Package 'LorMe'

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Title Lightening One-Code Resolving Microbial Ecology Solution

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Maintainer Ningqi Wang <2434066068@qq.com>

Description Provides a robust collection of functions tailored for microbial ecology analysis, encompassing both data analysis and visualization. It introduces an encapsulation feature that streamlines the process into a summary object. With the initial configuration of this summary object, users can execute a wide range of analyses with a single line of code, requiring only two essential parameters for setup. The package delivers comprehensive outputs including analysis objects, statistical outcomes, and visualization-ready data, enhancing the efficiency of research workflows. Designed with user-friendliness in mind, it caters to both novices and seasoned researchers, offering an intuitive interface coupled with adaptable customization options to meet diverse analytical needs.

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Encoding UTF-8

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Author Ningqi Wang [aut, cre, cph], Yaozhong Zhang [aut], Gaofei Jiang [aut]

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2 Contents

Contents

Index

| Alpha_diversity_calculator | 3 |
|---------------------------------------|---|
| | 4 |
| anova_report | 6 |
| - <i>C</i> - | 8 |
| - | 0 |
| - | 1 |
| | 2 |
| · · · · · · · · · · · · · · · · · · · | 4 |
| 1 –1 | 6 |
| 1- 7 | 7 |
| 1— V | 9 |
| differential_bar | |
| Dimension_reduction | |
| Facet_group | |
| Filter_function | |
| indicator_analysis | |
| kruskal_report | |
| LorMe | |
| manhattan | |
| Module_abundance | |
| Module_composition | |
| nc | |
| NC_remove | |
| network_analysis | |
| network_analysis2 | |
| network_stat | |
| network_visual | |
| network_visual_re | |
| network_withdiff | |
| object_config | |
| structure_plot | |
| sub_tax_summary | |
| tax_summary | |
| testotu | 7 |
| theme_zg | |
| Three_group | 8 |
| Top_taxa | |
| Two_group | |
| t_test_report | |
| volcano_plot | |
| wilcox_test_report | |
| | - |

65

Alpha_diversity_calculator

Calculate alpha diversity based on tax summary object

Description

Calculate alpha diversity for each sample

Usage

```
Alpha_diversity_calculator(taxobj, taxlevel, prefix = "")
```

Arguments

taxobj Configured tax summary objects. See in object_config.

 $tax level \\ tax onomy levels used for visualization. \\ Must be one of c ("Domain", "Phylum", "Class", "Order", "Family", "Class", \\ Tax of the context of$

prefix A character string as prefix of diversity index. Default:""

Value

'Alpha_diversity_calculator' returns alpha-diversity of each sample in format of column table (dataframe) combined with group information in meta file.

```
###data preparation####
data("Two_group")
require(ggplot2)

###analysis####
Alpha_results<- Alpha_diversity_calculator(taxobj = Two_group,taxlevel = "Base")

#Check data frame contained alpha diversity
head(Alpha_results$alphaframe,5)

#Check contained statistics and plot list
names(Alpha_results$plotlist)

#Check statistics for Shannon
Alpha_results$plotlist$Plotobj_Shannon$Statistics

#Extract plot for Shannon
Alpha_results$plotlist$Plotobj_Shannon$Barplot
Alpha_results$plotlist$Plotobj_Shannon$Boxplot
Alpha_results$plotlist$Plotobj_Shannon$Violinplot</pre>
```

Alpha_diversity_calculator2

Calculate alpha diversity based on tax summary object or dataframe table

Description

Calculate alpha diversity of each sample

Usage

```
Alpha_diversity_calculator2(
  taxobj = NULL,
  taxlevel = NULL,
  prefix = "",
  input,
  inputformat,
  reads
)
```

Arguments

tax summary objects computed by tax_summary. Default:NULL.

taxlevel taxonomy levels used for visualization. Must be one of c("Domain", "Phylum", "Class", "Order", "Family", "Order", "Orde

prefix A character string as prefix of diversity index. Default:""

input Reads or relative abundance of OTU/Taxa/gene data frame,see details in input-

format. (Useless when taxobj is set).

inputformat (Useless when taxobj is set) 1:data frame with first column of OTUID and last

column of taxonomy

2:data frame with first column of OTUID/taxonomy

3:data frame of all numeric

reads If the input data frame were from reads table or not(relative abundance ta-

ble).(Useless when taxobj is set).

Value

when tax taxobj is set, returns column table with group information combined with for alphadiversity of each sample, else returns data frame for alpha-diversity of each sample

Note

1. When input data frame is in relative abundance table, Chao and ACE are not available

Author(s)

Wang Ningqi 2434066068@qq.com

```
### Data preparation ####
data(testotu)
groupinformation <- data.frame(</pre>
  group = c(rep("a", 10), rep("b", 10)),
  factor1 = rnorm(10),
  factor2 = rnorm(mean = 100, 10),
  subject = factor(c(1:10, 1:10))
)
# Summary OTU table into genus table and phylum table
testtax_summary <- tax_summary(</pre>
  groupfile = groupinformation,
  inputtable = testotu[, 2:21],
  reads = TRUE,
  taxonomytable = testotu[, c(1, 22)]
)
### Use taxsummary object as input ###
Alpha <- Alpha_diversity_calculator2(</pre>
  taxobj = testtax_summary,
  taxlevel = "Base"
head(Alpha)
# In genus level
Alpha <- Alpha_diversity_calculator2(
  taxobj = testtax_summary,
  taxlevel = "Genus",
  prefix = "Genus"
)
head(Alpha)
### Input dataframe from reads table ###
Alpha <- Alpha_diversity_calculator2(</pre>
  input = testotu,
  prefix = "Bacterial",
  inputformat = 1,
  reads = TRUE
)
### Input dataframe from relative abundance table ###
if (!require(magrittr)) install.packages("magrittr")
library(magrittr)
Alpha <- Filter_function(
  input = testotu,
  threshold = 0,
  format = 1
) %>%
  Alpha_diversity_calculator2(
    input = .,
    prefix = "Bacterial",
```

6 anova_report

```
inputformat = 1,
  reads = FALSE
)
head(Alpha)
```

anova_report

Print Analysis of Variance report

Description

Print Analysis of Variance report

Usage

```
anova_report(
  data,
  treatment_col,
  value_col,
  prior = FALSE,
  comparison_method = "Auto",
  equally_rep = TRUE,
  report = TRUE
)
```

Arguments

data Data frame containing the treatment, value and other information.
treatment_col Numeric indicating where treatment locates (column number) in data.
value_col Numeric indicating where treatment value (column number) in data.

prior logical. Whether conducted prior comparisons.

comparison_method

Default would automaticly choose method. Method of multiple comparison, must

be one of "SNK", "Tukey", "bonferroni", "LSD" or "Scheffe".

equally_rep Logical. Whether all treatments have same number of replication.

report Logical. If print report to console. Default:TRUE

Value

anova_report returns list of:

- 1)basic data description
- 2)ANOVA model
- 3)summary of ANOVA model
- 4)model of multiple comparison
- 5) difference of multiple comparison
- 6)letters of multiple comparison, which could be use for visualization.

anova_report 7

```
#' Data loading from 'agricolae' package
data("cotton", package = "agricolae")
#' ANOVA report with default settings
anova_results <- anova_report(</pre>
 data = cotton,
  treatment_col = 3,
 value\_col = 5
## Here returns NULL because no significance among groups
## To conduct prior comparisons
anova_results <- anova_report(</pre>
 data = cotton,
  treatment_col = 3,
 value\_col = 5,
 prior = TRUE
## Here found no difference among groups, thus change to a more sensitive method
## (maybe illegal, but only as an example)
anova_results <- anova_report(</pre>
 data = cotton,
 treatment_col = 3,
 value\_col = 5,
 prior = TRUE,
 comparison_method = "LSD"
#' Data loading 'iris' dataset
data("iris")
#' ANOVA report for 'iris' dataset
anova_results <- anova_report(</pre>
  data = iris,
  treatment\_col = 5,
 value\_col = 2
### Extract return
### Basic data description
print(anova_results$basicdata)
### ANOVA model
print(anova_results$anova_model)
### Summary of ANOVA model
print(anova_results$anova_summary)
```

8 auto_signif_test

```
### Model of multiple comparison
print(anova_results$multiple_comparison_model)

### Difference of multiple comparison
print(anova_results$comparison_results)

### Letters of multiple comparison, which could be used for visualization
print(anova_results$comparison_letters)
}
```

auto_signif_test

Automatic significance test

Description

Automatically conduct significance testing

Usage

```
auto_signif_test(
  data,
  treatment_col,
  value_col,
  paired,
  subject_col,
  prior = FALSE,
  comparison_method = NULL,
  equally_rep = TRUE,
  output = "console",
  output_dir = "./",
  filename = "auto_signif_test",
  report = TRUE
)
```

Arguments

data Data frame containing the treatment, value and other information.

treatment_col Numeric indicating where treatment locates (column number) in data.

value_col Numeric indicating where treatment value (column number) in data.

paired Logical indicating whether you want a paired t-test.

subject_col Only meaningful when Pair is ture. Numeric indicating where subject of treat-

ment (column number) in data.

prior logical. Whether conducted prior comparisons.

comparison_method

Character string. Only use for more than 2 treatment. Default would automatically choose method. Method of multiple comparison,must be one of "SNK", "Tukey", "bonferroni", "LSD" or "Scheffe".

auto_signif_test 9

| equally_rep | Logical indicating Whether all treatments have same number of replication. |
|-------------|---|
| output | A character string indicating output style. Default: "console", which print the report in console. And "file" is available to output report into text-file. |
| output_dir | Default:"./". Available only when output="file". The direction of output file. |
| filename | A character string indicating file name of output file. Only work when output set as 'file'. |
| report | Logical. If print report to console. Default:TRUE |

Value

auto_signif_test returns results of significant test and print report in console or file. See details in example.

See results return in t_test_report, wilcox_test_report, anova_report, kruskal_report.

Note

1.when choose output="file", once caused error that terminate the program, use 'sink()' to end the written of exist files.

