

Package ‘pctax’

February 9, 2026

Type Package

Title Professional Comprehensive Omics Data Analysis

Version 0.1.7

Description Provides a comprehensive suite of tools for analyzing omics data. It includes functionalities for alpha diversity analysis, beta diversity analysis, differential abundance analysis, community assembly analysis, visualization of phylogenetic tree, and functional enrichment analysis. With a progressive approach, the package offers a range of analysis methods to explore and understand the complex communities. It is designed to support researchers and practitioners in conducting in-depth and professional omics data analysis.

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Encoding UTF-8

RoxygenNote 7.3.2

Depends R (>= 4.2.0)

LazyData true

Imports pcutils (>= 0.2.5), dplyr, readr, ggplot2 (>= 3.2.0), vegan, magrittr, grDevices, RColorBrewer, ggrepel, reshape2, stats, tibble, utils, ggpubr, patchwork, ggnewscale, ade4, scales

Suggests picante, httr, NST, permute, aplot, ggfun, pheatmap, MASS, Rtsne, mixOmics, geosphere, phyloseq, phyloseqGraphTest, plotly, umap, Hmisc, minpack.lm, bbmle, snow, foreach, doSNOW, tidytree, ggtree, ggtreeExtra, vctrs, zoo, ape, DESeq2, limma, ALDEx2, Mfuzz, edgeR, methods, randomForest, knitr, rmarkdown, MetaNet, showtext, jsonlite, prettydoc, readxl, stringr, ggExtra, clipr, zetadiv, ggforce, gggenes, mediation

VignetteBuilder knitr

BugReports <https://github.com/Asa12138/pctax/issues>

URL <https://github.com/Asa12138/pctax>

ByteCompile true

biocViews Microbiome, Software, Visualization

NeedsCompilation no

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Repository CRAN

Date/Publication 2026-02-09 08:20:08 UTC

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add_strip	<i>add strips for a tree plot</i>
-----------	-----------------------------------

Description

add strips for a tree plot

Usage

```
add_strip(trp, some_tax, flat_n = 5, strip_params = NULL)
```

Arguments

- trp tree plot from ggtree
- some_tax some tax you want to add strip
- flat_n flat the text when taxa number more than flat_n.
- strip_params parameters parse to [geom_strip](#)

Value

tree plot

Examples

```
data(otutab, package = "pcutils")
# run yourself
if (interactive()) {
  ann_tree(taxonomy, otutab) -> tree
  easy_tree(tree) -> p
  some_tax <- table(taxonomy$Phylum) %>%
    sort(decreasing = TRUE) %>%
    head(5) %>%
    names()
  add_strip(p, some_tax)
}
```

add_tax	<i>Add taxonomy for a pc_otu object</i>
---------	---

Description

Add taxonomy for a pc_otu object

Usage

```
add_tax(pc, taxonomy)
```

Arguments

pc	a pc_otu object
taxonomy	a taxonomy data.frame, look out the rownames of taxonomy and otutab should matched!

Value

pc_otu

Examples

```
data(otutab, package = "pcutils")
pc_tax1 <- pc_otu(otutab, metadata)
pc_tax1 <- add_tax(pc_tax1, taxonomy)
```

ALDEX	<i>ALDEX</i>
-------	--------------

Description

ALDEX

Usage

```
ALDEX(otutab, group_df)
```

Arguments

otutab	otutab
group_df	a dataframe with rowname same to dist and one group column

Value

diff

References

<https://cloud.tencent.com/developer/article/1621879>

all_ec_info	<i>all element cycle information.</i>
-------------	---------------------------------------

Description

all element cycle information.

Format

a list contains four tables.

ec_node chemicals

ec_link reactions

ec_gene genes

ec_path reactions labels

all_sp_la_zh_name	<i>all species latin names and chinese names</i>
-------------------	--

Description

all species latin names and chinese names.

Format

a dataframe.

latin_name latin name

chinese_name chinese name

ann_tree	<i>Annotate a tree</i>
----------	------------------------

Description

Annotate a tree

Easy way to plot a phylogenetic tree

Usage

```
ann_tree(f_tax, otutab = NULL, level = ncol(f_tax), ignore_pattern = NULL)
```

```
easy_tree(
  tree,
  highlight = "Phylum",
  colorfill = "color",
  topN = NULL,
  pal = NULL,
  name_prefix = FALSE,
  basic_params = NULL,
  add_abundance = TRUE,
  color_name = "abundance",
  add_tiplab = TRUE,
  fontsize = NULL
)
```

Arguments

f_tax	taxonomy dataframe
otutab	otutab, rowname==rowname(taxonomy)
level	1~7
ignore_pattern	An optional regular expression pattern to match tip or node labels for dropping.
tree	result from ann_tree
highlight	highlight which level, one of tree\$level
colorfill	"color" or "fill"
topN	topN to show
pal	color pal
name_prefix	keep the prefix like "k__" or "p__" in the label? Default: FALSE
basic_params	parameters parse to ggtree
add_abundance	logical
color_name	color name
add_tiplab	logical
fontsize	tip label fontsize

Value

a treedata
a ggplot

Examples

```
if (interactive()) {  
  data(otutab, package = "pcutils")  
  ann_tree(taxonomy, otutab) -> tree  
  # run yourself  
  easy_tree(tree, add_abundance = FALSE) -> p  
  p  
}
```

aor	<i>Calculate Abundance-occupancy_relationship</i>
-----	---

Description

Calculate Abundance-occupancy_relationship
Plot a AOR

Usage

```
aor(otutab, ...)  
  
## S3 method for class 'data.frame'  
aor(  
  otutab,  
  top_r = 0.7,  
  ocup_n = ceiling(0.8 * ncol(otutab)),  
  special_n = ceiling(0.1 * ncol(otutab)),  
  ...  
)  
  
## S3 method for class 'AOR'  
plot(x, ...)
```

Arguments

otutab	otutab
...	add
top_r	percentage of top relative abundance
ocup_n	percentage of top occupied
special_n	how many occupancy define as specialists
x	AOR object

Value

AOR
ggplot

References

Barberán, A., Bates, S. T., Casamayor, E. & Fierer, N. (2012) Using network analysis to explore co-occurrence patterns in soil microbial communities.

Examples

```
data(otutab, package = "pcutils")
aor(otutab) -> AOR
plot(AOR)
```

as.b_dist	<i>Transfer dist to b_dist</i>
-----------	--------------------------------

Description

Transfer dist to b_dist
Plot dist
Plot b_dist

Usage

```
as.b_dist(dist, group_df = NULL)

## S3 method for class 'dist'
plot(x, group_df = NULL, ...)

## S3 method for class 'b_dist'
plot(x, mode = 1, c_group = "inter", ...)
```

Arguments

dist	a dist object
group_df	a dataframe with rowname same to dist and one group column
x	a b_dist
...	additional
mode	1~3
c_group	"inter" or "intra" or both to plot

Value

- a b_dist with annotation by group
- a pheatmap
- a ggplot or pheatmap

Examples

```
data(otutab, package = "pcutils")
mat_dist(otutab) %>% as.b_dist(., group_df = metadata["Group"]) -> aa
plot(aa)
plot(aa, mode = 2)
```

as.dist.b_dist	<i>Transfer b_dist to dist</i>
----------------	--------------------------------

Description

Transfer b_dist to dist

Usage

```
## S3 method for class 'b_dist'
as.dist(m, diag = FALSE, upper = FALSE)
```

