns

Usage

```
df_char_to_factor(df)
```

Arguments

df A sample x condition data frame

Value

A sample x condition data frame

Examples

```
df_char_to_factor(matrix(seq_len(12)))
```

differential_abundance

Differential abundance analysis

Description

Differential abundance analysis

Usage

```
differential_abundance(
  MAE,
  tax_level,
  input_da_condition = c(),
  input_da_condition_covariate = NULL,
  min_num_filter = 5,
  input_da_padj_cutoff = 0.05,
  method = "DESeq2"
)
```

Arguments

MAE	A multi-assay experiment object
tax_level	The taxon level used for organisms
input_da_condition	Which condition is the target condition
input_da_condition_covariate	Covariates added to linear function
min_num_filter	Minimum number reads mapped to this microbe
input_da_padj_cutoff	adjusted pValue cutoff
method	choose between DESeq2 and limma

Value

A output dataframe

Examples

```
data_dir = system.file("extdata/MAE.rds", package = "animalcules")
toy_data <- readRDS(data_dir)
differential_abundance(toy_data,
tax_level="phylum",
input_da_condition=c("DISEASE"),
min_num_filter = 2,
input_da_padj_cutoff = 0.5,
method = "DESeq2")
```

dimred_pca

Dimensionality reduction through PCA

Description

Dimensionality reduction through PCA

Usage

```
dimred_pca(
  MAE,
  tax_level,
  color,
  shape = NULL,
  pcx = 1,
  pcy = 2,
  pcz = NULL,
  datatype = c("logcpm", "relabu", "counts")
)
```

Arguments

MAE	A multi-assay experiment object
tax_level	The taxon level used for organisms
color	A condition to color data points by e.g. "AGE"
shape	A condition to shape data points by e.g. "SEX"
pcx	Principal component on the x-axis e.g. 1
pcy	Principal component on the y-axis e.g. 2
pcz	Principal component on the z-axis e.g. 3
datatype	Datatype to use e.g. c("logcpm", "relabu", "counts")

Value

A list with a plotly object and summary table

Examples

```
data_dir = system.file("extdata/MAE.rds", package = "animalcules")
toy_data <- readRDS(data_dir)
result <- dimred_pca(toy_data,
                      tax_level="genus",
                      color="AGE",
                      shape="DISEASE",
                      pcx=1,
                      pcy=2,
                      datatype="logcpm")
result$plot
result$table
```

dimred_pcoa

Dimensionality reduction through PCoA

Description

Dimensionality reduction through PCoA

Usage

```
dimred_pcoa(
  MAE,
  tax_level,
  color,
  shape = NULL,
  axx = 1,
  axy = 2,
  axz = NULL,
  method = c("bray", "jaccard")
)
```

Arguments

MAE	A multi-assay experiment object
tax_level	The taxon level used for organisms
color	A condition to color data points by e.g. "AGE"
shape	A condition to shape data points by e.g. "SEX"
axx	Principle coordinate on the x-axis e.g. 1
axy	Principle coordinate on the y-axis e.g. 2
axz	Principle coordinate on the z-axis e.g. 2
method	Method to use e.g. c("bray", "jaccard")

Value

A list with a plotly object and summary table

Examples

```
data_dir = system.file("extdata/MAE.rds", package = "animalcules")
toy_data <- readRDS(data_dir)
result <- dimred_pcoa(toy_data,
                       tax_level="genus",
                       color="AGE",
                       shape="DISEASE",
                       axx=1,
                       axy=2,
                       method="bray")
result$plot
result$table
```

dimred_tsne

Dimensionality reduction through t-SNE

Description

Dimensionality reduction through t-SNE

Usage

```
dimred_tsne(
  MAE,
  tax_level,
  color,
  shape = NULL,
  k = c("2D", "3D"),
  initial_dims = 30,
  perplexity = 10,
  datatype = c("logcpm", "relabu", "counts"),
  tsne_cache = NULL
)
```

Arguments

MAE	A multi-assay experiment object
tax_level	The taxon level used for organisms
color	A condition to color data points by e.g. "AGE"
shape	A condition to shape data points by e.g. "SEX"
k	Plot dimensions e.g. c("2D", "3D")