2.Please confirm your data is in format of dataframe, else may cause bug! (e.g. Do not use 'read.xlsx' to load data into tibble format)

```
### Here shows different types of experimental design ###
data("cotton", package = "agricolae")
### Two randomly designed groups ###
sig_results <- auto_signif_test(</pre>
  data = cotton,
  treatment_col = 1,
  value\_col = 5
### Two paired design groups ###
sig_results <- auto_signif_test(</pre>
  data = cotton,
  treatment_col = 1,
  value\_col = 5,
  paired = TRUE,
  subject\_col = 2
)
### More than two randomly designed groups ###
sig_results <- auto_signif_test(</pre>
  data = cotton,
  treatment_col = 2,
  value\_col = 5
head(sig_results) # Check outputs
```

10 circulation_lm

```
### Conduct prior comparisons ###
sig_results <- auto_signif_test(</pre>
  data = cotton,
  treatment_col = 2,
  value\_col = 5,
  prior = TRUE
)
head(sig_results) # Check outputs
print(sig_results$basicdata) # Check statistical summary
print(sig_results$anova_model) # Extract ANOVA model
print(sig_results$anova_summary) # Check ANOVA summary
print(sig_results$multiple_comparison_model) # Extract multiple comparison model
print(sig_results$comparison_results) # Check between-group comparison
print(sig_results$comparison_letters) # Check letters (can be used in visualization)
## Change multiple comparison method (maybe not illegal!!)
sig_results <- auto_signif_test(</pre>
  data = cotton,
  treatment_col = 2,
  value\_col = 5,
  prior = TRUE,
  comparison_method = "LSD"
)
head(sig_results) # Check outputs
print(sig_results$comparison_letters) # Note that letters become different
```

circulation_lm

Circulation of fitting Linear Models

Description

Using circulation to fit linear models between one dependent variable and series of independent variable

Usage

```
circulation_lm(y, xframe, margin)
```

Arguments

y Dependent variable

xframe Matrix or data frame of independent variable

margin A vector of 1 or 2 indicates arrangement of xframe. 1:by rows 2:by columns

Details

if row names(for margin 1) and column names(for margin 2) are not given, ID column of return data frame will be row/column numbers.

color_scheme 11

Value

Data frame contains lm statistics of all Independent Variable

Note

Other arguments used in function lm were set as default. See in 1m.

Author(s)

```
Wang Ningqi2434066068@qq.com
```

Examples

```
data(testotu)
###using margin 1, arrange by rows##
dep=testotu[1,2:21]
in_dep=testotu[-1,2:21]
lm_stat<-circulation_lm(y = dep,xframe = in_dep,margin = 1)
lm_stat
###using margin 2, arrange by column##
dep=testotu[,2]
in_dep=testotu[,3:21]
lm_stat<-circulation_lm(y = dep,xframe = in_dep,margin = 2)
lm_stat</pre>
```

color_scheme

Get color scheme

Description

color_scheme() can generate color scheme from nine color scheme database and expand into color-Ramp

Usage

```
color_scheme(Plan, expand = NULL, names = NULL, show = TRUE)
```

Arguments

| Plan | Character, 'Plan1' to 'Plan10' are optional. |
|--------|---|
| expand | Numeric, default:NULL. Numeric indicating numbers to expand color scheme into colorRamp |
| names | Character string. Names to assign for color scheme. |
| show | Logical. If show assigned color in plot panel. Default:TRUE. |

Value

If parameter 'names' is not given, 'color_scheme' returns character string including color scheme. When 'names' is set, 'color_scheme' returns named vector of color scheme.

Note

1.Parameter 'names' is strongly recommended to assign for fixed color scheme, see details in gg-plot::scale_color_manual

Examples

```
### Commonly used example ###
my_color <- color_scheme(</pre>
  Plan = "Plan1",
  names = c("Treatment1", "Treatment2")
### Generate colorRamp still based on 'Plan1'
my_color <- color_scheme(</pre>
  Plan = "Plan1",
  expand = 4,
  names = c("Treatment1", "Treatment2", "Treatment3", "Treatment4")
)
### View color scheme from plan1 to plan10 in 'Plots' interface ###
color_scheme(Plan = "Plan1")
color_scheme(Plan = "Plan2")
color_scheme(Plan = "Plan3")
color_scheme(Plan = "Plan4")
color_scheme(Plan = "Plan5")
color_scheme(Plan = "Plan6")
color_scheme(Plan = "Plan7")
color_scheme(Plan = "Plan8")
color_scheme(Plan = "Plan9")
color_scheme(Plan = "Plan10")
```

combine_and_translate Combine data for visualization

Description

Combine group information and index into data frame for visualization(scatter, bar plot, alluvial,box plot etc.).

Usage

```
combine_and_translate(inputframe, groupframe, itemname, indexname, inputtype)
```

combine_and_translate

13

Arguments

inputframe Data frame of index ,sample ID in column,requires all numeric(e.g. result from

Alpha_diversity_calculator or Top_taxa function)

groupframe Data frame of group information(and other abiotic/geographic factors)

itemname A character string of your inputframe itemname
indexname A character string of your inputframe indexname

inputtype If sample ID were in row and index in column in inputframe.

Value

key-value pairs data frame

Author(s)

Wang Ningqi2434066068@qq.com

```
{
 require(magrittr)
 data(testotu)
 ## Data preparation ##
 Alpha <- Alpha_diversity_calculator2(
   input = testotu,
   prefix = "Bacterial",
   inputformat = 1,
    reads = TRUE
 )
 topotu <- data.frame(</pre>
   Top_taxa(
      input = testotu,
     n = 10,
     inputformat = 1,
      outformat = 1
   )[, -1],
    row.names = paste0(rep("otu", 11), 1:11)
 groupinformation1 <- data.frame(</pre>
   group = c(rep("a", 10), rep("b", 10)),
   factor1 = rnorm(10),
    factor2 = rnorm(mean = 100, 10)
 ### Use inputtype = FALSE ###
 head(Alpha)
 combine_and_translate(
   Alpha, groupinformation1,
```

14 community_plot

```
itemname = "Alpha", indexname = "index",
  inputtype = FALSE
)

### Use inputtype = TRUE ###
head(topotu)
combine_and_translate(
  topotu, groupinformation1,
  itemname = "OTU", indexname = "reads",
  inputtype = TRUE
)
}
```

community_plot

Generate Community Composition Plot Based on Tax_summary Object

Description

Microbial community composition visualization in format of barplot, areaplot and alluvialplot

Usage

```
community_plot(
  taxobj,
  taxlevel,
  n = 10,
  palette = "Spectral",
  nrow = NULL,
  rmprefix = NULL
)
```

Arguments

| taxobj | Configured tax summary objects. See in object_config. |
|----------|--|
| taxlevel | Character. taxonomy levels used for visualization. Must be one of c("Domain", "Phylum", "Class", "Order", |
| n | Numeric. Top n taxa remained according to relative abundance. Default:10 |
| palette | Character. Palette for visualization,default: "Spectral",recommended to use "Paired" for more than 15 tax. |
| nrow | Numeric. Number of rows when wrap panels, default: NULL. |
| rmprefix | Numeric. Removed prefix character in taxonomy annotation.Default:NULL. See details in example. |

Value

community_plot2 returns three ggplot objects, two data frame used in visualization and one character of filled mapping colors

community_plot 15

Author(s)

Wang Ningqi2434066068@qq.com

```
require(magrittr)
 ### Data preparation ###
 data("Two_group")
 ## Use taxonomy summary objects
 phylum10 <- community_plot(</pre>
   taxobj = Two_group,
   taxlevel = "Phylum",
   n = 10,
   rmprefix = "p__"
 )
 phylum10$barplot # Check bar plot
 phylum10$areaplot # Check area plot
 phylum10$alluvialplot # Check alluvial plot
 phylum10$Top10Phylum %>% head(10) # Check top taxa data frame
 phylum10\$Grouped\_Top10Phylum~\%>\%~head(10)~\#~Check~grouped~top~taxa~data~frame
 print(phylum10$filled_color) # Check mapping colors
 # Double facet
 data("Facet_group")
 # Using palette by default
 phylum10 <- community_plot(</pre>
    taxobj = Facet_group,
   taxlevel = "Phylum",
   n = 10,
   rmprefix = " p__"
 phylum10$barplot
 phylum10$areaplot
 phylum10$alluvialplot
 # Another example
 genus20 <- community_plot(</pre>
   taxobj = Facet_group,
   taxlevel = "Genus",
   n = 20,
   palette = "Paired",
   rmprefix = " g__"
 genus20$alluvialplot
}
```

16 compare_plot

| compare_plot | Comparison plot generator This function help generate comparsion plot including bar plot, box plot, and violin plot |
|--------------|---|
| | ριοι ιπετιατίτες σαν ριοι, σολ ριοι, απα νιοιτίτ ριοι |

Description

Comparison plot generator This function help generate comparsion plot including bar plot, box plot, and violin plot

Usage

```
compare_plot(
  inputframe,
  treat_location,
  value_location,
  aes_col = NULL,
  point = TRUE,
  facet_location = NULL,
  ylab_text = NULL
)
```

Arguments

```
inputframe A data frame contain information for visualization.

treat_location Numeric. Treatment column number in inputframe.

value_location Numeric. Value column number in inputframe.

aes_col Named character string, default:NULL. A set of aesthetic character to map treatment to.

point Logical. If draw point on bar, box and violin plot. Default:TRUE.

facet_location Numeric, default:NULL. Facet column number in inputframe.

ylab_text Character. Text for y axis.
```

Value

A list contained plot and statistics

```
results$Boxplot
results$Violinplot
iris$Treat2=rep(c(rep("A",25),rep("B",25)),3)
results=compare_plot(inputframe=iris,treat_location=5,
                     value_location=1,facet_location = 6,
                     ylab_text = "Sepal Length")
#Check statistics
results$Statistics
#Extract plot
results$Barplot
results$Boxplot
results$Violinplot
#Extract combined plot
results$All_Barplot
results$All_Boxplot
results$All_Violinplot
```

Deseq_analysis

Deseq Analysis Function

Description

This function performs a differential expression analysis using the DESeq2 package. It is designed to work with microbiome data and can handle paired or non-paired samples.

Usage

```
Deseq_analysis(
  taxobj,
  taxlevel,
  comparison = NULL,
  cutoff,
  control_name,
  paired = FALSE,
  subject = NULL
)
```

Arguments

taxlevel

taxobj Configured tax summary objects. See in object_config.

The state of the s

The taxonomic level for the analysis. Must be one of c("Domain", "Phylum", "Class", "Order", "Family", "Ge

comparison A vector of conditions to compare. Default: NULL, all unique conditions are

compared (only for Two groups).

| cutoff | The log2 fold change cutoff for considering as differential taxon. |
|--------------|--|
| control_name | Character. The name of the control group for the comparison. |
| paired | Logical. Should the samples be treated as paired? Default: False |
| subject | Optional. The subject identifier for paired samples. Default: Null |

Value

A data frame with the results of the differential expression analysis.

Note

- 1. Regulation is judged by cutoff of q-value(adjust p value). Detail see in DESeq
- 2. For more than two groups in taxobj, the 'comparison' must be assigned.
- 3. The function requires the 'DESeq2', 'S4Vectors', and 'tibble' packages.