Arguments

- | | |
|-------|---|
| m | a b_dist object |
| diag | logical value indicating whether the diagonal of the distance matrix should be printed by print.dist. |
| upper | logical value indicating whether the upper triangle of the distance matrix should be printed by print.dist. |

Value

dist

a_diversity	<i>Calculate a_diversity of otutab</i>
-------------	--

Description

Calculate a_diversity of otutab

Usage

```
a_diversity(otutab, ...)

## S3 method for class 'data.frame'
a_diversity(
  otutab,
  method = c("richness", "shannon"),
  tree = NULL,
  digits = 4,
  ...
)

## S3 method for class 'pc_otu'
a_diversity(otutab, method = "all", tbl = "otutab", ...)

## S3 method for class 'numeric'
a_diversity(otutab, ...)
```

Arguments

otutab	numeric
...	pass to a_diversity.data.frame
method	one of "all", "richness", "chao1", "ace", "gc", "shannon", "simpson", "pd", "pielou"
tree	a iphylo object match the rownames of otutab
digits	maintance how many digits
tbl	which table

Value

a a_res object

Examples

```
data(otutab, package = "pcutils")
a_diversity(otutab) -> a_res
plot(a_res, "Group", metadata)
```

batch_mediate	<i>Batch mediate</i>
---------------	----------------------

Description

Batch mediate

Usage

```
batch_mediate(data, mediator_df, nsims = 500, conf.level = 0.95)
```

Arguments

data	data.frame with two columns: X (independent variable) and Y (dependent variable).
mediator_df	data.frame with mediators, each column representing a different mediator variable.
nsims	Number of bootstrap simulations for estimating confidence intervals (default is 500).
conf.level	Confidence level for the confidence intervals (default is 0.95).

Value

data.frame

Examples

```
set.seed(123)
n <- 200
X <- rnorm(n)
M1 <- 0.5 * X + rnorm(n)
M2 <- 0.3 * X + rnorm(n)
M3 <- 0.1 * X + rnorm(n)
Y <- 0.3 * X + 0.4 * M1 + 0.2 * M2 + rnorm(n)
data <- data.frame(X, Y)
mediators <- data.frame(M1, M2, M3)
if (requireNamespace("mediation")) {
  results <- batch_mediate(data, mediators, nsims = 99)
  print(results)
}
```

bbtt	<i>ggdotchart for diff analysis</i>
------	-------------------------------------

Description

ggdotchart for diff analysis

Usage

```
bbtt(res, pvalue = "glm.eBH", topN = 20)
```

Arguments

res	result of ALDEX or kwtest
pvalue	the name of pvaule
topN	topN

Value

ggplot

before_tree	<i>Before df2tree check</i>
-------------	-----------------------------

Description

Before df2tree check

Usage

```
before_tree(f_tax)
```

Arguments

f_tax	table
-------	-------

Value

table

Examples

```
wrong_taxdf <- data.frame(
  kingdom = c(rep(c("A", "B"), each = 4), "C", NA),
  "phylum" = c("A", "a", "b", "c", "c", "c", "d", NA, NA, "e")
)
before_tree(wrong_taxdf)
```

b_analyse

*Beta_diversity Ordination: dimensionality reduction***Description**

Species abundance data can be preprocessed with Hellinger transformation or chord transformation data before PCA analysis. Because the Hellinger distance or chord distance with-without data is equal to $\sqrt{2}\sqrt{1 - \text{Ochiai similarity}}$, therefore, the sorting diagram (type 1 scale) of PCA analysis after Hellinger transformation or chord transformation with-without data is internal sample The distance between the squares is the Ochiai distance. $\sqrt{2}\sqrt{1 - \text{Ochiai similarity}}$ is a distance measure, which is also suitable for the analysis of species data. The processed data is then used for pca without norm.

Usage

```
b_analyse(otutab, ...)

## S3 method for class 'data.frame'
b_analyse(
  otutab,
  norm = TRUE,
  method = c("pca"),
  group = NULL,
  dist = "bray",
  ndim = 2,
  scale = FALSE,
  ...
)
```

Arguments

otutab	an otutab data.frame, samples are columns, taxa are rows.
...	add
norm	should normalized or not? (hellinger)
method	one of "pca", "pcoa", "ca", "dca", "nmms", "plsda", "tsne", "umap", "lda", "all"
group	if needed, give a group vector
dist	if use pcoa or nmms, your can choose a dist method (default: bray) or input a distance matrix.
ndim	how many dimension be kept? (default:2). 3 for b_res_3d()
scale	scale, default: FALSE

Value

b_res object

References

<https://www.jianshu.com/p/9694c0b6302d> <https://zhuanlan.zhihu.com/p/25501130>

Examples

```
data(otutab, package = "pcutils")
b_analyse(otutab, method = "pca") -> b_res
plot(b_res, "Group", metadata)
```

b_NTI1

Calculate beta_NTI

Description

Calculate beta_NTI

Usage

```
b_NTI1(
  phylo,
  otutab,
  beta.reps = 9,
  weighted = TRUE,
  threads = 1,
  verbose = TRUE
)
```

Arguments

phylo	a phylo object
otutab	otutab
beta.reps	how many simulation performed?
weighted	logical
threads	use how many threads to calculate (default:4)
verbose	verbose

Value

a dist: b_NTI

b_res_3d	<i>3D plot for b_res</i>
----------	--------------------------

Description

3D plot for b_res

Usage

```
b_res_3d(b_res, Group, metadata = NULL, ...)
```

Arguments

b_res	a b_res object
Group	group vector for color
metadata	metadata contain Group
...	add

Value

plotly list

Examples

```
if (requireNamespace("plotly")) {  
  data(otutab, package = "pcutils")  
  b_analyse(otutab, method = "pca", ndim = 3) -> b_res  
  b_res_3d(b_res, "Group", metadata)  
}
```

check_taxonkit	<i>Check taxonkit</i>
----------------	-----------------------

Description

Check taxonkit

Usage

```
check_taxonkit(print = TRUE)
```

Arguments

print	print
-------	-------

Value

taxonkit path

See Also

Other Rtaxonkit: [download_taxonkit_dataset\(\)](#), [install_taxonkit\(\)](#), [name_or_id2df\(\)](#), [taxonkit_filter\(\)](#), [taxonkit_lca\(\)](#), [taxonkit_lineage\(\)](#), [taxonkit_list\(\)](#), [taxonkit_name2taxid\(\)](#), [taxonkit_reformat\(\)](#)

convert_taxon_name	<i>Convert taxon names between Chinese and Latin</i>
--------------------	--

Description

Convert taxon names between Chinese and Latin

Usage

```
convert_taxon_name(input_names, mode = "latin_to_chinese", fuzzy = FALSE)
```

Arguments

input_names	input names
mode	conversion mode, "latin_to_chinese" or "chinese_to_latin"
fuzzy	whether to use fuzzy matching, default is FALSE

Value

character vector of converted names

Examples

```
convert_taxon_name(c("Escherichia coli", "Clostridioides difficile"))
```

cor_net	<i>Correlation network, species-interaction network for omics</i>
---------	---

Description

Correlation network, species-interaction network for omics

Usage

```
cor_net()
```

Value

No value

df2tree	<i>From a dataframe to construct a phylo</i>
---------	--

Description

NOTE: this function will do before_tree first.

Usage

```
df2tree(data, edge_df = FALSE, ignore_pattern = NULL)
```

Arguments

data dataframe
 edge_df if the data is edge_df ?
 ignore_pattern An optional regular expression pattern to match tip or node labels for dropping.