<code>initial_dims</code>	The number of dimensions to use in reduction method
<code>perplexity</code>	Optimal number of neighbors
<code>datatype</code>	Datatype to use e.g. c("logcpm", "relabu", "counts")
<code>tsne_cache</code>	Pass the cached data back into the function

Value

A list with a plotly object and cached data

Examples

```
data_dir = system.file("extdata/MAE.rds", package = "animalcules")
toy_data <- readRDS(data_dir)
results <- dimred_tsne(toy_data,
                        tax_level="phylum",
                        color="AGE",
                        shape="GROUP",
                        k="3D",
                        initial_dims=30,
                        perplexity=10,
                        datatype="logcpm")
results$plot
```

dimred_umap

Dimensionality reduction through PCA

Description

Dimensionality reduction through PCA

Usage

```
dimred_umap(
  MAE,
  tax_level,
  color,
  shape = NULL,
  cx = 1,
  cy = 2,
  cz = NULL,
  n_neighbors = 15,
  metric = c("euclidean", "manhattan"),
  n_epochs = 200,
  init = c("spectral", "random"),
  min_dist = 0.1,
  datatype = c("logcpm", "relabu", "counts"))
)
```

Arguments

MAE	A multi-assay experiment object
tax_level	The taxon level used for organisms
color	A condition to color data points by e.g. "AGE"
shape	A condition to shape data points by e.g. "SEX"
cx	Component on the x-axis e.g. 1
cy	Component on the y-axis e.g. 2
cz	Component on the z-axis e.g. 3
n_neighbors	Number of nearest neighbors
metric	Distance function e.g. c("euclidean", "manhattan")
n_epochs	Number of iterations
init	Initial embedding using eigenvector e.g c("spectral", "random")
min_dist	Determines how close points appear in the final layout
datatype	Datatype to use e.g. c("logcpm", "relabu", "counts")

Value

A list with a plotly object and summary table

Examples

```
data_dir = system.file("extdata/MAE.rds", package = "animalcules")
toy_data <- readRDS(data_dir)
result <- dimred_umap(toy_data,
                       tax_level="genus",
                       color="AGE",
                       shape="DISEASE",
                       cx=1,
                       cy=2,
                       datatype="logcpm")
result$plot
```

diversities

Get alpha diversity

Description

Get alpha diversity

Usage

```
diversities(counts_table, index = "all", zeroes = TRUE)
```

Arguments

- counts_table A dataframe with organism x sample
- index One of inverse_simpson,gini_simpson,shannon,fisher,coverage,unit
- zeroes A boolean for whether to ignore zero values

Value

A list of alpha diversity

Examples

```
diversities(matrix(seq_len(12), nrow = 3), index="shannon")
```

diversities_help	<i>Get alpha diversity</i>
------------------	----------------------------

Description

Get alpha diversity

Usage

```
diversities_help(counts_table, index = "all", zeroes = TRUE)
```

Arguments

- counts_table A dataframe with organism x sample
- index one of inverse_simpson,gini_simpson,shannon,fisher,coverage,unit
- zeroes A boolean for whether to ignore zero values

Value

A list of alpha diversity

Examples

```
diversities_help(matrix(seq_len(12), nrow = 3), index='shannon')
```

diversity_beta_boxplot
Beta diversity boxplot

Description

Beta diversity boxplot

Usage

```
diversity_beta_boxplot(  
  MAE,  
  tax_level,  
  input_beta_method,  
  input_select_beta_condition  
)
```

Arguments

MAE	A multi-assay experiment object
tax_level	The taxon level used for organisms
input_beta_method	bray, jaccard
input_select_beta_condition	Which condition to group samples

Value

A plotly object

Examples

```
data_dir = system.file('extdata/MAE.rds', package = 'animalcules')  
toy_data <- readRDS(data_dir)  
p <- diversity_beta_boxplot(toy_data,  
                           tax_level = 'genus',  
                           input_beta_method = 'bray',  
                           input_select_beta_condition = 'DISEASE')  
p
```

diversity_beta_heatmap

Description

Beta diversity heatmap

Usage

```
diversity_beta_heatmap(  
  MAE,  
  tax_level,  
  input_beta_method,  
  input_bdhm_select_conditions,  
  input_bdhm_sort_by = c("nosort", "conditions"))  
)
```