Author(s)

```
Wang Ningqi 2434066068@qq.com
```

See Also

```
DESeqDataSetFromMatrix, DESeq, DataFrame, as_tibble
```

```
if (requireNamespace("DESeq2", quietly = TRUE) &&
    requireNamespace("S4Vectors", quietly = TRUE) &&
    requireNamespace("tibble", quietly = TRUE)) {
    ### Data preparation ###
   data("Two_group")
    ### Deseq analysis ###
   deseq_results <- Deseq_analysis(</pre>
      taxobj = Two_group,
      taxlevel = "Genus",
      cutoff = 1,
      control_name = "Control"
   # Visualization of volcano plot ##
   volcano_plot <- volcano_plot(</pre>
      inputframe = deseq_results,
      cutoff = 1,
      aes_col = Two_group$configuration$treat_col
   volcano_plot$FC_FDR
   volcano_plot$Mean_FC
    # Visualization of Manhattan plot ##
```

```
manhattan_object <- manhattan(</pre>
    inputframe = deseq_results,
    taxlevel = "Phylum",
    control_name = "Control",
    mode = "most",
    top_n = 10,
    rmprefix = "p__"
 )
 manhattan_object$manhattan # Tradition manhattan plot
 manhattan_object$manhattan_circle # Circular manhattan plot
  # For object with more than two groups
  ### Data preparation ###
  data("Three_group")
  # Specific comparison
  deseq_results_BFCF <- Deseq_analysis(</pre>
    taxobj = Three_group,
    taxlevel = "Genus",
    comparison = c("BF", "CF"),
    cutoff = 1,
    control_name = "CF"
 )
  volcano_plot <- volcano_plot(</pre>
    inputframe = deseq_results_BFCF,
    cutoff = 1,
    aes_col = Three_group$configuration$treat_col
 volcano_plot$FC_FDR
} else {
 message(
    "The 'DESeq2', 'S4Vectors', and/or 'tibble' package(s) are not installed. ",
    "Please install them to use all features of Deseq_analysis."
 )
}
```

Deseq_analysis2

Deseq analysis

Description

Deseq analysis

Usage

Deseq_analysis2(inputframe, condition, cutoff, control_name, paired, subject)

Arguments

inputframe Otu/gene/taxa table with all integer numeric variables.Rownames must be Otu/gene/taxa

names, colnames must be sample names with control in front and treatment be-

hind. Reads table is recommended.

condition A character string which indicates group of samples cutoff threshold of log2(Foldchange).Detail see in DESeq control_name A character indicating the control group name

paired Logical to determine if paired comparision would be used. TRUE or FALSE.

subject A character string which indicates paired design of samples

Value

Statistics dataframe of all otu/gene/taxa

Note

- 1. Inputframe must be all integer numeric variables without NA/NAN/inf! In case your data is not an integer one,a practical method is to multiply them in equal proportion(eg. x 1e6) then round them into integer
- 2. Regulation is judged by cutoff of qvalue(adjust p value). Detail see in DESeq
- 3. Set cutoff as 1 is recommened. In case of too few taxa(eg. Phylum level deseq), cutoff can be set to 0.
- 4. if control_name is not given, the control group will be set according to ASCII
- 5. The function requires the 'DESeq2', 'S4Vectors', and 'tibble' packages.

Author(s)

Wang Ningqi 2434066068@qq.com

See Also

DESeqDataSetFromMatrix, DESeq, DataFrame, as_tibble

```
### Data preparation ###
data(testotu)
rownames(testotu) <- testotu[, 1]
inputotu <- testotu[, -c(1, ncol(testotu))]
head(inputotu)
group <- c(rep("a", 10), rep("b", 10))

### DESeq analysis ###
if (requireNamespace("DESeq2", quietly = TRUE) &&
    requireNamespace("S4Vectors", quietly = TRUE) &&
    requireNamespace("tibble", quietly = TRUE)) {</pre>
```

differential_bar 21

```
Deseqresult <- Deseq_analysis2(</pre>
      inputframe = inputotu,
      condition = group,
      cutoff = 1,
      control_name = "b"
    ### Paired DESeq analysis ###
    subject <- factor(c(1:10, 1:10))</pre>
   Deseqresult <- Deseq_analysis2(</pre>
      inputframe = inputotu,
      condition = group,
      cutoff = 1,
      control_name = "b",
      paired = TRUE,
      subject = subject
 }
}
```

differential_bar

Generate Differential Bar Plot and Error bar Plot

Description

Generate Differential Bar Plot and Error bar Plot

Usage

```
differential_bar(
  taxobj,
  taxlevel,
  comparison = NULL,
  rel_threshold = 0.005,
  anno_row = "taxonomy",
  aes_col = NULL,
  limit_num = NULL
)
```

Arguments

taxobj Configured tax summary objects. See in object_config.

taxlevel Taxonomy levels used for visualization. Must be one of c("Domain", "Phylum", "Class", "Order", "Family",

comparison A vector of conditions to compare. Default: NULL, all unique conditions are

compared (only for Two groups).

rel_threshold Threshold filtering taxa for differential analysis. Default:0.005

22 differential_bar

| anno_row | Default: 'taxonomy'. Rownames for visualization. Options are 'taxonomy' for showing taxonomic information and 'ID' for showing taxonomic ID. |
|-----------|--|
| aes_col | A named vector of colors to be used in the plots. |
| limit_num | Numeric. The maximum number of significant results to display. Default: NULL, showing all differential taxa. |

Value

A list containing the bar plot, source data for the bar plot, difference plot, and source data for the difference plot.

Note

The differential analysis is performed using two-sided Welch's t-test. The p-values are adjusted using the 'BH' (i.e., FDR) method.

```
{
  # Data preparation
  data("Two_group")
  # Simple mode
  diff_results <- differential_bar(</pre>
   taxobj = Two_group,
   taxlevel = "Genus"
  print(diff_results$Barplot) # Print Barplot
  head(diff_results$Barplot_sourcedata) # Show source data of barplot
  print(diff_results$Differenceplot) # Print Differential errorbar plot
 head(diff_results$Differenceplot_sourcedata) # Show source data of Differential errorbar plot
  require(patchwork)
  diff_results$Barplot|diff_results$Differenceplot
  # Displaying ID
  diff_results <- differential_bar(</pre>
    taxobj = Two_group,
    taxlevel = "Base",
    anno_row = "ID"
  print(diff_results$Barplot)
  # Threshold adjustment
  diff_results <- differential_bar(</pre>
    taxobj = Two_group,
    taxlevel = "Base",
    rel_{threshold} = 0.001
  print(diff_results$Barplot)
  # Limit the displaying number
```

Dimension_reduction 23

```
diff_results <- differential_bar(</pre>
    taxobj = Two_group,
    taxlevel = "Base",
    rel_threshold = 0.001,
   limit_num = 10
 print(diff_results$Barplot)
 # For object with more than two groups
 # Data preparation
 data("Three_group")
 # Specific comparison
 Three_group_col <- Three_group$configuration$treat_col</pre>
 diff_results <- differential_bar(</pre>
    taxobj = Three_group,
    taxlevel = "Genus",
    comparison = c("BF", "CF"),
    aes_col = Three_group_col
 print(diff_results$Barplot)
}
```

Dimension_reduction

Dimension_reduction: PCA, PCOA, and NMDS Analysis

Description

Performs dimension reduction analysis using PCA, PCOA, or NMDS.

Usage

```
Dimension_reduction(inputframe, group, format)
```

Arguments

 $input frame \qquad \quad An \ OTU/gene/taxa \ table \ with \ all \ numeric \ variables \ and \ no \ NA/NAN/inf \ values.$

group Group information with the sample order the same as in inputframe. The format of analysis: 1 for PCA, 2 for PCOA, 3 for NMDS.

Value

A list containing data frames and other statistics for dimension reduction analysis.

Note

Inputframe should be a numeric matrix without NA/NAN/inf values.

The row names of inputframe should be set as OTU/gene/taxa annotations for further analysis.

The results are combined into a list for output. Use as.data.frame(result[[1]]) to extract the data frame, and \$result\$ to extract other statistics. See examples for details.

24 Facet_group

Author(s)

Wang Ningqi 2434066068@qq.com

Examples

```
### Data preparation ###
data(testotu)
rownames(testotu) <- testotu[, 1]</pre>
inputotu <- testotu[, -c(1, ncol(testotu))]</pre>
head(inputotu)
groupinformation1 <- data.frame(</pre>
  group = c(rep("a", 10), rep("b", 10)),
  factor1 = rnorm(10),
  factor2 = rnorm(mean = 100, 10)
)
### PCA ###
PCAresult <- Dimension_reduction(inputotu, groupinformation1, 1)</pre>
PCAframe <- PCAresult$outframe # Extract data for visualization
head(PCAresult$data.pca$rotation,5) # OTU coordinates
### PCOA ###
PCOAresult <- Dimension_reduction(inputotu, groupinformation1, 2)</pre>
PCOAframe <- PCOAresult$outframe # Extract data for visualization
head(PCOAresult$PCOA$values,2) # Explanation of first two axis
### NMDS ###
NMDSresult <- Dimension_reduction(inputotu, groupinformation1, 3)</pre>
NMDSframe <- NMDSresult$outframe # Extract data for visualization
# Here we got a warning of `stress is (nearly) zero: you may have insufficient data`,
# so make sure you have sufficient data for NMDS
print(NMDSresult$NMDSstat$stress) # Extract stress of NMDS
```

Facet_group

Tax summary object with Facet 2x2 Groups

Description

Enraptured summary object with facet 2x2 Groups. Configuration has been assigned.

Usage

Facet_group

Format

Tax summary object with configuration

Filter_function 25

| Filter_function Filter OTU/ASV/metagenomic profile/gene profile by threshold | |
|--|--|
|--|--|

Description

Sequenced data of taxonomy&gene still remains some sequencing error which we needed to be wiped off before analyzing. Here we provide function including four formats to wipe them clean.