Value

phylo object

Examples

```
data(otutab, package = "pcutils")
df2tree(taxonomy) -> tax_tree
print(tax_tree)
# check all nodes matched!
if (requireNamespace("picante")) {
  picante::match.phylo.comm(tax_tree, t(otutab)) -> nn
  nrow(nn$comm) == nrow(t(otutab))
}
```

df2tree1	<i>From a dataframe to construct a phylo (save nwk)</i>
----------	---

Description

NOTE: this function will transfer all space to _

Usage

```
df2tree1(taxa)
```

Arguments

taxa dataframe

Value

phylo object

Examples

```
if (requireNamespace("ape")) {
  data(otutab, package = "pcutils")
  df2tree1(taxonomy) -> tax_tree
  print(tax_tree)
}
```

diff_da	<i>Difference analysis</i>
---------	----------------------------

Description

Difference analysis

Usage

```
diff_da(
  otutab,
  group_df,
  ctrl = NULL,
  method = "deseq2",
  log = TRUE,
  add_mini = NULL,
  ...
)
```

Arguments

otutab	otutab
group_df	a dataframe with rowname same to dist and one group column
ctrl	the control group, one level of groups
method	one of "deseq2", "edger", "limma", "t.test", "wilcox.test"
log	do log transfer for limma?
add_mini	add_mini when calculate the logFC. e.g (10+0.1)/(0+0.1), default 0.5*min(abundance)
...	other parameters

Value

a dataframe

Examples

```
if (requireNamespace("limma")) {
  data(otutab, package = "pcutils")
  diff_da(otutab, metadata["Group"], method = "limma") -> res
  volcano_p(res)
  volcano_p(res, mode = 2)
}
```

download_taxonkit_dataset

Download taxonkit dataset

Description

Download taxonkit dataset

Usage

```
download_taxonkit_dataset(make_sure = FALSE, taxdump_tar_gz = NULL)
```

Arguments

make_sure make sure to do this

taxdump_tar_gz your download taxdump_tar_gz file from <https://ftp.ncbi.nih.gov/pub/taxonomy/taxdump.tar.gz>

Value

No value

See Also

Other Rtaxonkit: [check_taxonkit\(\)](#), [install_taxonkit\(\)](#), [name_or_id2df\(\)](#), [taxonkit_filter\(\)](#), [taxonkit_lca\(\)](#), [taxonkit_lineage\(\)](#), [taxonkit_list\(\)](#), [taxonkit_name2taxid\(\)](#), [taxonkit_reformat\(\)](#)

drop_tips_update

Drop Tips and Update a Phylogenetic Tree

Description

This function iteratively removes specified tips (or tips matching a pattern) from a phylogenetic tree without collapsing internal nodes or singleton nodes.

Usage

```
drop_tips_update(tr, drop_name, pattern = NULL)
```

Arguments

<code>tr</code>	A phylogenetic tree of class <code>phylo</code> .
<code>drop_name</code>	A character vector of tip or node names to drop. If missing and <code>pattern</code> is provided, names matching the pattern will be dropped.
<code>pattern</code>	An optional regular expression pattern to match tip or node labels for dropping.

Value

A `phylo` object with specified tips removed.

Examples

```
if (requireNamespace("ape")) {
  library(ape)
  tr <- rtree(10)
  plot(tr)
  # Drop tips containing "t1" or "t2" in their label
  tr2 <- drop_tips_update(tr, pattern = "t1|t2")
  plot(tr2)

  # Alternatively, specify tips directly
  tr3 <- drop_tips_update(tr, drop_name = c("t3", "t5"))
  plot(tr3)
}
```

envfitt

Envfit test for RDA result

Description

Envfit test for RDA result

Usage

```
envfitt(phy.rda, env, ...)
```

Arguments

<code>phy.rda</code>	a rda result
<code>env</code>	environmental factors
<code>...</code>	add

Value

`g_test` object

See Also[envfit](#)**Examples**

```
data(otutab, package = "pcutils")
env <- metadata[, 6:10]
# RDA
myRDA(otutab, env) -> phy.rda
envfitt(phy.rda, env) -> envfit_res
plot(envfit_res)
```

geo_sim

*Lm for sample similarity and geographical distance***Description**

Lm for sample similarity and geographical distance

Usage

```
geo_sim(otutab, geo, method = "bray", spe_nwk = NULL, ...)
```

Arguments

otutab	an otutab data.frame, samples are columns, taxa are rows.
geo	a two-columns dataframe, first is latitude, second is longitude
method	Dissimilarity index, partial match to "bray", "euclidean"...see vegdist ; unifrac
spe_nwk	a phylo tree if use unifrac...
...	additional

Value

a ggplot

References

Graco-Roza, C. et al. (2022) Distance decay 2.0 - A global synthesis of taxonomic and functional turnover in ecological communities. *Glob Ecol Biogeogr* 31, 1399–1421.

Examples

```
if (requireNamespace("geosphere")) {
  library(ggplot2)
  data(otutab, package = "pcutils")
  metadata[, c("lat", "long")] -> geo
  geo_sim(otutab, geo) -> geo_res
}
```

get_all_sp_la_zh_name *get all species Latin and Chinese name from the CCTCC database*

Description

get all species Latin and Chinese name from the CCTCC database

Usage

```
get_all_sp_la_zh_name(
  download_dir = "~/Documents/",
  each_verbose = FALSE,
  max_requests = 50,
  max_id = 30609,
  failure_ids = NULL
)
```

Arguments

download_dir	default
each_verbose	each_verbose
max_requests	default 50
max_id	default 30609, try to make sure on the website
failure_ids	failure_ids

Value

No value

get_diff_type *Get mean and type*

Description

Get mean and type

Usage

```
get_diff_type(otutab, group_df)
```

Arguments

otutab	otutab
group_df	a dataframe with rowname same to dist and one group column

Value

No value

gp_dis_density	<i>Group inter-intra density</i>
----------------	----------------------------------

Description

Group inter-intra density

Usage

```
gp_dis_density(otutab, group)
```

Arguments

otutab	an otutab data.frame, samples are columns, taxa are rows.
group	group vector

Value

ggplot

Examples

```
data(otutab, package = "pcutils")
gp_dis_density(otutab, metadata["Group"])
```

grap_p_test	<i>Performs graph-based permutation tests</i>
-------------	---

Description

Performs graph-based permutation tests

Usage

```
grap_p_test(otutab, metadata, group = "Group", nperm = 999, ...)
```

Arguments

otutab	an otutab data.frame, samples are columns, taxa are rows.
metadata	metadata
group	one group name in columns of metadata
nperm	numbers of permutations to perform
...	additional

Value

ggplot

Examples

```
if (requireNamespace("phyloseqGraphTest") && requireNamespace("phyloseq")) {  
  data(otutab, package = "pcutils")  
  grap_p_test(otutab, metadata, "Group")  
}
```

install_taxonkit	<i>Install taxonkit</i>
------------------	-------------------------

Description

Install taxonkit

Usage

```
install_taxonkit(make_sure = FALSE, taxonkit_tar_gz = NULL)
```

Arguments

make_sure make sure to do this

taxonkit_tar_gz your download taxonkit_tar_gz file from <https://github.com/shenwei356/taxonkit/releases/>

Value

No value

See Also

Other Rtaxonkit: [check_taxonkit\(\)](#), [download_taxonkit_dataset\(\)](#), [name_or_id2df\(\)](#), [taxonkit_filter\(\)](#), [taxonkit_lca\(\)](#), [taxonkit_lineage\(\)](#), [taxonkit_list\(\)](#), [taxonkit_name2taxid\(\)](#), [taxonkit_reformat\(\)](#)

kwtest	<i>KW test</i>
--------	----------------

Description

KW test

Usage

```
kwtest(otutab, group_df, method = "kruskal.test")
```

Arguments

otutab	otutab
group_df	a dataframe with rowname same to dist and one group column
method	"kruskal.test", see compare_means

Value

res

Examples

```
data(otutab, package = "pcutils")
kwtest(otutab, metadata["Group"]) -> res
bbtt(res, pvalue = "p.format")
```

load_mpa_df	<i>Load a metaphlan format data.frame</i>
-------------	---

Description

Load a metaphlan format data.frame

Usage

```
load_mpa_df(mpa_df, sum_unidentified = TRUE, rank_prefixes = pc_rank_prefixes)
```

Arguments

mpa_df	metaphlan format data.frame, rownames are taxon, all value are numeric.
sum_unidentified	logical, whether to sum the unidentified reads to the correspond level.
rank_prefixes	a named vector of rank prefixes, default is pc_rank_prefixes.