Arguments

MAE	A multi-assay experiment object
tax_level	The taxon level used for organisms
input_beta_method	bray, jaccard
input_bdhm_select_conditions	Which condition to group samples
input_bdhm_sort_by	Sorting option e.g. "nosort", "conditions"

Value

A `plotly` object

Examples

diversity_beta_test *Beta diversity test (by default we use bray-curtis distance)*

Description

Beta diversity test (by default we use bray-curtis distance)

Usage

```
diversity_beta_test(
  MAE,
  tax_level,
  input_beta_method,
  input_select_beta_condition,
  input_select_beta_stat_method,
  input_num_permutation_permanova = 999
)
```

Arguments

MAE	A multi-assay experiment object
tax_level	The taxon level used for organisms
input_beta_method	bray, jaccard
input_select_beta_condition	Which condition to group samples
input_select_beta_stat_method	PERMANOVA,Kruskal-Wallis,Wilcoxon test
input_num_permutation_permanova	number of permutations

Value

A plotly object

Examples

```
data_dir = system.file('extdata/MAE.rds', package = 'animalcules')
toy_data <- readRDS(data_dir)
p <- diversity_beta_test(toy_data,
  tax_level = 'genus',
  input_beta_method = 'bray',
  input_select_beta_condition = 'DISEASE',
  input_select_beta_stat_method = 'PERMANOVA',
  input_num_permutation_permanova = 999)
p
```

do_alpha_div_test *Alpha diversity statistical test*

Description

Alpha diversity statistical test

Usage

```
do_alpha_div_test(
  MAE,
  tax_level,
  condition,
  alpha_metric = c("inverse_simpson", "gini_simpson", "shannon", "fisher", "coverage",
    "unit"),
  alpha_stat = c("Wilcoxon rank sum test", "T-test", "Kruskal-Wallis")
)
```

Arguments

MAE	A multi-assay experiment object
tax_level	The taxon level used for organisms
condition	Which condition to group samples
alpha_metric	Which alpha diversity metric to use
alpha_stat	Which stat test to use

Value

A dataframe

Examples

```
data_dir = system.file("extdata/MAE.rds", package = "animalcules")
toy_data <- readRDS(data_dir)
p <- do_alpha_div_test(toy_data,
  tax_level = "genus",
  condition = "DISEASE",
  alpha_metric = "shannon",
  alpha_stat = "Wilcoxon rank sum test")
p
```

<code>filter_categorize</code>	<i>Categorize continuous variables</i>
--------------------------------	--

Description

Categorize continuous variables

Usage

```
filter_categorize(
  sam_table,
  sample_condition,
  new_label,
  nbins = NULL,
  bin_breaks = c(),
  bin_labels = c()
)
```

Arguments

<code>sam_table</code>	A sample x condition dataframe
<code>sample_condition</code>	Continuous variable to categorize
<code>new_label</code>	Column name for categorized variable
<code>nbins</code>	Auto select ranges for n bins/categories
<code>bin_breaks</code>	Manually select ranges for bins/categories
<code>bin_labels</code>	Manually label bins/categories

Value

A list with an updated sample table and before/after plots

Examples

```
library(SummarizedExperiment)
data_dir = system.file('extdata/MAE.rds', package = 'animalcules')
toy_data <- readRDS(data_dir)
microbe <- MultiAssayExperiment::experiments(toy_data)[[1]]
samples <- as.data.frame(colData(microbe))
result <- filter_categorize(samples,
  sample_condition = 'AGE',
  new_label='AGE_GROUP',
  bin_breaks=c(0,55,75,100),
  bin_labels=c('Young','Adult','Elderly'))
result$sam_table
result$plot.unbinned
result$plot.binned
```

filter_summary_bar_density

Data visualization by bar plot / density plot

Description

Data visualization by bar plot / density plot

Usage

```
filter_summary_bar_density(  
  MAE,  
  samples_discard = NULL,  
  filter_type,  
  sample_condition  
)
```

Arguments

MAE	A multi-assay experiment object
<code>samples_discard</code>	The list of samples to filter
<code>filter_type</code>	Either 'By Microbes' or 'By Metadata'
<code>sample_condition</code>	Which condition to check e.g. 'SEX'