Usage

```
Filter_function(input, threshold, format, report = TRUE)
```

Arguments

| 8 | |
|-----------|---|
| input | Data frame of absolute abundance of standard OTU table, with the first column of OTUID and the final column of taxonomy annotation. If your data frame is gene table or not a standard OTU table, please manually transformed into a standard input data frame. |
| threshold | threshold of filter. Relative abundance for format 1 and 4, reads number for format 2, sample size for format 3 $$ |
| format | 1:filter OTU/gene below overall-sample relative abundance threshold(<) 2:filter OTU/gene below overall-sample reads threshold(<) 3:filter OTU/gene reads 0 over threshold sample size(>) 4:filter OTU/gene below relative abundance threshold in each sample(<) |
| report | Logical. If print report to console. Default:TRUE |
| | |

Value

Dataframe of OTU/gene in format of absolute abundacne(reads) or relative abundance(%)

Author(s)

Wang Ningqi

```
### Data frame with absolute abundance (reads)###
### And first column of OTUID and last column of taxonomy ###
data(testotu)

#### If your data frame does not contain the OTUID column or taxonomy column,
#### you can add a simulated column to fit the input format like testotu ##

### 1. Filter OTU with total relative abundance below 0.0001###
filtered_otu <- Filter_function(
   input = testotu,
   threshold = 0.0001,</pre>
```

26 indicator_analysis

```
format = 1
)
### 2. Filter OTU with total reads below 20 ###
filtered_otu <- Filter_function(</pre>
 input = testotu,
 threshold = 20,
 format = 2
)
### 3. Filter OTU reads 0 over (>=) 11 samples ###
filtered_otu <- Filter_function(</pre>
 input = testotu,
 threshold = 11,
 format = 3
)
### 4. Filter OTU with relative abundance below 0.0001 in each sample ###
filtered_otu <- Filter_function(</pre>
 input = testotu,
 threshold = 0.0001,
 format = 4
)
```

indicator_analysis

Indicator Analysis

Description

Performs the indicator analysis based on taxonomic summary object

Usage

```
indicator_analysis(taxobj, taxlevel, func = "r.g", reads = FALSE)
```

Arguments

taxobj Configured tax summary objects. See in object_config.

taxlevel taxonomy levels used for visualization. Must be one of c("Domain", "Phylum", "Class", "Order", "Family", "Gunc Default: "r.g". The function to use for the indicator analysis, see in multipatt

A logical value indicating whether the input data is in terms of raw reads (TRUE)

or relative abundance (FALSE)

Value

A data frame with the results of the indicator analysis, including adjusted p-values, tags and taxonomic information.

kruskal_report 27

Note

This function depends on the following packages: indicspecies, permute. These packages are not automatically loaded and should be installed before using this function.

See Also

```
multipatt, how
```

Examples

```
data("Two_group")
if (requireNamespace("indicspecies", quietly = TRUE) &&
    requireNamespace("permute", quietly = TRUE)) {
    set.seed(999)
    indicator_results <- indicator_analysis(
        taxobj = Two_group,
        taxlevel = "Genus"
    )
    head(indicator_results)
}</pre>
```

kruskal_report

Print Kruskal-Wallis Rank Sum Test report

Description

Print Kruskal-Wallis Rank Sum Test report

Usage

```
kruskal_report(
  data,
  treatment_col,
  value_col,
  prior = FALSE,
  comparison_method = "Auto",
  equally_rep = TRUE,
  report = TRUE
)
```

Arguments

data Data frame containing the treatment, value and other information.

treatment_col Numeric indicating where treatment locates (column number) in data.

value_col Numeric indicating where treatment value (column number) in data.

prior logical. Whether conducted prior comparisons.

28 kruskal_report

comparison_method

Default would automaticly choose method. Method of multiple comparison, must

be one of "SNK" or "Tukey".

equally_rep Logical. Whether all treatments have same number of replication.

report Logical. If print report to console. Default:TRUE

Value

kruskal_report returns list with

1)basic data description

2)summary of Kruskal-Wallis Rank Sum Test

3)model of multiple comparison

4) difference of multiple comparison

5)letters of multiple comparison, which could be use for visualization.

```
data("cotton",package ="agricolae" )
kruskal_results=kruskal_report(data = cotton,treatment_col =3,value_col = 5)
##here returns NULL because no significance among groups
##to conduct prior comparisons.
kruskal_results=kruskal_report(data = cotton,treatment_col =3,value_col = 5,prior = TRUE)
data("iris")
kruskal_results=kruskal_report(data = iris,treatment_col = 5,value_col = 2)
###extract return##
###basic data description
kruskal_results$basicdata
###summry of Kruskal-Wallis Rank Sum Test
kruskal_results$Kruskal_Wallis_summary
###model of multiple comparision
kruskal_results$muiltiple_comparision_model
###difference of multiple comparision
kruskal_results$comparision_results
###letters of multiple comparision, which could be use for visualization.
kruskal_results$comparison_letters
```

LorMe 29

| LorMe | LorMe package: Lightening One-Code Resolving Microbial Ecology Program |
|-------|---|
| | |

Description

LorMe package summarizes a series of functions normally used in microbiome analysis analysis.

Details

```
_PACKAGE
```

#Basic functions####

auto_signif_test Automatically conduct significance testing

compare_plot Comparison plot generator

Filter_function Filter OTU/ASV/metagenomic profile/gene profile by threshold

tax_summary Encapsulate meta file, feature tables and taxonomy annotation into tax summary object

sub_tax_summary subsets tax summary objects according to meta file

combine_and_translate Combine feature table with meta file and transform into a recognizable data frame for visualization.

color_scheme generate color scheme from nine color scheme database and expand into colorRamp theme_zg A classic theme for ggplot.

#Community features####

Alpha_diversity_calculator Calculator for alpha diversity of each sample.

Dimension_reduction Dimension reduction analysis including PCA,PCOA and NMDS

structure_plot A fast view of microbial structure with PCA plot,PCOA plot and NMDS plot.

Top_taxa Calculate most abundant taxon

community_plot A fast view of microbial community with bar plot, alluvial plot and area plot.

#Differential analysis####

Deseq_analysis Performs a differential expression analysis

indicator_analysis Performs the indicator analysis based on taxonomic summary object

differential_bar Generate Differential Bar Plot and errorbar plot

volcano_plot Generate volcano plot base on Deseq_analysis or indicator_analysis results

manhattan Generate Manhattan Plot base on Deseq_analysis or indicator_analysis results

#Network analysis####

network_analysis A convenient and fast network analysis function, with output results suitable for cytoscape and gephi

network_withdiff Meta network analysis integrating differential taxon into a network analysis network_visual Visualizes a network based on network object from network_analysis

30 manhattan

```
network_visual_re Re-visualize or adjust network plot from network_visual or network_withdiff
Module_composition Pie chart for network module composition
Module_abundance Calculate network module abundance for each sample
nc Calculate network Natural Connectivity
NC_remove Conduct natural connectivity analysis
#Correlation analysis###
circulation_lm Quick test using circulation to fit linear models between one dependent variable
```

and series of independent variable tbRDA_analysis RDA analysis including co-linearity diagnostics and necessary statistics.

Author(s)

Wang Ningqi

manhattan

Manhattan Plot Generator

Description

Generate Manhattan Plot base on Deseq_analysis or indicator_analysis results

Usage

```
manhattan(
  inputframe,
  taxlevel = "Phylum",
  control_name,
  mode = "all",
  top_n = NULL,
  palette = "Set1",
  select_tax = NULL,
  rmprefix = NULL
)
```

Arguments

select_tax

rmprefix

| 9 | |
|--------------|---|
| inputframe | A data frame generated from Deseq_analysis or indicator_analysis |
| taxlevel | Taxonomy levels used for visualization. Must be one of c("Domain", "Phylum", "Class", "Order", "Family", " |
| control_name | Character. The name of the control group for the comparison. |
| mode | The mode for selecting which taxa to plot: "all" for all taxa, "most" for the top N taxa, and "select" for specific taxa selection |
| top_n | The number of top taxa to plot when mode is set to "most" |
| palette | Character. Palette for visualization, default: "Set1". Optional palette same as 'RColorBrewer'. "Plan1" to "Plan10" were also optional, see in color_scheme |

A vector of taxa to be selected for plotting when mode is "select".

A string prefix to be removed from the taxonomic annotation. Default: NULL.

manhattan 31

Value

a list containing the Manhattan plot, circular Manhattan plot, source data, and color assignments

```
# Data preparation
data("Two_group")
# DESeq analysis
deseq_results <- Deseq_analysis(</pre>
  taxobj = Two_group,
  taxlevel = "Base",
 cutoff = 1,
  control_name = "Control"
)
# Indicator analysis
indicator_results <- indicator_analysis(</pre>
  taxobj = Two_group,
  taxlevel = "Genus"
# Show all with Manhattan plot
 manhattan_object <- manhattan(</pre>
    inputframe = deseq_results,
    taxlevel = "Phylum",
    control_name = "Control"
 print(manhattan_object$manhattan) # Tradition Manhattan plot
 print(manhattan_object$manhattan_circle) # Circular Manhattan plot
 print(manhattan_object$sourcedata) # Source data for plot
 print(manhattan_object$aes_color) # Aesthetic color for plot
# Top 8 Phyla with most taxon
  manhattan_object <- manhattan(</pre>
    inputframe = indicator_results,
    taxlevel = "Phylum",
    control_name = "Control",
    mode = "most",
    top_n = 8,
    palette = "Set1"
 print(manhattan_object$manhattan)
# Specific phyla
# Top nine dominant phyla
  community <- community_plot(</pre>
    taxobj = Two_group,
    taxlevel = "Phylum",
    n = 9,
    palette = "Paired",
```

32 Module_abundance

```
rmprefix = "p__"
)

manhattan_object <- manhattan(
   inputframe = indicator_results,
   taxlevel = "Phylum",
   control_name = "Control",
   mode = "select",
   palette = community$filled_color,
   select_tax = names(community$filled_color),
   rmprefix = "p__"
)
   print(manhattan_object$manhattan)
   print(manhattan_object$manhattan_circle)
}</pre>
```

Module_abundance

Calculate network module abundance for each sample

Description

Calculate network module abundance for each sample

Usage

```
Module_abundance(network_obj, No.module)
```

Arguments

network_obj Network analysis results generated from network_analysis

No.module Numeric or numeric vector of No.module

Value

A list containing module abundance in metafile and column table of corresponding data frame

```
#data preparation
data("Two_group")
##network analysis
network_results<- network_analysis(taxobj = Two_group,taxlevel = "Genus",n = 10,threshold = 0.8)
require(ggplot2)
#one module
moduleframe=Module_abundance(network_obj =network_results,No.module = 3 )
moduleframe$rowframe #combine into metafile
moduleframe$columnframe #column table
#statistics</pre>
```

Module_composition 33

```
moduleframe$plotlist$Plotobj_Module3$Statistics
#extract plot
moduleframe$plotlist$Plotobj_Module3$Barplot
moduleframe$plotlist$Plotobj_Module3$Boxplot
moduleframe$plotlist$Plotobj_Module3$Violinplot

#multiple modules
moduleframe=Module_abundance(network_results,c(1,3,6))
moduleframe$rowframe
moduleframe$columnframe #column table can be used in ggplot visualization
#same as above to extract plots and statistics
moduleframe$plotlist$Plotobj_Module6$Barplot
```

Module_composition

Pie chart for network module composition

Description

This function analyzes the composition of modules within a network object, providing a visual and data summary based on taxonomic levels.