Value

a list

load_N_data	<i>Load N-cycle data</i>
-------------	--------------------------

Description

Load N-cycle data

Usage

```
load_N_data()
```

Value

list

References

Tu, Q., Lin, L., Cheng, L., Deng, Y. & He, Z. (2019) NCycDB: a curated integrative database for fast and accurate metagenomic profiling of nitrogen cycling genes. *Bioinformatics* 35, 1040–1048.
Kuypers, M. M. M., Marchant, H. K. & Kartal, B. (2018) The microbial nitrogen-cycling network. *Nat Rev Microbiol* 16, 263–276.

mat_dist	<i>Calculate distance for otutab</i>
----------	--------------------------------------

Description

Calculate distance for otutab

Usage

```
mat_dist(otutab, method = "bray", spe_nwk = NULL)
```

Arguments

otutab	an otutab data.frame, samples are columns, taxa are rows.
method	Dissimilarity index, partial match to "bray", "euclidean"...see vegdist ; unifrac
spe_nwk	a phylo tree if use unifrac...

Value

dist

Examples

```
data(otutab, package = "pcutils")
mat_dist(otutab)
```

micro_sbatch	<i>Microbiome sbatch</i>
--------------	--------------------------

Description

Microbiome sbatch

Usage

```
micro_sbatch(
  work_dir = "/share/home/jianglab/pengchen/work/asthma/",
  step = "fastp",
  all_sample_num = 40,
  array = 1,
  partition = "cpu",
  cpus_per_task = 1,
  mem_per_cpu = "2G"
)
```

Arguments

work_dir	work_dir
step	"fastp","rm_human","megahit","prodigal","salmon-quant",...
all_sample_num	all sample number
array	array number
partition	partition
cpus_per_task	cpus_per_task
mem_per_cpu	mem_per_cpu, "2G"

Value

No value

multi_bar	<i>Difference analysis</i>
-----------	----------------------------

Description

Difference analysis

Usage

```
multi_bar(  
  otutab,  
  group_df,  
  mode = 1,  
  text_df = NULL,  
  text_x = NULL,  
  text_angle = -90,  
  errorbar = "bottom"  
)
```

Arguments

otutab	otutab
group_df	a dataframe with rowname same to dist and one group column
mode	1~2
text_df	text_df
text_x	text_x
text_angle	text_angle
errorbar	top, bottom, none

Value

ggplot

Examples

```
data(otutab, package = "pcutils")  
multi_bar(otutab[1:10, ], metadata["Group"])
```

multi_conf	<i>Difference analysis</i>
------------	----------------------------

Description

Difference analysis

Usage

```
multi_conf(otutab, group_df)
```

Arguments

otutab	otutab
group_df	a dataframe with rowname same to dist and one group column

Value

ggplot

Examples

```
data(otutab, package = "pcutils")
multi_conf(otutab[1:10, 1:12], metadata["Group"])
```

myRDA	<i>RDA</i>
-------	------------

Description

RDA

Usage

```
myRDA(
  otutab,
  env,
  norm = TRUE,
  scale = FALSE,
  choose_var = FALSE,
  direction = "forward",
  nperm = 499,
  verbose = TRUE,
  method = "rda",
  dist = "bray"
)
```

```

myCCA(
  otutab,
  env,
  norm = TRUE,
  scale = FALSE,
  choose_var = FALSE,
  nperm = 499,
  verbose = TRUE
)

myCAP(
  otutab,
  env,
  norm = TRUE,
  scale = FALSE,
  choose_var = FALSE,
  nperm = 499,
  verbose = TRUE,
  dist = "bray"
)

```

Arguments

otutab	an otutab data.frame, samples are columns, taxa are rows.
env	environmental factors
norm	should normalize? (default:TRUE)
scale	should scale species? (default:FALSE)
choose_var	should choose variables? use forward step
direction	The direction of the stepwise selection, "both", "forward" or "backward", default is "forward"
nperm	number of permutation
verbose	verbose
method	"rda", "cca", "cap", "dbrda"
dist	The name of the dissimilarity (or distance) index for "cap" or "dbrda", for vegdist

Value

rda/cca

See Also

[vegdist](#); [unifrac](#)

Examples

```
data(otutab, package = "pcutils")
env <- metadata[, 6:10]
# RDA
myRDA(otutab, env) -> phy.rda
RDA_plot(phy.rda, "Group", metadata)
```

name_or_id2df

*Transfer taxon name or taxid to the lineage dataframe***Description**

Transfer taxon name or taxid to the lineage dataframe

Usage

```
name_or_id2df(
  name_or_id,
  mode = "name",
  add_prefix = TRUE,
  fill_miss_rank = TRUE,
  data_dir = NULL
)
```

Arguments

name_or_id	name or taxid
mode	"id" or "name"
add_prefix	add_prefix
fill_miss_rank	fill_miss_rank
data_dir	directory containing nodes.dmp and names.dmp (default "/Users/asa/.taxonkit")

Value

dataframe

See Also

Other Rtaxonkit: [check_taxonkit\(\)](#), [download_taxonkit_dataset\(\)](#), [install_taxonkit\(\)](#), [taxonkit_filter\(\)](#), [taxonkit_lca\(\)](#), [taxonkit_lineage\(\)](#), [taxonkit_list\(\)](#), [taxonkit_name2taxid\(\)](#), [taxonkit_reformat\(\)](#)

Examples

```
## Not run:
name_or_id2df(c("Homo sapiens", "Akkermansia muciniphila ATCC BAA-835"))

## End(Not run)
```

ncm

*Sloan Neutral Model***Description**

Sloan Neutral Model

Plot ncm_res

Usage

```
ncm(otutab, model = "nls")

## S3 method for class 'ncm_res'
plot(
  x,
  mycols = c(Above = "#069870", Below = "#e29e02", In = "#1e353a"),
  text_position = NULL,
  pie_text_params = list(size = 2.5),
  ...
)
```

Arguments

otutab	an otutab data.frame, samples are columns, taxa are rows.
model	fit method, one of "nls", "mle"
x	a ncm_res object
mycols	mycols
text_position	text_position
pie_text_params	pie text parameters
...	add

Value

ncm_res
ggplot

References

Sloan, W. TRUE. et al. (2006) Quantifying the roles of immigration and chance in shaping prokaryote community structure. *Environmental Microbiology* 8, 732–740.