Value

A `plotly` object

Examples

filter_summary_pie_box

Data visualization by pie chart / box plot

Description

Data visualization by pie chart / box plot

Usage

```
filter_summary_pie_box(  
  MAE,  
  samples_discard = NULL,  
  filter_type,  
  sample_condition  
)
```

Arguments

MAE	A multi-assay experiment object
samples_discard	The list of samples to filter
filter_type	Either 'By Microbes' or 'By Metadata'
sample_condition	Which condition to check e.g. 'SEX'

Value

A plotly object

Examples

```
data_dir = system.file('extdata/MAE.rds', package = 'animalcules')  
toy_data <- readRDS(data_dir)  
result <- filter_summary_pie_box(toy_data,  
  samples_discard = c('subject_2', 'subject_4'),  
  filter_type = 'By Microbes',  
  sample_condition = 'SEX')  
result
```

<code>find_biomarker</code>	<i>Identify biomarkers</i>
-----------------------------	----------------------------

Description

Identify biomarkers

Usage

```
find_biomarker(
  MAE,
  tax_level,
  input_select_target_biomarker,
  nfolds = 3,
  nrepeats = 3,
  seed = 99,
  percent_top_biomarker = 0.2,
  model_name = c("logistic regression", "random forest")
)
```

Arguments

MAE	A multi-assay experiment object
tax_level	The taxon level used for organisms
input_select_target_biomarker	Which condition is the target condition
nfolds	number of splits in CV
nrepeats	number of CVs with different random splits
seed	for repeatable research
percent_top_biomarker	Top importance percentage to pick biomarker
model_name	one of 'logistic regression', 'random forest'

Value

A list

Examples

```
data_dir = system.file('extdata/MAE.rds', package = 'animalcules')
toy_data <- readRDS(data_dir)
p <- find_biomarker(toy_data,
  tax_level='family',
  input_select_target_biomarker=c('DISEASE'),
  nfolds = 3,
  nrepeats = 3,
```

```
    seed = 99,  
    percent_top_biomarker = 0.2,  
    model_name = 'logistic regression')  
p
```

find_taxonomy

Find the taxonomy for unlimited tids

Description

Find the taxonomy for unlimited tids

Usage

```
find_taxonomy(tids)
```

Arguments

tids Given taxonomy ids

Value

A list of taxon levels with information

Examples

```
taxonLevels <- find_taxonomy(tids=1200)
```

find_taxonomy_300

Find the taxonomy for maximum 300 tids

Description

Find the taxonomy for maximum 300 tids

Usage

```
find_taxonomy_300(tids)
```

Arguments

tids Given taxonomy ids

Value

`taxondata` Data with the taxonomy information

Examples

```
taxonLevels <- find_taxonomy_300(tids=1200)
```

find_taxon_mat

Find the Taxonomy Information Matrix

Description

Find the Taxonomy Information Matrix

Usage

```
find_taxon_mat(names, taxonLevels)
```

Arguments

<code>names</code>	Row names of the taxonomy matrix
<code>taxonLevels</code>	Taxon Levels of all tids

Value

`taxmat` Taxonomy Information Matrix

Examples

```
ids <- c("ti|54005", "ti|73001", "ti|573", "ti|228277", "ti|53458")
tids <- c("54005", "73001", "573", "228277", "53458")
taxonLevels <- find_taxonomy(tids)
tax_table <- find_taxon_mat(ids, taxonLevels)
```

gini_simpson	<i>Get alpha diversity using gini</i>
--------------	---------------------------------------

Description

Get alpha diversity using gini

Usage

```
gini_simpson(x)
```

Arguments

x	A list of counts
---	------------------

Value

A single value

Examples

```
gini_simpson(seq_len(10))
```

grep_tid	<i>Grep the tid from the given identifier string</i>
----------	--

Description

Greps the tid from the given identifier string

Usage

```
grep_tid(id)
```

Arguments

id	Given identifier string
----	-------------------------

Value

tid string

Examples

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

inverse_simpson *Get alpha diversity using inverse simpson*

Description

Get alpha diversity using inverse simpson

Usage

```
inverse_simpson(x)
```

Arguments

x A list of counts

Value

A single value

Examples

```
inverse_simpson(seq_len(10))
```

is_categorical *Check if object is categorical*

Description

Check if object is categorical

Usage

```
is_categorical(v)
```

Arguments

v A single value

Value

Boolean

Examples

```
nums <- 2  
is_categorical(nums)
```

is_integer0	<i>check if integer(0)</i>
-------------	----------------------------