Usage

```
Module_composition(
  network_obj,
  No.module,
  taxlevel = "Phylum",
  mode = "all",
  top_n = NULL,
  palette = "Set1",
  select_tax = NULL,
  rmprefix = NULL
)
```

Arguments

| network_obj | Network analysis results generated from network_analysis |
|-------------|--|
| No.module | Numeric or numeric vector of No.module |
| taxlevel | Taxonomy levels used for visualization. Must be one of c("Domain", "Phylum", "Class", "Order", "Family", " |
| mode | The mode for selecting which taxa to plot: "all" for all taxa, "most" for the top N taxa, and "select" for specific taxa selection |
| top_n | The number of top taxa to plot when mode is set to "most" |
| palette | Character. Palette for visualization,default: "Set1". See optional palette in same |

as 'RColorBrewer'. And "Plan1" to "Plan10" were also optional, see in color_scheme select_tax

A vector of taxa to be selected for plotting when mode is "select".

rmprefix A string prefix to be removed from the taxonomic annotation

Module_composition

Value

34

The function returns a list containing pie chart of specific module, corresponding source data and color assignments

```
#Data loading
data("Two_group")
# Network analysis
network_Two_group <- network_analysis(</pre>
   taxobj = Two_group,
   taxlevel = "Genus",
  reads = TRUE,
  n = 8,
  threshold = 0.7
 )
 # Show all taxa
 module_results <- Module_composition(</pre>
   network_obj = network_Two_group,
  No.module = c(2, 5),
   taxlevel = "Phylum"
 )
 print(module_results$Module5$Pie)
 print(module_results$Module2$Pie) # View pie chart
 head(module_results$Module2$source_data_Module2) # View source data for pie chart
 print(module_results$aes_color) # Check aesthetic color
 # Show taxa with top five frequency
 module_results <- Module_composition(</pre>
  network_obj = network_Two_group,
  No.module = c(2, 5),
   taxlevel = "Phylum",
  mode = "most",
   top_n = 5
 )
 print(module_results$Module2$Pie_plot_Module2)
 # Show specific taxa
 community <- community_plot(</pre>
   taxobj = Two_group,
  taxlevel = "Phylum",
  n = 5,
  palette = "Paired"
 ) # Get top 5 dominant phyla
 top5_phyla <- names(community$filled_color)</pre>
 module_results <- Module_composition(</pre>
  network_obj = network_Two_group,
  No.module = c(2, 5),
   taxlevel = "Phylum",
```

nc 35

```
mode = "select",
 palette = community$filled_color,
  select_tax = top5_phyla
print(module_results$Module2$Pie_plot_Module2)
# Specific taxa with no prefix 'p__'
module_results <- Module_composition(</pre>
  network_obj = network_Two_group,
 No.module = 2,
  taxlevel = "Phylum",
 mode = "select",
  select_tax = c("Proteobacteria", "Actinobacteria")
print(module_results$Module2$Pie_plot_Module2)
# Remove 'p__' prefix
module_results <- Module_composition(</pre>
 network_obj = network_Two_group,
 No.module = 2,
 taxlevel = "Phylum",
 mode = "most",
  top_n = 5,
 palette = "Set2",
 rmprefix = "p__"
print(module_results$Module2$Pie_plot_Module2)
```

Calculate Network Natural Connectivity

nc

Description

Calculate Network Natural Connectivity

Usage

```
nc(adj_matrix)
```

Arguments

adj_matrix Adjacency data frame or matrix. Can be calculated from network_analysis

Value

Numeric value of natural connectivity

NC_remove

Examples

```
{
    ### Data preparation ###
    data("Two_group")

### One input network analysis ###
    network_results <- network_analysis(
    taxobj = Two_group,
    taxlevel = "Base",
    reads = FALSE,
    n = 10,
    threshold = 0.6
)

# Convert network results to a data frame for the adjacency matrix
    network_matrix <- as.data.frame(network_results[[3]]) # Complete adjacency matrix

# Check initial natural connectivity
    nc_initial <- nc(network_matrix)
    print(nc_initial) # Print the initial natural connectivity
}</pre>
```

NC_remove

Natural connectivity analysis

Description

Natural connectivity analysis

Usage

```
NC_remove(input, num, seed = 1)
```

Arguments

input Network adjacency matrix. Can be generated by network_analysis

num Max number of removed nodes. Default: Automatically match max number that

can be removed.

seed Random seed Number to be set. Default: 1. See in set. seed

Value

NC_remove returns data frame with removed nodes and corresponding natural connectivity

Author(s)

Wang Ningqi2434066068@qq.com

network_analysis 37

Examples

```
{
  ### Data preparation ###
  data("Two_group")
  ### One input network analysis ###
  network_results <- network_analysis(</pre>
    taxobj = Two_group,
    taxlevel = "Base",
    reads = FALSE,
    n = 10,
    threshold = 0.6
  network_matrix <- as.data.frame(network_results[[3]]) # Complete adjacency matrix</pre>
  # Check initial natural connectivity
  nc <- nc(network_matrix)</pre>
  # Conduct natural connectivity analysis
  nc_remove <- NC_remove(input = network_matrix)</pre>
  head(nc_remove)
  tail(nc_remove)
  # Set target number for natural connectivity analysis
  nc_remove <- NC_remove(input = network_matrix, num = 400)</pre>
}
```

network_analysis

Conduct Network analysis based on tax summary object

Description

Conduct Network analysis based on tax summary object

Usage

```
network_analysis(
  taxobj,
  taxlevel,
  reads = FALSE,
  n,
  threshold,
  rel_threshold = 0,
  method = "spearman",
  display = TRUE
)
```

38 network_analysis

Arguments

| taxobj | tax summary objects computed by tax_summary. |
|---------------|---|
| taxlevel | taxonomy levels used for analysis. Must be one of c("Domain","Phylum","Class","Order","Family","Gen |
| reads | Logical,default:FALSE. Taxonomy abundance type used in analysis.FALSE for relative abundance, TRUE for absolute abundance. |
| n | Numeric. Number of sample size indicating kept asv/otu/gene/taxa appearing. Recommended to set more than half of total sample size. |
| threshold | Numeric.Threshold of absolute correlation value (r value for pearson method and rho value for spearman method). |
| rel_threshold | Numeric.Threshold of relative abundance included in the network analysis.Default:0 |
| method | Character, default: "spearman". A character indicating which correlation coefficient method to be computed. One of "pearson" or "spearman" |
| display | Logical, default:TRUE. If display a preview plot of network based on igraph. FALSE for the first attempt is recommended in case of too many vertices and edges. |

Details

- 1. We had optimized the correlation algorithm to achieve a faster running speed. It takes less than 2 minute to calculate dataframe correlation and p value which more than 400 samples and 10000 OTUs for computer with dual Core i5 processor. However, too many vertices(>2000) or links(>10000) may slow the statistical process and visualization,so we recommend that in your first attempt,set display paramter as F to have a preview. Then you can adjust your n/threshold/method paramter to generate a suitable visualization network
- 2. We display a preview plot so as to adjusting your network. Generally a global figure (like we show in examples) with less than 1000 vertices and 5000 edges/links is recommended. Further more, we recommend you to output the statistics and adjacency table and use software like cytoscape or gephi for better visualization.

Value

One list contains nodes information table, adjacency column table, adjacency matrix and 'igraph' object.

Note

- 1. Replicates should be at least 5, more than 8 is recommend.
- 2. In case of too many edges/links or not a global network plot, you can stop the process immediately to provent wasting too much time.

```
{
  ### Data preparation ###
  data("Two_group")
  set.seed(999)
```

network_analysis2 39

```
## Analysis
 network_results <- network_analysis(</pre>
    taxobj = Two_group,
    taxlevel = "Genus",
   n = 10,
    threshold = 0.8
 )
 # Nodes information table
 network_nodes <- network_results$Nodes_info</pre>
 head(network_nodes)
 # Adjacency table
 network_adjacency <- network_results$Adjacency_column_table</pre>
 head(network_adjacency)
 # Complete adjacency matrix
 network_matrix <- network_results$Adjacency_matrix</pre>
 print(network_matrix[1:10, 1:10])
 # igraph object
 igraph_object <- network_results$Igraph_object</pre>
 network_stat(igraph_object) # In case you want to see statistics again
 # or do other analysis based on igraph.
}
```

network_analysis2

Conduct Network analysis

Description

A convenient and fast network analysis function, with output results suitable for cytoscape and gephi

Usage

```
network_analysis2(
  input,
  inputtype,
  n,
  threshold,
  method = "spearman",
  display = TRUE,
  input2,
  input2type
)
```

40 network_analysis2

Arguments

input Input dataframe with otu/gene/taxa in row and sample ID in column,at least 5

replicates (more than 8 replicates are recommened).

input type Input dataframe type

1:dataframe with first column of OTUID and last column of taxonomy

2:dataframe with first column of OTUID/taxonomy

3:dataframe of all numeric

n Only keep otu/gene/taxa appearing in n sample size

threshold Threshold of correlation r value

method A character string indicating which correlation coefficient is to be computed.

One of "pearson" or "spearman"

display If display a preview plot of network based on igraph. FALSE for the first attempt

is recommended in case of too many vertices and edges.

input2 A second input data frame with otu/gene/taxa in row and sample ID in column.

Default:NULL

input2type The second input data frame type. Details the same as above. Default:NULL

Details

1. We had optimized the correlation algorithm to achieve a faster running speed. It takes less than 2 minute to calculate dataframe correlation and p value which more than 400 samples and 10000 OTUs for computer with dual Core i5 processor. However, too many vertices(>2000) or links(>10000) may slow the statistical process and visualization,so we recommend that in your first attempt,set display paramter as F to have a preview. Then you can adjust your n/threshold/method paramter to generate a suitable visualization network

2. We display a preview plot so as to adjusting your network. Generally a global figure (like we show in examples) with less than 1000 vertices and 5000 edges/links is recommended. Further more, we recommend you to output the statistics and adjacency table and use software like cytoscape or gephi for better visualization.

Value

One list contains a statistics table of network vertices/nodes and an adjacency table. One preview plot of network in the plot interface and an igraph object(named igraph1) in global environment.

Note

- 1. Replicates should be at least 5, more than 8 is recommend.
- 2. In case of too many edges/links or not a global network plot, you can stop the process immediately to provent wasting too much time.