Examples

```
if (requireNamespace("Hmisc") && requireNamespace("minpack.lm")) {
  data(otutab, package = "pcutils")
  ncm(otutab) -> ncm_res
  plot(ncm_res)
}
```

nst	<i>Calculate NST for each group</i>
-----	-------------------------------------

Description

Calculate NST for each group

Usage

```
nst(otutab, group_df, threads = 1, file = NULL, rep = 20, save = FALSE)
```

Arguments

otutab	an otutab data.frame, samples are columns, taxa are rows.
group_df	a dataframe with rowname and one group column
threads	default:4
file	filename to save
rep	repeat numbers: suggest 999
save	save the file

Value

a b_dist object, dis is MSTij

References

Ning, D., Deng, Y., Tiedje, J. M. & Zhou, J. (2019) A general framework for quantitatively assessing ecological stochasticity. *Proceedings of the National Academy of Sciences* 116, 16892–16898.

Examples

```
if (requireNamespace("NST")) {
  library(ggplot2)
  data(otutab, package = "pcutils")
  nst(otutab, metadata["Group"]) -> nst_res
  plot(nst_res, c_group = "intra") + geom_hline(yintercept = 0.5, lty = 2) + ylab("NST")
}
```

nti_rc	<i>Calculate b_NTI and RC_bray for each group</i>
--------	---

Description

Calculate b_NTI and RC_bray for each group
 Plot NTI_RC object

Usage

```
nti_rc(
  otutab,
  phylo,
  group_df,
  threads = 1,
  file = NULL,
  rep = 20,
  save = FALSE
)

## S3 method for class 'NTI_RC'
plot(x, ...)
```

Arguments

otutab	an otutab data.frame, samples are columns, taxa are rows.
phylo	a phylo object
group_df	a dataframe with rowname and one group column
threads	default:4
file	filename to save
rep	repeat numbers: suggest 999
save	save the file
x	NTI_RC object
...	pass to stackplot

Value

a b_dist object, dis is MSTij
 ggplot

References

Ning, D., Deng, Y., Tiedje, J. M. & Zhou, J. (2019) A general framework for quantitatively assessing ecological stochasticity. *Proceedings of the National Academy of Sciences* 116, 16892–16898.

Examples

```

if (requireNamespace("NST") && requireNamespace("pctax")) {
  data(otutab, package = "pcutils")
  pctax::df2tree(taxonomy) -> phylo
  nti_rc(otutab, phylo, metadata["Group"]) -> nti_res
  plot(nti_res)
}

```

pc_otu

Create a pc_otu class object

Description

Create a pc_otu class object

Usage

```
pc_otu(otutab = data.frame(), metadata = data.frame(), taxonomy = NULL, ...)
```

Arguments

otutab	an otutab data.frame, samples are columns, taxa are rows.
metadata	a metadata data.frame, samples are rows
taxonomy	a taxonomy data.frame, look out the rowname of taxonomy and otutab should matched!
...	add

Value

pc_otu

Examples

```

data(otutab, package = "pcutils")
pc_tax1 <- pc_otu(otutab, metadata)
print(pc_tax1)

```

pc_tax1	<i>test data (pc_otu class) for pc_tax package.</i>
---------	---

Description

an otutab, metadata and a taxonomy table.

Format

a pc_otu contains an otutab, metadata and a taxonomy table.

tbls contains otutable rawdata

metas contains metadata

otus contains taxonomy table

pc_valid	<i>Judge pc_otu is valid or not</i>
----------	-------------------------------------

Description

Judge pc_otu is valid or not

Usage

```
pc_valid(pc)
```

Arguments

pc a pc_otu object

Value

logical

permanova

*Permanova between a otutab and a variable***Description**

Permanova between a otutab and a variable

Usage

```
permanova(
  otutab,
  envs,
  norm = TRUE,
  each = TRUE,
  method = "adonis",
  dist = "bray",
  nperm = 999,
  ...
)
```

Arguments

otutab	an otutab data.frame, samples are columns, taxa are rows.
envs	factors need to test
norm	should normalize?(default:TRUE)
each	test factor one by one, rather than whole
method	adonis/mrpp/anosim/mantel
dist	if use pcoa or nmds, your can choose a dist method (default: bray)
nperm	numbers of permutations to perform
...	additional

Value

a g_test object with these columns

group	the test group or factor
r	relationship
r2	model R-square
p_value	model test p_value
sig	whether significant

Examples

```
data(otutab, package = "pcutils")
permanova(otutab, metadata[, c(2:10)]) -> adonis_res
print(adonis_res)
plot(adonis_res)
```

plot.a_res	<i>Plot a_res object</i>
------------	--------------------------

Description

Plot a_res object

Usage

```
## S3 method for class 'a_res'
plot(x, group, metadata, ...)
```

Arguments

x	a a_res object
group	one of colname of metadata
metadata	metadata
...	additional parameters for group_box or my_lm

Value

patchwork object,you can change theme with &

See Also

[a_diversity](#)

plot.b_res	<i>Plot a b_res</i>
------------	---------------------

Description

Plot a b_res

Usage

```
## S3 method for class 'b_res'
plot(
  x,
  Group,
  metadata = NULL,
  Group2 = NULL,
  mode = 1,
  bi = FALSE,
  Topn = 10,
```

```

    rate = 1,
    margin = FALSE,
    margin_label = TRUE,
    permanova_res = NULL,
    text_param = list(),
    box_margin = TRUE,
    box_param = list(),
    pal = NULL,
    sample_label = TRUE,
    stat_ellipse = TRUE,
    ellipse_level = 0.95,
    add_centroid_label = TRUE,
    coord_fix = FALSE,
    bi_text_size = 3,
    ...
)

```

Arguments

x	a b_res object
Group	group vector for color
metadata	metadata contain Group
Group2	mapping point shape
mode	plot mode:1~3
bi	plot variables segments?
Topn	how many variables to show?
rate	segments length rate
margin	plot the margin boxplot?
margin_label	margin_label, TRUE
permanova_res	permanova result
text_param	text_param for annotate
box_margin	margin plot box or density?
box_param	box_param for group_box
pal	colors for group
sample_label	plot the labels of samples?
stat_ellipse	plot the stat_ellipse?
ellipse_level	the level of stat_ellipse, default 0.95
add_centroid_label	add the centroid label in mode 3?
coord_fix	fix the coordinates y/x ratio
bi_text_size	biplot text size
...	add

Value

a ggplot

See Also

[b_analyse](#)

plot.g_test	<i>Plot g_test</i>
-------------	--------------------

Description

Plot g_test

Usage

```
## S3 method for class 'g_test'
plot(x, ...)
```

Arguments

x	a g_test object
...	add

Value

ggplot

See Also

[permanova](#)

plot.pro_res	<i>Plot pro_res</i>
--------------	---------------------

Description

Plot pro_res

Usage

```
## S3 method for class 'pro_res'
plot(x, group, metadata = NULL, pal = NULL, ...)
```

Arguments

x	pro_res
group	group
metadata	metadata
pal	pal
...	add

Value

a ggplot

plot.time_cm	<i>Plot time_cm</i>
--------------	---------------------

Description

Plot time_cm

Usage

```
## S3 method for class 'time_cm'  
plot(x, mem_thr = 0.6, ...)
```

Arguments

x	time_cm
mem_thr	membership threshold
...	add

Value

ggplot

plot_checkm2_res	<i>Visualize CheckM2 Genome Quality Assessment Results</i>
------------------	--

Description

This function creates a scatter plot showing genome completeness vs contamination with optional marginal density plots to display distributions.