Description

check if integer(0)

Usage

is_integer0(x)

Arguments

x	A single value
---	----------------

Value

Boolean

Examples

```
nums <- 2
is_integer0(nums)
```

is_integer1	<i>check if integer(1)</i>
-------------	----------------------------

Description

check if integer(1)

Usage

is_integer1(x)

Arguments

x	A single value
---	----------------

Value

Boolean

Examples

```
nums <- 2
is_integer1(nums)
```

mae_pick_organisms *Modify organisms of multi-assay experiment object*

Description

Modify organisms of multi-assay experiment object

Usage

```
mae_pick_organisms(MAE, isolate_organisms = NULL, discard_organisms = NULL)
```

Arguments

MAE A multi-assay experiment object

isolate_organisms

Isolate specific organisms e.g. ti|001, ti|002

discard_organisms

Discard specific organisms e.g. ti|001, ti|002

Value

A multi-assay experiment object

Examples

```
data_dir = system.file('extdata/MAE.rds', package = 'animalcules')
toy_data <- readRDS(data_dir)
subset <- mae_pick_organisms(toy_data,
isolate_organisms=c('ti|001', 'ti|002'))
```

mae_pick_samples *Modify samples of multi-assay experiment object*

Description

Modify samples of multi-assay experiment object

Usage

```
mae_pick_samples(MAE, isolate_samples = NULL, discard_samples = NULL)
```

Arguments

MAE	A multi-assay experiment object
isolate_samples	Isolate specific samples e.g. c('SAM_01', 'SAM_02')
discard_samples	Discard specific samples e.g. c('SAM_01', 'SAM_02')

Value

A multi-assay experiment object

Examples

```
data_dir = system.file('extdata/MAE.rds', package = 'animalcules')
toy_data <- readRDS(data_dir)
subset <- mae_pick_samples(toy_data,
isolate_samples=c('subject_9',
'subject_14'))
```

pct2str

Converts decimal percentage to string with specified digits

Description

Converts decimal percentage to string with specified digits

Usage

```
pct2str(v, digits = 2)
```

Arguments

v	A single value
digits	number of digits

Value

Boolean

Examples

```
nums <- 0.23
pct2str(nums)
```

percent	<i>Format decimals to percentages</i>
----------------	---------------------------------------

Description

Format decimals to percentages

Usage

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

Arguments

x	An array of decimals
digits	number of digits
format	f

Value

An array of formatted strings

Examples

```
nums <- c(0.42, 0.15, 0.4, 0.563, 0.2)
percent(nums)
```

read_pathoscope_data	<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

```
read_pathoscope_data(
  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

relabu_barplot

Plot bar plots of sample and group level relative abundance

Description

Plot bar plots of sample and group level relative abundance

Usage

```

relabu_barplot(
  MAE,
  tax_level,
  order_organisms = c(),
  sort_by = c("nosort", "conditions", "organisms", "alphabetically"),
  group_samples = FALSE,
  group_conditions = "ALL",
  sample_conditions = c(),
  isolate_samples = c(),
  discard_samples = c(),
  show_legend = TRUE
)

```

Arguments

```

MAE           A multi-assay experiment object
tax_level     The taxon level used for organisms
order_organisms
                  A character list of organisms to send to top
sort_by        Sort bars by one of c("nosort", "conditions", "organisms", "alphabetically")
group_samples  A bool specifying whether to group samples

```

```

group_conditions
    Group by one or more conditions e.g. "ALL" or "SEX"
sample_conditions
    Plot associatied conditions with samples.
isolate_samples
    Isolate specific samples e.g. c("SAM_01", "SAM_02")
discard_samples
    Discard specific samples e.g. c("SAM_01", "SAM_02")
show_legend     A bool specifying whether or not to show organisms legend

```

Value

A plotly object

Examples