Author(s)

Wang Ningqi 2434066068@qq.com

network_analysis2 41

```
{
 ### Data preparation ###
 data(testotu)
 rownames(testotu) <- testotu[, 1]</pre>
 inputotu <- testotu[, -c(1, ncol(testotu))]</pre>
 head(inputotu)
 set.seed(999)
 ### One input network analysis ###
 network_result <- network_analysis2(</pre>
    inputotu,
    3,
    10,
    0.9,
    "spearman",
    TRUE
 )
 # Nodes information table
 network_nodes <- network_result$Nodes_info</pre>
 head(network_nodes)
 # Adjacency table
 network_adjacency <- network_result$Adjacency_column_table</pre>
 head(network_adjacency)
 # Complete adjacency matrix
 network_matrix <- network_result$Adjacency_matrix</pre>
 print(network_matrix[1:10, 1:10])
 # igraph object
 igraph_object <- network_result$Igraph_object</pre>
 network_stat(igraph_object) # In case you want to see statistics again
 # or do other analysis based on igraph.
 ### Two inputs network analysis ###
 inputotu1 <- inputotu[1:456, ]</pre>
 inputotu2 <- inputotu[524:975, ]</pre>
 network_result <- network_analysis2(</pre>
    input = inputotu1,
    inputtype = 3,
    input2 = inputotu2,
    input2type = 3,
   n = 10,
   threshold = 0.85,
   method = "spearman",
   display = TRUE
 #### Incorrect demonstration !! ###
     network_result <- network_analysis2(inputotu, 3, 3, 0.8, "spearman", TRUE)</pre>
```

42 network_visual

```
}
# Total edges/links: 10199
# Total vertices: 826
# Too many edges and not a global network
}
```

network_stat

Igraph network statistics

Description

Igraph network statistics

Usage

```
network_stat(input, report = TRUE)
```

Arguments

input An igraph object.

report Logical. If print report to console. Default:TRUE

Value

network statistics

Author(s)

Wang Ningqi 2434066068@qq.com

network_visual

Network Visualization

Description

Visualizes a network based on a network object from network_analysis.

network_visual 43

Usage

```
network_visual(
  network_obj,
  mode = "major_module",
  major_num = 5,
  taxlevel = NULL,
  select_tax = NULL,
  palette = "Set1",
  vertex.size = 6
)
```

Arguments

| network_obj | A network analysis results object generated from ${\tt network_analysis}$. |
|-------------|--|
| mode | The visualization mode, optionally "major_module" or "major_tax". |
| major_num | The number of major modules to display in the network. |
| taxlevel | Taxonomy levels used for visualization when mode is "major_tax". |
| select_tax | A vector of taxa to be selected for displaying in "major_tax" mode. |
| palette | Character. Palette for visualization. |
| vertex.size | Numeric. The size of the vertices. |

Value

A list containing the configured igraph object and the coordinates of the vertices, with network visualization displayed in the plots panel.

```
# Data preparation
data("Two_group")
set.seed(999)
# Analysis
network_results <- network_analysis(</pre>
 taxobj = Two_group,
 taxlevel = "Species",
 n = 10,
  threshold = 0.6
)
# Default mode
network_visual_obj <- network_visual(network_obj = network_results)</pre>
# View again
network_visual_re(network_visual_obj)
# More modules
network_visual_obj <- network_visual(</pre>
```

network_visual_re

```
network_obj = network_results,
   major_num = 10
 # Specific tax
 # Generate top 5 phyla for displaying
 community <- community_plot(</pre>
    taxobj = Two_group,
    taxlevel = "Phylum",
   n = 5,
   palette = "Paired"
 display_phyla <- names(community$filled_color)</pre>
 network_visual_obj <- network_visual(</pre>
   network_obj = network_results,
   mode = "major_tax",
    taxlevel = "Phylum",
    select_tax = display_phyla,
    palette = community$filled_color
 # Another sample for specific tax
 network_visual_obj <- network_visual(</pre>
   network_obj = network_results,
   mode = "major_tax",
    taxlevel = "Phylum",
    select_tax = "p__Proteobacteria"
 )
}
```

Description

Re-visualize network plot from network_visual or network_withdiff

Usage

```
network_visual_re(
  network_visual_obj,
  module_paint = FALSE,
  module_num = NULL,
  module_palette = c("aquamarine3", "antiquewhite2", "goldenrod2"),
  vertex.size = 6,
  vertex.shape = "circle"
)
```

network_withdiff 45

Arguments

network_visual_obj

Network object from network_visual or network_withdiff

module_paint Logical. If network module should be painted. Only work for network object

from network_withdiff.

module_num Numeric indicating which module to be painted.

module_palette Character string with at least two elements. Palette for painting modules.

vertex.size Numeric. The size of the vertices, default:6. Only for network object from

network_visual

vertex.shape Character. The shape of vertices, default: "circle"

Value

NULL but visualization in plot panel.

| network_withdiff Network Analysis with Differential Species | |
|---|--|
|---|--|

Description

Meta network analysis integrating differential taxon into a network analysis

Usage

```
network_withdiff(network_obj, diff_frame, aes_col = NULL, tag_threshold = 5)
```

Arguments

| network_obj | Network analysis results generated from network_analysis |
|-------------|--|
| diff_frame | Differential analysis results generated from indicator_analysis or Deseq_analysis. |

aes_col A named vector of colors to be used to highlight differential taxon vertices

tag_threshold Numeric. A threshold for the minimum number of differential taxon to display.

Value

A list containing the configured igraph object, vertices coordinates, parameters, and tag statistics.

46 network_withdiff

```
# Data preparation
data("Two_group")
set.seed(999)
# Analysis
network_results <- network_analysis(</pre>
  taxobj = Two_group,
  taxlevel = "Genus",
 n = 10,
 threshold = 0.8
indicator_results <- indicator_analysis(</pre>
  taxobj = Two_group,
  taxlevel = "Genus"
deseq_results <- Deseq_analysis(</pre>
  taxobj = Two_group,
  taxlevel = "Genus",
  cutoff = 1,
  control_name = "Control"
# Visualize
network_diff_obj <- network_withdiff(</pre>
 network_obj = network_results,
 diff_frame = indicator_results
# Check contained tags for each model
print(network_diff_obj$tag_statistics$sum_of_tags)
# Check contained different tags for each model
print(network_diff_obj$tag_statistics$detailed_tags)
# Re-visualize
network_visual_re(
  network_visual_obj = network_diff_obj,
  module_paint = TRUE,
 module_num = c(1, 4)
) # Show module with most Treatment indicators
my_module_palette <- color_scheme(</pre>
 c("#83BA9E", "#F49128"),
  5
)
network_visual_re(
 network_visual_obj = network_diff_obj,
  module_paint = TRUE,
  module_num = c(1, 4, 6, 3, 8),
  module_palette = my_module_palette
) # Show module with most Treatment indicators
# Available also for DESeq analysis results
```

network_withdiff 47

```
network_diff_obj <- network_withdiff(</pre>
  network_obj = network_results,
 diff_frame = deseq_results
)
# Parameter adjustment
network_diff_obj <- network_withdiff(</pre>
  network_obj = network_results,
 diff_frame = indicator_results,
  tag_threshold = 20
) # The 'tag_threshold' set too high
network_diff_obj <- network_withdiff(</pre>
  network_obj = network_results,
  diff_frame = indicator_results,
  tag_threshold = 10
) # Set lower
# Check contained tags for each model
print(network_diff_obj$tag_statistics$sum_of_tags)
# Check contained different tags for each model
print(network_diff_obj$tag_statistics$detailed_tags)
network_diff_obj <- network_withdiff(</pre>
  network_obj = network_results,
  diff_frame = indicator_results,
  tag_threshold = 1
) # Set too low
# Another example
data("Three_group")
network_results <- network_analysis(</pre>
  taxobj = Three_group,
  taxlevel = "Genus",
 n = 15,
  threshold = 0.9
)
indicator_results <- indicator_analysis(</pre>
  taxobj = Three_group,
  taxlevel = "Genus"
tag_color <- c(</pre>
  "CF" = "#F8766D",
  "CF_OF" = "#FFFF00",
  "OF" = "\#00BA38",
  "OF_BF" = "#800080",
  "BF" = "\#619CFF",
  "CF_BF" = "#00FFFF"
)
network_diff_obj <- network_withdiff(</pre>
  network_obj = network_results,
  diff_frame = indicator_results,
  aes_col = tag_color,
```

48 object_config

```
tag_threshold = 10
)

# Re-visualize
print(network_diff_obj$tag_statistics$detailed_tags)
network_visual_re(
    network_visual_obj = network_diff_obj,
    module_paint = TRUE,
    module_num = c(8, 10, 11)
) # Show module with most BF indicators
network_visual_re(
    network_visual_obj = network_diff_obj,
    module_paint = TRUE,
    module_num = c(1, 6, 8)
) # Show module with most BF and OF_BF indicators
}
```

object_config

Set taxonomy summary configuration

Description

This function set taxonomy summary configuration by assigning treatment column number, facet column number, replication column number, treatment mapping color, treatment order and facet order.

Usage

```
object_config(
  taxobj,
  treat_location,
  facet_location = NULL,
  rep_location,
  subject_location = NULL,
  treat_col = NULL,
  treat_order = NULL,
  facet_order = NULL
)
```

Arguments

taxobj Taxonomic summary object generated by tax_summary
treat_location Numeric. Treatment column number in metafile/groupinformation.
facet_location Numeric, default:NULL. Facet column number in metafile/groupinformation.
rep_location Numeric. Replication column number in metafile/groupinformation.

object_config 49

subject_location

Numeric, default: NULL. Subject column number in metafile/groupinformation

(used for pairwise experiment).

treat_col Named character string, default:NULL. A set of aesthetic character to map treat-

ment to.

treat_order Character string, default:NULL. The character string indicating treatment dis-

playing order.

facet_order Character string, default:NULL. The character string indicating facet displaying

order.

Value

object_config returns taxonomy summary object with configuration.

Author(s)

Wang Ningqi2434066068@qq.com

```
{
  ### Data preparation ###
  data(testotu)
  groupinformation <- data.frame(</pre>
    group = c(rep("a", 10), rep("b", 10)),
    factor1 = rnorm(10),
    factor2 = rnorm(mean = 100, 10),
    subject = factor(c(1:10, 1:10)),
    group2 = c(rep("e", 5), rep("f", 5), rep("e", 5), rep("f", 5))
  )
  ### Packaging metafile, community data, and taxonomy table ###
  test_object <- tax_summary(</pre>
    groupfile = groupinformation,
    inputtable = testotu[, 2:21],
    reads = TRUE,
    taxonomytable = testotu[, c(1, 22)]
  )
  ### Object configuration ###
  test_object_plan1 <- object_config(</pre>
    taxobj = test_object,
    treat_location = 1,
    rep_location = 4
  ### Facet configuration ###
  test_object_plan2 <- object_config(</pre>
    taxobj = test_object,
    treat_location = 1,
    rep_location = 4,
```

50 structure_plot

```
facet_location = 5
)
}
```

structure_plot

Visualize microbial community composition structure based on tax summary object

Description

Function for visualization of microbial structure with PCAplot, PCoAplot and NMDSplot

Usage

```
structure_plot(
  taxobj,
  taxlevel,
  ptsize = 2,
  diagram = NULL,
  ellipse.level = 0.85,
  facet_row = NULL
)
```

Arguments

| taxobj | Configured tax summary objects. See in object_config. |
|---------------|--|
| taxlevel | taxonomy levels used for visualization. Must be one of c("Domain", "Phylum", "Class", "Order", "Family", "Order", "Family", "Class", "Order", "Family", "Class", "Order", "Family", "Order", "Orde |
| ptsize | Numeric, default: 2. Size of point in plot. See geom_point for details. |
| diagram | Character, default: NULL. A character indicating group diagram, should be in c("ellipse", "stick", "polygon"). |
| ellipse.level | Numeric, default: 0.85. The level at which to draw an ellipse, or, if type = "euclid", the radius of the circle to be drawn. See stat_ellipse for details. |
| facet_row | Numeric, default: NULL. Number of rows when wrap panels. See facet_wrap for details. |

Value

Microbial structure analysis object.