Usage

```
plot_checkm2_res(
  checkm2_df,
  add_marginal = TRUE,
  marginal_type = "density",
  point_size = 0.6,
  base_size = 14,
  quality_thresholds = list(high_comp = 90, high_contam = 5, med_comp = 70, med_contam =
    10),
  filter_data = TRUE,
  min_quality_score = 50,
  min_completeness = 50,
  max_contamination = 10
)
```

Arguments

checkm2_df	Data frame. CheckM2 results containing at least columns: 'Completeness', 'Contamination', and 'Name'.
add_marginal	Logical. Whether to add marginal density plots using ggExtra. Default is TRUE.
marginal_type	Character. Type of marginal plot: "density", "histogram", "boxplot", or "violin". Default is "density".
point_size	Numeric. Size of points in scatter plot. Default is 0.6.
base_size	Numeric. Base font size for the plot. Default is 14.
quality_thresholds	List. Custom thresholds for quality classification. Default list(high_comp = 90, high_contam = 5, med_comp = 70, med_contam = 10)
filter_data	Logical. Whether to filter low-quality genomes. Default is TRUE.
min_quality_score	Numeric. Minimum quality score for filtering. Default is 50.
min_completeness	Numeric. Minimum completeness for filtering. Default is 50.
max_contamination	Numeric. Maximum contamination for filtering. Default is 10.

Value

A ggplot object or ggExtra plot object if marginal plots are added.

plot_contigs_quality *Visualize Contigs Quality Metrics*

Description

This function creates a scatter plot to visualize the quality metrics of contigs from geNomad and CheckV analysis results.

Usage

```
plot_contigs_quality(  
  genomad_out_res,  
  show_valid = FALSE,  
  point_size = 1.4,  
  point_alpha = 0.6,  
  base_size = 15  
)
```

Arguments

genomad_out_res	List. Output object from pre_genomad function containing virus summary data.
show_valid	Logical. Whether to display only valid (high-quality) viral sequences. Default is FALSE (show all sequences).
point_size	Numeric. Size of the points in the plot. Default is 1.4.
point_alpha	Numeric. Transparency of the points (0-1). Default is 0.6.
base_size	Numeric. Base font size for the plot. Default is 15.

Value

A ggplot object displaying contigs quality metrics.

plot_element_cycle	<i>Plot element cycle</i>
--------------------	---------------------------

Description

Plot element cycle

Usage

```
plot_element_cycle(  
  cycle = "Nitrogen cycle",  
  anno_df = NULL,  
  only_anno = FALSE,  
  cell_fill = NA,  
  cell_color = "orange",  
  use_chinese = FALSE,  
  chemical_size = 7,  
  chemical_bold = TRUE,  
  chemical_color = "black",  
  chemical_label = TRUE,  
  reaction_width = 1,  
  reaction_arrow_size = 4,  
  reaction_arrow_closed = TRUE,  
  gene_or_ko = "gene",  
  gene_size = 3,  
  gene_x_offset = 0.3,  
  gene_y_offset = 0.15,  
  gene_label = TRUE,  
  gene_color = NULL,  
  gene_bold = TRUE,  
  gene_italic = TRUE,  
  gene_label_fill = "white"  
)
```

Arguments

cycle	one of c("Carbon cycle", "Nitrogen cycle", "Phosphorus cycle", "Sulfur cycle", "Iron cycle")
anno_df	anno_df, columns should contains Gene or KO and Group
only_anno	only show genes in anno_df?
cell_fill	cell fill color
cell_color	cell border color
use_chinese	use chinese label?
chemical_size	chemical text size
chemical_bold	chemical text bold

chemical_color	chemical text color
chemical_label	chemical text in geom_label or geom_text?
reaction_width	reaction line width
reaction_arrow_size	reaction arrow size
reaction_arrow_closed	reaction arrow closed?
gene_or_ko	"gene" or "ko"
gene_size	gene text size
gene_x_offset	gene_x_offset
gene_y_offset	gene_y_offset
gene_label	gene text in geom_label or geom_text?
gene_color	gene text color
gene_bold	gene text bold?
gene_italic	gene text italic?
gene_label_fill	gene label fill color

Value

ggplot

Examples

```
if (requireNamespace("ggforce")) plot_element_cycle()
```

plot_gtdb_tr

Plot GTDB-Tk Phylogenetic Tree with Taxonomic Coloring

Description

This function creates a circular phylogenetic tree visualization from GTDB-Tk results, with tips colored by specified taxonomic level. It can optionally represent only one genome per species.

Usage

```
plot_gtdb_tr(
  gtdb_res,
  tree,
  tax_level = "Phylum",
  represented = TRUE,
  layout = "fan",
  branch_size = 0.2,
  color_na = "black"
)
```

Arguments

gtddb_res	A data frame containing GTDB-Tk classification results, typically the output from pre_gtddb_tk function.
tree	A phylogenetic tree object (phylo class) containing all genomes.
tax_level	Character specifying the taxonomic level for coloring tips. Default is "Phylum". Other options include "Class", "Order", "Family", "Genus", "Species".
represented	Logical indicating whether to include only one representative genome per species. Default is TRUE.
layout	Character specifying the tree layout. Options include "fan", "circular", "rectangular", etc. Default is "fan".
branch_size	Numeric value for branch line width. Default is 0.2.
color_na	Character specifying color for tips with missing taxonomic information. Default is "black".

Value

A ggplot object containing the phylogenetic tree visualization.

plot_N_cycle	<i>Plot the N-cycling pathway and genes</i>
--------------	---

Description

Plot the N-cycling pathway and genes

Usage

```
plot_N_cycle(
  my_N_genes = NULL,
  just_diff = FALSE,
  path_col = NULL,
  type_col = c(up = "red", down = "blue", none = NA),
  fill_alpha = 0.5,
  arrow_size = 0.1,
  line_width = 1,
  title = "Nitrogen cycling",
  legend.position = c(0.85, 0.15)
)
```

Arguments

my_N_genes	dataframe, "Gene_families", "type" should in colnames of my_N_genes
just_diff	logical, just plot the different genes?
path_col	colors of pathways

type_col	colors of types
fill_alpha	alpha, default 0.5
arrow_size	arrow_size, default 0.1
line_width	line_width, default 1
title	title, default "Nitrogen cycling"
legend.position	default c(0.85,0.15)

Value

ggplot

Examples

```

N_data <- load_N_data()
my_N_genes <- data.frame(
  `Gene_families` = sample(N_data$N_genes$Gene_families, 10, replace = FALSE),
  change = rnorm(10), check.names = FALSE
)
my_N_genes <- dplyr::mutate(my_N_genes,
  type = ifelse(change > 0, "up", ifelse(change < 0, "down", "none"))
)
plot_N_cycle(my_N_genes, just_diff = FALSE, fill_alpha = 0.2)
# ggsave(filename = "test.pdf", width = 14, height = 10)

```

plot_one_phage

*Plot Individual Phage Genome Structure with Annotations***Description**

This function creates a circular genome map for a single phage, displaying gene annotations, functional categories, and other genomic features. It automatically extracts topology information and provides flexible parameter customization.