```

data_dir = system.file("extdata/MAE.rds", package = "animalcules")
toy_data <- readRDS(data_dir)
p <- relabu_barplot(toy_data,
                     tax_level="family",
                     order_organisms=c('Retroviridae'),
                     sort_by="organisms",
                     sample_conditions=c('SEX', 'AGE'),
                     show_legend=TRUE)
p

```

relabu_boxplot	<i>Plot boxplots comparing different organism prevalence across conditions</i>
-----------------------	--

Description

Plot boxplots comparing different organism prevalence across conditions

Usage

```

relabu_boxplot(
  MAE,
  tax_level,
  condition,
  organisms = c(),
  datatype = c("counts", "relative abundance", "logcpm")
)

```

Arguments

MAE	A multi-assay experiment object
tax_level	The taxon level used for organisms
condition	Compare groups by condition e.g. 'SEX'
organisms	Include organisms for plotting.
datatype	counts, relative abundance,logcpm

Value

A plotly object

Examples

```
data_dir = system.file('extdata/MAE.rds', package = 'animalcules')
toy_data <- readRDS(data_dir)
p <- relabu_boxplot(toy_data,
                     tax_level='genus',
                     organisms=c('Escherichia', 'Actinomyces'),
                     condition='SEX',
                     datatype='logcpm')
p
```

relabu_heatmap

Plot heatmap of sample level counts in logcpm

Description

Plot heatmap of sample level counts in logcpm

Usage

```
relabu_heatmap(
  MAE,
  tax_level,
  sort_by = c("nosort", "conditions", "organisms", "alphabetically"),
  sample_conditions = c(),
  isolate_organisms = c(),
  isolate_samples = c(),
  discard_samples = c(),
  log_cpm = TRUE
)
```

Arguments

MAE	A multi-assay experiment object
tax_level	The taxon level used for organisms
sort_by	Sort bars by one of c('nosort', 'conditions', 'organisms', 'alphabetically')
sample_conditions	Plot conditions e.g. c('SEX', 'AGE')
isolate_organisms	Isolate specific organisms e.g. c('Hepacivirus')
isolate_samples	Isolate specific samples e.g. c('SAM_01', 'SAM_02')
discard_samples	Discard specific samples e.g. c('SAM_01', 'SAM_02')
log_cpm	Convert counts to logcpm

Value

A plotly object

Examples

```
data_dir = system.file('extdata/MAE.rds', package = 'animalcules')
toy_data <- readRDS(data_dir)
p <- relabu_heatmap(toy_data,
                      tax_level='genus',
                      sort_by='conditions',
                      sample_conditions=c('SEX', 'AGE'))
p
```

run_animalcules *Run animalcules shiny app*

Description

Run animalcules shiny app

Usage

```
run_animalcules(dev = FALSE)
```

Arguments

dev	Run the applicaiton in developer mode
-----	---------------------------------------

Value

The shiny app will open

Examples

```
## Not run:  
run_animalcules()  
  
## End(Not run)
```

shannon

Get alpha diversity using shannon

Description

Get alpha diversity using shannon

Usage

```
shannon(x)
```

Arguments

x A list of counts

Value

A single value

Examples

```
shannon(seq_len(10))
```

simpson_index

Get alpha diversity using simpson

Description

Get alpha diversity using simpson

Usage

```
simpson_index(x)
```

Arguments

x A list of counts

Value

A single value

Examples

```
simpson_index(seq_len(10))
```

upsample_counts

Upsample a counts table to a higher taxon level

Description

Upsample a counts table to a higher taxon level

Usage

```
upsample_counts(counts_table, tax_table, higher_level)
```

Arguments

counts_table	A organism x sample data frame of counts
tax_table	A organism x taxlev data frame of labels
higher_level	Higher taxon level to upsample to

Value

A organism x sample data frame of counts

Examples

```
toy_data <- readRDS(system.file("extdata/toy_data.rds", package = "animalcules"))
tax_table <- toy_data$tax_table
sam_table <- toy_data$sam_table
counts_table <- toy_data$counts_table
counts_table <- upsample_counts(counts_table, tax_table, "phylum")
```

<code>write_to_biom</code>	<i>Output biom</i>
----------------------------	--------------------

Description

Output biom

Usage

```
write_to_biom(MAE, path_to_output)
```

Arguments

<code>MAE</code>	A multi-assay experiment object
<code>path_to_output</code>	The folder to output biom file

Value

A message

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