Note

- 1. Do not use NMDS when warning: In metaMDS(t(inputframe)) :stress is (nearly) zero: you may have insufficient data
- 2. Ellipse not available when replicates less than 3, please use 'stick' or 'polygon' instead

structure_plot 51

```
###data preparation####
data("Two_group")
###analvsis####
set.seed(999)
community_structure<- structure_plot(taxobj = Two_group,taxlevel = "Base")</pre>
#check output list in console (not run)
 ######Output list##
 ####Plot#
 ####PCAplot:named as('PCA_Plot')(1/3)
 ####PCoAplot:named as('PCoA_Plot')(2/3)
 ####NMDSplot:named as('NMDS_Plot')(3/3)
 #####Analysis object#
 ####PCA object:named as('PCA_object')
 ####PCoA object:named as('PCoA_object')
 ####NMDS object:named as ('NMDS_object')
 #####Coordinates dataframe#
 ####PCA Coordinates dataframe:named as('PCA_coordinates')
 ####PCoA Coordinates dataframe:named as('PCoA_coordinates')
 ####NMDS Coordinates dataframe:named as('NMDS_coordinates')
 #####Done##
 #check PERMANOVA results
 community_structure$PERMANOVA_statistics
 #extract plot
 community_structure$PCA_Plot
 community_structure$PCoA_Plot
 community_structure$NMDS_Plot
 #extract object
 PCA_obj<- community_structure$PCA_object
 print(PCA_obj)
 #extract coordinates frame
 PCA_coord<- community_structure$PCA_coordinates
 head(PCA_coord)
 #stick plot
 set.seed(999)
community_structure<- structure_plot(taxobj = Two_group,taxlevel = "Base",diagram = "stick")</pre>
 community_structure$PCoA_Plot
 #faced form
 data("Facet_group")
 set.seed(999)
community_structure<- structure_plot(taxobj = Facet_group,taxlevel = "Genus",diagram = "stick")</pre>
 community_structure$PERMANOVA_statistics
 community_structure$PCA_Plot
 community_structure$PCoA_Plot
 community_structure$NMDS_Plot
```

52 sub_tax_summary

sub_tax_summary

Subsetting tax summary objects

Description

Subsetting tax summary objects

Usage

```
sub_tax_summary(taxobj, ..., specificnum = NULL, taxnum = NULL)
```

Arguments

tax summary objects computed by tax_summary.

... logical expression that are are defined in terms of the variables in Groupfile of

tax summary objects. See details in subset.

specific numbers indicating samples to keep based on Groupfile of tax summary

objects.

taxnum specific numbers indicating taxonomy to keep based on Base file

Value

Subset of tax summary objects. Same as tax_summary.

Author(s)

Wang Ningqi 2434066068@qq.com

```
data("Three_group")

# Check meta file
print(Three_group$Groupfile)

# Subsetting tax summary objects

# Select BF and OF groups
sub_testtax_summary <- sub_tax_summary(Three_group, Group %in% c("BF", "OF"))
print(sub_testtax_summary$Groupfile)

# Subsetting according to taxonomy

Proteo <- sub_tax_summary(
    Three_group,
    taxnum = which(Three_group$Base_taxonomy$Phylum == "p__Proteobacteria")
)
print(Proteo$Phylum_percent) # Check phylum table
print(Proteo$Genus_percent) # Check genus table</pre>
```

tax_summary 53

| tax_summary | Encapsulate meta file, feature tables and taxonomy annotation into tax summary object |
|-------------|---|
| | |

Description

The function packages meta file, feature tables and taxonomy annotation into tax summary object

Usage

```
tax_summary(
  groupfile,
  inputtable,
  reads = TRUE,
  taxonomytable,
  into = "standard",
  sep = ";",
  outputtax = c("Phylum", "Genus")
)
```

Arguments

groupfile A data frame containing treatment information

OTU/ASV/species data frame with all numeric. Samples ID should be in column names.

reads Logical.True for reads table and FALSE for percentage table. Default: TRUE taxonomytable Taxonomy annotation data frame, with first column OTU/ASV/TAX number ID and second column taxonomy annotation. See details in example.

into Names of separated taxonomy to create as character vector. Must select from

c("Domain", "Phylum", "Class", "Order", "Family", "Genus", "Species"). Shortcut input: 1) By default "standard": c("Domain" "Phylum" "Class" "Order" "Family" "Genus" "Phylum" "Class" "Order" "Family" "Genus" "Phylum" "Class" "Order" "Family" "Genus" "Family" "Family" "Genus" "Family" "Family" "Genus" "Family" "Fam

input:1)By default. "standard":c("Domain", "Phylum", "Class", "Order", "Family", "Genus", "Species"). Used for standard taxonomy annotation to OTU/ASV table. 2) "complete":c("Domain", "Kingdom", "Phylum", "Class", "Order", "Family", "Genus", "Species").

Used for complete taxonomy annotation to meta genomic table.

sep Separator of taxonomy table.Default: ";".

outputtax Names of output taxonomy level table. Default:c("Phylum", "Genus"). Shortcut

input is available with 'standard' and 'complete' same as above.

Value

One list containing taxonomy table data frame, containing reads and percentage table for each specified output. Full taxonomy annotation data frame is output in global environment.

Note

For taxonomy annotation with 'Kingdom' level, please set 'into' parameter as 'complete'!!!

54 tbRDA_analysis

Author(s)

Wang Ningqi 2434066068@qq.com

```
# Load data
 data(testotu)
 # Create group information data frame
 groupinformation <- data.frame(</pre>
    group = c(rep("a", 10), rep("b", 10)),
    factor1 = rnorm(10),
   factor2 = rnorm(mean = 100, 10),
    subject = factor(c(1:10, 1:10))
 )
 # Packaging data into a taxonomy summary object
 test_object <- tax_summary(</pre>
    groupfile = groupinformation,
    inputtable = testotu[, 2:21],
    reads = TRUE,
    taxonomytable = testotu[, c(1, 22)]
 )
 # Check integrated object
 print(test_object)
 # Extract genus relative abundance table
 test_Genus <- test_object$Genus_percent</pre>
 head(test_Genus)
 # Check corresponding taxonomy information of genus table
 test_Genus_tax <- test_object$Genus_taxonomy</pre>
 head(test_Genus_tax)
 # Summary base table into all taxonomy levels with standard output
 test_object <- tax_summary(</pre>
    groupfile = groupinformation,
    inputtable = testotu[, 2:21],
    reads = TRUE,
    taxonomytable = testotu[, c(1, 22)],
    outputtax = "standard"
 head(test_object$Species_percent) # View first 10 rows of species percentage
 head(test_object$Genus) # View first 10 rows of genus table
}
```

tbRDA_analysis 55

Description

RDA analysis including co-linearity diagnostics and necessary statistics.

Usage

```
tbRDA_analysis(otudata, envdata, collinearity, perm.test = TRUE)
```

Arguments

| otudata | Feature table of all numeric variable, with annotation in row names |
|--------------|--|
| envdata | Environmental factor of all numeric variable, with sample-ID in row names and environmental factor in column names |
| collinearity | If done collinearity diagnostics. Default, TRUE. |
| perm.test | Logical. If conduct permutation test. Default:TRUE. |

Value

Three permutation test result print ,one preview plot ,a RDA object(default name:otu.tab.1) and a summary of RDA object

Note

1. When Axis length in first axis more than 4, you should choose CCA instead of RDA.

Author(s)

```
Wang Ningqi 2434066068@qq.com
```

```
### Data preparation ###
library(vegan)
data(varechem)
head(varechem)
data(testotu)
require(tidyr); require(magrittr) ## Or use pipe command in "dplyr"
sep_testotu <- Filter_function(</pre>
 input = testotu,
  threshold = 0.0001,
 format = 1
) %>%
separate(
  ., col = taxonomy,
  into = c("Domain", "Phylum", "Order", "Family", "Class", "Genus", "Species"),
  sep = ";"
top10phylum <- aggregate(</pre>
 sep_testotu[, 2:21],
```

56 tbRDA_analysis

```
by = list(sep_testotu$Phylum),
  FUN = sum
) %>%
Top_taxa(
  input = .,
  n = 10,
  inputformat = 2,
  outformat = 1
)
rownames(top10phylum) <- top10phylum[, 1]</pre>
top10phylum <- top10phylum[, -1]</pre>
group <- data.frame(</pre>
  group = c(rep("a", 10), rep("b", 10)),
  factor1 = rnorm(10),
  factor2 = rnorm(mean = 100, 10)
### RDA analysis ###
set.seed(999)
RDAresult <- tbRDA_analysis(
  top10phylum,
  varechem[1:20, ],
  TRUE
)
# Environmental statistics
print(RDAresult$factor_statistics)
# Visualization using ggplot
rda_object <- RDAresult$rda_object</pre>
rda_summary <- RDAresult$rdasummary</pre>
rda_env <- as.data.frame(rda_summary$biplot)</pre>
rda_sample <- as.data.frame(rda_summary$sites)</pre>
rda_otu <- as.data.frame(rda_summary$species)</pre>
xlab <- paste0("RDA1:", round(RDAresult$rdasummary$concont$importance[2, 1], 4) * 100, "%")
ylab <- paste0("RDA2:", round(RDAresult$rdasummary$concont$importance[2, 2], 4) * 100, "%")
library(ggrepel)
# Create a sample RDA plot
RDAplot <- ggplot(data = rda_sample, aes(RDA1, RDA2)) +</pre>
  geom_point(aes(color = group$group), size = 2) +
  geom_point(data = rda_otu, pch = "+", color = "orange", size = 4) +
  geom_hline(yintercept = 0) +
  geom_vline(xintercept = 0) +
  geom_segment(data = rda_env, aes(x = 0, y = 0, xend = RDA1 \star 0.8, yend = RDA2 \star 0.8),
                arrow = arrow(angle = 22.5, length = unit(0.35, "cm")),
                linetype = 1, size = 0.6, colour = "red") +
  geom_text_repel(color = "red", data = rda_env,
                   aes(RDA1, RDA2, label = row.names(rda_env))) +
  labs(x = xlab, y = ylab, color = "Treatment",
        title = paste0("p = ", anova.cca(rda_object)["Model", "Pr(>F)"])) +
```

testotu 57

testotu

test otudata

Description

A dataset containing 20 samples and 1000 OTUs from soil to test,taxonomy information is covered randomly(not actual)

Usage

testotu

Format

A data frame with 1000 rows and 22 variables.

theme_zg

A classic theme for ggplot

Description

A classic theme for ggplot

Usage

```
theme_zg()
```

Value

ggplot theme

Note

Build inside the LorMe package, Please use theme_zg() as a theme directly

Top_taxa

| Three_group | Tax summary object with three groups | |
|-------------|--------------------------------------|--|
|-------------|--------------------------------------|--|

Description

Enraptured summary object with three groups. Configuration has been assigned.