Usage

```

plot_one_phage(
  one_phage,
  genomad_out_res,
  anno_table = NULL,
  y_var = "strand",
  fill_var = "COG",
  label_var = c("annotation_description"),
  label_wrap = 20,
  palette = "Set3"
)

```

Arguments

one_phage	Character. Phage sequence identifier (e.g., "k141_10408").
genomad_out_res	List. Output object from pre_genomad function containing virus summary and gene data.
anno_table	Data frame. Optional annotation table with gene information for left join. Must contain a 'gene' column for merging with genomad gene data.
y_var	Character. Variable for y-axis mapping. Default is "strand".
fill_var	Character. Variable for fill color mapping. Default is "COG".
label_var	Character vector. Columns to use for gene description labels.
label_wrap	Integer. Width for wrapping gene description labels. Default is 20.
palette	Character. Color palette for fill categories. Default is "Set3".

Details

The function automatically extracts topology information from the virus summary data and creates a polar coordinate visualization showing:

- Gene arrows indicating direction and position
- Flexible y-axis and fill color mappings
- Genome length and automatically detected topology information
- Gene descriptions with intelligent label placement

Value

A ggplot object displaying the circular phage genome map.

plot_two_tree	<i>Plot two trees in one plot</i>
---------------	-----------------------------------

Description

Plot two trees in one plot

Usage

```
plot_two_tree(
  tree1,
  tree2,
  edge_df = NULL,
  tree2_x = 10,
  filter_link = FALSE,
  tree1_param = list(),
  tree2_param = list(),
```

```

    line_param = list(),
    tree1_tip = FALSE,
    tip1_param = list(),
    tree2_tip = FALSE,
    tip2_param = list(),
    tree1_highlight = NULL,
    highlight1_param = list(),
    highlight1_scale = NULL,
    tree2_highlight = NULL,
    highlight2_param = list(),
    highlight2_scale = ggplot2::scale_fill_hue(na.value = NA)
  )

```

Arguments

tree1	phylo object
tree2	phylo object
edge_df	dataframe with edge information, containing "from" and "to" columns
tree2_x	x position of tree2
filter_link	filter the link between tree1 and tree2
tree1_param	parameters for geom_tree
tree2_param	parameters for geom_tree
line_param	parameters for geom_line
tree1_tip	tree tip label
tip1_param	parameters for geom_tiplab
tree2_tip	tree tip label
tip2_param	parameters for geom_tiplab
tree1_highlight	tree1 highlight data.frame
highlight1_param	parameters for geom_highlight
highlight1_scale	scale_fill_ for highlight1
tree2_highlight	tree2 highlight data.frame
highlight2_param	parameters for geom_highlight
highlight2_scale	scale_fill_ for highlight2

Value

ggplot object

Examples

```

if (requireNamespace("ggtree")) {
  data(otutab, package = "pcutils")
  df2tree(taxonomy[1:50, ]) -> tax_tree
  df2tree(taxonomy[51:100, ]) -> tax_tree2
  link <- data.frame(from = sample(tax_tree$tip.label, 20), to = sample(tax_tree2$tip.label, 20))
  plot_two_tree(tax_tree, tax_tree2, link)
}

```

```
pre_assembly_stats      Prepare the result from assembly_stats (.json file)
```

Description

Prepare the result from assembly_stats (.json file)

Usage

```
pre_assembly_stats(jsonfiles)
```

Arguments

jsonfiles the directory contains .json file

Value

data.frame

```
pre_fastp              Prepare the result from fastp (.json file)
```

Description

Prepare the result from fastp (.json file)

Usage

```
pre_fastp(jsonfiles, prefix = c("Raw", "Clean"))
```

Arguments

jsonfiles the directory contains .json file
 prefix default c("Raw", "Clean"), for the before filtering and after filtering.

Value

data.frame

```
pre_format_report
```

Preprocess MPA Species Abundance and Taxonomy Data

Description

This function reads a species abundance profile from kraken2 format_report output, performs filtering to remove low-abundance and unwanted taxa, and extracts a clean taxonomy table for downstream analysis.

Usage

```
pre_format_report(dir = "", exclude = NULL, relative_threshold = 1e-04)
```

Arguments

<code>dir</code>	Character. Path to the directory containing the input file "mpa_profile_species.txt". Default is an empty string (current directory).
<code>exclude</code>	Character. Pattern to exclude specific taxa (e.g., "g__Streptococcus"). Uses <code>grepl()</code> for pattern matching. Default is NULL (no exclusion).
<code>relative_threshold</code>	Numeric. Relative abundance threshold for filtering low-abundance taxa. Taxa with mean relative abundance below this threshold will be removed. Default is 1e-4.

Details

The function performs the following steps:

1. Reads the Metaphlan species profile table
2. Removes "unclassified" entries and "cellular_organisms" category
3. Filters out taxa matching the exclude pattern (if provided)
4. Applies relative abundance filtering using `pcutils::rm_low`
5. Extracts and formats taxonomy information from the Metaphlan-style names
6. Cleans species names by removing the "s__" prefix

Value

A list with two components:

<code>species</code>	A matrix of filtered species abundance data
<code>species_taxonomy</code>	A data.frame of taxonomy information for each species

pre_genomad

*Preprocess geNomad and CheckV Output Results***Description**

This function automatically processes geNomad output files by detecting sample names from the directory structure and optionally integrates CheckV quality assessment results.

Usage

```
pre_genomad(
  genomad_out_dir = "",
  checkV_out_dir = NULL,
  provirus = TRUE,
  filter = TRUE,
  checkV_out_prefix = NULL,
  min_length = 1000,
  min_completeness = 50
)
```

Arguments

genomad_out_dir	Character. Path to the geNomad output directory. This directory should contain sample-specific subdirectories with the pattern "*.contigs_summary".
checkV_out_dir	Character. Optional path to the CheckV output directory. If provided, quality summary will be integrated. Default is NULL.
provirus	Logical. Whether to identify and separate provirus sequences. Default is TRUE.
filter	Logical. Whether to apply quality filtering to viral sequences. Default is TRUE.
checkV_out_prefix	Character. Optional prefix to remove from CheckV contig IDs.
min_length	Numeric. Minimum sequence length for filtering. Default is 1000.
min_completeness	Numeric. Minimum completeness score for CheckV filtering. Default is 50.

Details

The function automatically detects sample names by searching for directories with the pattern "*.contigs_summary" within the genomad_out_dir. It then extracts the sample name by removing the ".contigs_summary" suffix.

Value

An object of class "virus_res" containing four components:

sample	Detected sample name
--------	----------------------

```

virus_summary   Integrated data frame with geNomad and optional CheckV results
virus_genes     Gene-level annotations from geNomad
valid_virus     Filtered high-quality viral sequences

```

Examples

```

## Not run:
# Basic usage - sample name will be automatically detected
virus_results <- pre_genomad(genomad_out_dir = "~/Documents/R/Lung_vrome/data/genomad_out2/")

# Access the detected sample name
sample_name <- virus_results$sample
print(paste("Detected sample:", sample_name))

## End(Not run)

```

pre_gtdb_tk	<i>Preprocess GTDB-Tk Classification Results</i>
-------------	--

Description

This function reads and processes the output files from a GTDB-Tk classify workflow. It combines bacterial (bac120) and archaeal (ar53) classification summaries and phylogenetic trees (if available) into a unified format.

Usage

```
pre_gtdb_tk(classify_dir)
```

Arguments

```

classify_dir    A character string specifying the path to the GTDB-Tk classify output di-
                  rectory. This directory should contain files like gtdbtk.bac120.summary.tsv,
                  gtdbtk.bac120.classify.tree, etc.

```

Details

The function performs the following steps:

1. Checks if the provided directory exists and contains the necessary *.summary.tsv files.
2. Reads the bacterial backbone tree.
3. If an archaeal tree file exists, it binds it to the bacterial tree.
4. Reads and combines all *.summary.tsv files in the directory.
5. Parses the semicolon-separated classification string into separate columns for each taxonomic rank.
6. Ensures the resulting taxonomy table has standard ranks (Domain, Phylum, Class, Order, Family, Genus, Species).