Usage

Three_group

Format

Tax summary object with configuration

| Top_taxa | Calculate top taxa and others | |
|----------|-------------------------------|--|
| | | |

Description

Top taxa is widely used in data analysis,here we provide a simple function to calculate which simplify your R script.

Usage

```
Top_taxa(input, n, inputformat, outformat)
```

Arguments

| input | Reads or relative abundance(recommended) of OTU/Taxa/gene data frame,see details in inputformat |
|-------------|---|
| n | Top n taxa remained according to relative abundance |
| inputformat | 1:data frame with first column of OTUID and last column of taxonomy 2:data frame with first column of OTUID/taxonomy (recommended!!!) 3:data frame of all numeric, with row names of OTUID/taxonomy |
| outformat | return outformat the same as inputformat return data frame of all numeric with OTU/gene/taxa ID in row names(not available for inputformat 1). |

Value

Data frame with top n taxa

Top_taxa 59

Author(s)

Wang Ningqi2434066068@qq.com

```
### Data preparation ####
data(testotu)
require(tidyr); require(magrittr) ## Or use pipe command in "dplyr"
testotu.pct <- data.frame(</pre>
  OTU.ID = testotu[, 1],
  sweep(testotu[, -c(1, 22)], 2, colSums(testotu[, -c(1, 22)]), "/"),
  taxonomy = testotu[, 22]
)
sep_testotu <- Filter_function(</pre>
  input = testotu,
  threshold = 0.0001,
  format = 1
) %>%
  separate(
    ., col = taxonomy,
    into = c("Domain", "Phylum", "Order", "Family", "Class", "Genus", "Species"),
    sep = ";"
  )
phylum <- aggregate(</pre>
  sep_testotu[, 2:21], by = list(sep_testotu$Phylum), FUN = sum
)
phylum1 <- data.frame(row.names = phylum[, 1], phylum[, -1])</pre>
##### Input format 1, top 100 OTU #####
top100otu <- Top_taxa(</pre>
  input = testotu.pct,
 n = 100,
  inputformat = 1,
  outformat = 1
)
##### Input format 2, top 15 phylum #####
head(phylum)
top15phylum <- Top_taxa(</pre>
  input = phylum,
  n = 15,
  inputformat = 2,
  outformat = 1
)
##### Input format 3, top 15 phylum #####
head(phylum1)
top15phylum <- Top_taxa(</pre>
```

60 t_test_report

```
input = phylum1,
  n = 15,
  inputformat = 3,
  outformat = 1
)
```

Two_group

Tax summary object with two groups

Description

Enraptured summary object with two groups. Configuration has been assigned.

Usage

Two_group

Format

Tax summary object with configuration

t_test_report

Print Student's t-Test report

Description

Print Student's t-Test report

Usage

```
t_test_report(
  data,
  treatment_col,
  value_col,
  paired,
  subject_col,
  report = TRUE
)
```

Arguments

data Data frame containing the treatment, value and other information.

treatment_col Numeric indicating where treatment locates (column number) in data.

value_col Numeric indicating where treatment value (column number) in data.

paired Logical indicating whether you want a paired t-test.

subject_col Only meaningful when Pair is ture. Numeric indicating where subject of treat-

ment (column number) in data.

report Logical. If print report to console. Default:TRUE

volcano_plot 61

Value

t_test_report returns list containing:

- 1. data frame of basic data descrption
- 2. results of student's t-Test

Examples

```
{
  ### Data preparation ###
  testdata <- data.frame(</pre>
    treatment = c(rep("A", 6), rep("B", 6)),
    subject = rep(c(1:6), 2),
    value = c(rnorm(6, 2), rnorm(6, 1))
  )
  # Perform t-test (unpaired)
  t_test_result <- t_test_report(</pre>
    data = testdata,
    treatment_col = 1,
    value\_col = 3
  )
  # Perform paired t-test
  t_test_result <- t_test_report(</pre>
    data = testdata,
    treatment_col = 1,
    value\_col = 3,
    paired = TRUE,
    subject\_col = 2
  )
  ### Basic data description ###
  print(t_test_result[[1]])
  print(t_test_result$basicdata)
  ### T-test results ###
  print(t_test_result[[2]])
  print(t_test_result$t.test_results)
}
```

volcano_plot

Generate Volcano plot base on Deseq_analysis or indicator_analysis results

Description

Generate Volcano plot base on Deseq_analysis or indicator_analysis results

62 volcano_plot

Usage

```
volcano_plot(inputframe, cutoff = NULL, aes_col = c("#FE5C5C", "#75ABDE"))
```

Arguments

inputframe A data frame containing the results based on Deseq_analysis or indicator_analysis

(only two group indicators)

cutoff A numeric value specifying the fold change cutoff, should be the same as in

Deseq_analysis

aes_col A named vector of colors to be used in the plots

Value

A list of two ggplot objects, one for the fold change versus adjusted p-value plot and another for the mean abundance versus fold change or enrichment factor plot.

Author(s)

Wang Ningqi 2434066068@qq.com

```
###data prepration###
 # Load data
 data("Two_group")
 # Define color based on treatment column
 mycolor <- Two_group$configuration$treat_col</pre>
 ### DESeq analysis ###
 deseq_results <- Deseq_analysis(</pre>
    taxobj = Two_group,
    taxlevel = "Genus",
   cutoff = 1,
    control_name = "Control"
 )
 ### Or indicator analysis ###
 indicator_results <- indicator_analysis(</pre>
    taxobj = Two_group,
    taxlevel = "Genus"
 )
 # Create volcano plot for DESeq results
 volcano_plot <- volcano_plot(</pre>
    inputframe = deseq_results,
   cutoff = 1,
   aes_col = mycolor
 )
```

wilcox_test_report 63

```
print(volcano_plot$FC_FDR) # Fold Change and FDR values
print(volcano_plot$Mean_FC) # Mean Fold Change values

# Create volcano plot for indicator results
volcano_plot <- volcano_plot(
   inputframe = indicator_results,
   cutoff = 1,
   aes_col = mycolor
)
print(volcano_plot$FC_FDR) # Fold Change and FDR values
print(volcano_plot$Mean_FC) # Mean Fold Change values
}</pre>
```

wilcox_test_report

Print Wilcoxon Rank Sum and Signed Rank Tests reprot

Description

Print Wilcoxon Rank Sum and Signed Rank Tests reprot

Usage

```
wilcox_test_report(
  data,
  treatment_col,
  value_col,
  paired = FALSE,
  subject_col = NULL,
  report = TRUE
)
```

Arguments

Data frame containing the treatment, value and other information.

treatment_col Numeric indicating where treatment locates (column number) in data.

value_col Numeric indicating where treatment value (column number) in data.

Logical indicating whether you want a paired test.

subject_col Only meaningful when Pair is ture. Numeric indicating where subject of treatment (column number) in data.

report Logical. If print report to console. Default:TRUE

Value

wilcox_test_report returns data frame of basic data description.

64 wilcox_test_report

```
# Data preparation
 testdata <- data.frame(</pre>
   treatment = c(rep("A", 6), rep("B", 6)),
   subject = rep(1:6, 2),
   value = c(rnorm(6, 2), rnorm(6, 1))
 # Wilcoxon test (unpaired)
 wilcox_result <- wilcox_test_report(</pre>
   data = testdata,
   treatment_col = 1,
   value\_col = 3
 )
 # Wilcoxon signed rank test (paired)
 wilcox_result <- wilcox_test_report(</pre>
   data = testdata,
   treatment_col = 1,
   value_col = 3,
   paired = TRUE,
   subject\_col = 2
 ### Basic data description ###
 print(wilcox_result)
}
```

Index

| * datasets Facet_group, 24 testotu, 57 Three_group, 58 | Module_abundance, 30, 32 Module_composition, 30, 33 multipatt, 26, 27 |
|--|---|
| Two_group, 60 | nc, <i>30</i> , 35 NC_remove, <i>30</i> , 36 |
| Alpha_diversity_calculator, 3, 29 Alpha_diversity_calculator2, 4 anova_report, 6, 9 as_tibble, 18, 20 | network_analysis, 29, 32, 33, 35, 36, 37, 42, 43, 45 network_analysis2, 39 network_stat, 42 |
| auto_signif_test, 8, 29 | network_visual, 29, 30, 42, 44, 45 network_visual_re, 30, 44 |
| circulation_lm, 10, 30 color_scheme, 11, 29, 30, 33 | network_withdiff, 29, 30, 44, 45, 45 |
| combine_and_translate, 12, 29 community_plot, 14, 29 | object_config, 3, 14, 17, 21, 26, 48, 50 |
| compare_plot, 16, 29 | set.seed, 36 stat_ellipse, 50 |
| DataFrame, 18, 20 | structure_plot, 29, 50 |
| DESeq, 18, 20 | sub_tax_summary, 29, 52 |
| Deseq_analysis, 17, 29, 30, 45, 62 Deseq_analysis2, 19 | subset, 52 |
| DESeqDataSetFromMatrix, 18, 20 | t_test_report, 9, 60 |
| differential_bar, 21, 29 | tax_summary, 4, 29, 38, 48, 52, 53 |
| Dimension_reduction, 23, 29 | tbRDA_analysis, 30, 54 testotu, 57 |
| Facet_group, 24 | theme_zg, 29, 57 |
| $facet_wrap, 50$ | Three_group, 58 |
| Filter_function, 25, 29 | Top_taxa, 29, 58 |
| <pre>geom_point, 50</pre> | Two_group, 60 |
| how, 27 | volcano_plot, 29, 61 |
| indicator_analysis, 26, 29, 30, 45, 62 | wilcox_test_report, 9, 63 |
| kruskal_report, 9, 27 | |
| lm, <i>11</i> LorMe, 29 | |
| manhattan, 29, 30 | |