Value

A list with two components:

gtddb_res A data frame containing the combined taxonomic classification for all genomes. The classification column is parsed into standard taxonomic ranks (Domain to Species).

tree A phylogenetic tree (phylo object) combining the bacterial and (if present) archaeal trees.

pre_tax_table	<i>Complete a taxonomy table</i>
---------------	----------------------------------

Description

Complete a taxonomy table

Usage

```
pre_tax_table(  
  tax_table,  
  tax_levels = c("k", "p", "c", "o", "f", "g", "s", "st"),  
  na_tax = "Unclassified|uncultured|Ambiguous|Unknown|unknown|metagenome|Unassig",  
  ignore.case = TRUE,  
  na_repalce = "Unknown",  
  add_prefix = TRUE  
)
```

Arguments

tax_table	taxonomy table
tax_levels	a vector whose length longer than ncol(taxdf), use to be prefix. Default: c("k", "p", "c", "o", "f", "g", "s", "st")
na_tax	grepl some words and turn to na_repalce, default: "Unclassified uncultured Ambiguous Unknown unknown metagenome Unassig"
ignore.case	ignore.case for na_tax
na_repalce	defalut: Unknown
add_prefix	add prefix? logical

Value

a good taxonomy table

References

MicrobiotaProcess

Examples

```
taxmat <- matrix(sample("onelevel", 7 * 2, replace = TRUE), nrow = 2, ncol = 7) %>% as.data.frame()  
colnames(taxmat) <- c("Kingdom", "Phylum", "Class", "Order", "Family", "Genus", "Species")  
pre_tax_table(taxmat)
```

<code>print.pc_otu</code>	<i>Print</i>
---------------------------	--------------

Description

Print

Usage

```
## S3 method for class 'pc_otu'  
print(x, ...)
```

Arguments

<code>x</code>	<code>pc_otu</code>
<code>...</code>	<code>add</code>

Value

No value

<code>print.virus_res</code>	<i>Print method for virus_res objects</i>
------------------------------	---

Description

Print method for virus_res objects

Usage

```
## S3 method for class 'virus_res'  
print(x, ...)
```

Arguments

<code>x</code>	An object of class virus_res
<code>...</code>	Additional arguments (not used)

procrustes_analyse	<i>Procrustes Rotation of Two Configurations and PROTEST</i>
--------------------	--

Description

Procrustes Rotation of Two Configurations and PROTEST

Usage

```
procrustes_analyse(b_res1, b_res2, nperm = 999, ...)
```

Arguments

b_res1	Target matrix
b_res2	Matrix to be rotated
nperm	numbers of permutations to perform
...	additional

Value

pro_res

Examples

```
data(otutab, package = "pcutils")
b_analyse(otutab, method = "pca") -> b_res1
b_analyse(otutab * abs(rnorm(10))), method = "pca") -> b_res2
pro_res <- procrustes_analyse(b_res1, b_res2)
plot(pro_res, "Group", metadata)
```

rarefaction	<i>Rarefy a otutab</i>
-------------	------------------------

Description

Rarefy a otutab

Usage

```
rarefaction(otutab, sample = NULL)
```

Arguments

otutab	otutab
sample	number

Value

a rarefied otutab

Examples

```
data(otutab, package = "pcutils")
rarefaction(otutab)
```

rare_curve_sample	<i>Rare the sample</i>
-------------------	------------------------

Description

Rare the sample
Plot a rare curve

Usage

```
rare_curve_sample(otutab, rep = 30, count_cutoff = 1)

## S3 method for class 'rare_res'
plot(x, ...)
```

Arguments

otutab	otutab
rep	repeats number
count_cutoff	cutoff to be 0
x	AOR object
...	add

Value

ggplot
ggplot

Examples

```
data(otutab, package = "pcutils")
a <- rare_curve_sample(otutab)
plot(a)
```

rare_curve_species	<i>Rare the species</i>
--------------------	-------------------------

Description

Rare the species

Usage

```
rare_curve_species(  
  otutab,  
  step = 2000,  
  method = "richness",  
  mode = 2,  
  reps = 3,  
  threads = 1,  
  verbose = TRUE  
)
```

Arguments

otutab	otutab
step	default 2000
method	one of "richness", "chao1", "ace", "gc", "shannon", "simpson", "pd", "pielou"
mode	1 for little table, 2 for big
reps	reps
threads	use how many threads to calculate (default:1)
verbose	verbose

Value

ggplot

Examples

```
data(otutab, package = "pcutils")  
a <- rare_curve_species(otutab, mode = 1)  
plot(a)
```

RCbray1

*Calculate RCbray-curtis***Description**

Calculate RCbray-curtis

Usage

```
RCbray1(
  otutab,
  reps = 9,
  threads = 1,
  classic_metric = TRUE,
  split_ties = TRUE
)
```

Arguments

otutab	otutab
reps	how many simulation performed?
threads	use how many threads to calculate (default:4)
classic_metric	standardizes the metric to range from -1 to 1
split_ties	adds half of the number of null observations that are equal to the observed number of shared species to the calculation- this is highly recommended

Details

Parallelized version of the Raup-Crick algorithm for "abundance" data (Stegen et al. 2013).

Value

a dist

Examples

```
if (requireNamespace("picante")) {
  data(otutab, package = "pcutils")
  df2tree(taxonomy) -> phylo
  b_NTI1(phylo, otutab) -> bnti_res
  RCbray1(otutab, reps = 9) -> rc_res

  data.frame(
    type = factor(c("Homo_S", "Heter_S", "Homo_D", "D_limit", "Undominated"),
      levels = c("Homo_S", "Heter_S", "Homo_D", "D_limit", "Undominated")
    ),
    number = c(
```

```

        sum(bnti_res < (-2)), sum(bnti_res > 2),
        sum((abs(bnti_res) < 2) & (abs(rc_res) < 0.95)),
        sum((abs(bnti_res) < 2) & (rc_res < (-0.95))),
        sum((abs(bnti_res) < 2) & (rc_res > 0.95))
    )
) -> com_pro
pcutils::gghuan(com_pro, reorder = FALSE)
}

```

RDA_plot

Plot RDA res

Description

Plot RDA res

Usage

```

RDA_plot(
  phy.rda,
  Group,
  metadata = NULL,
  Group2 = NULL,
  env_rate = 1,
  mode = 1,
  tri = FALSE,
  Topn = 10,
  rate = 1,
  margin = FALSE,
  box_margin = TRUE,
  pal = NULL,
  sample_label = TRUE,
  stat_ellipse = TRUE,
  ellipse_level = 0.95,
  coord_fix = FALSE,
  bi_text_size = 3,
  env_text_param = NULL,
  ...
)

```

Arguments

phy.rda	rda/cca object
Group	group vector for color
metadata	metadata contain Group
Group2	mapping point shape

env_rate	default 1
mode	plot mode:1~3
tri	plot variables segments?
Topn	how many variables to show?
rate	segments length rate
margin	plot the margin boxplot?
box_margin	margin plot box or density?
pal	colors for group
sample_label	plot the labels of samples?
stat_ellipse	plot the stat_ellipse?
ellipse_level	the level of stat_ellipse, defalut 0.95
coord_fix	fix the coordinates y/x ratio
bi_text_size	biplot text size
env_text_param	parameters pass to geom_text
...	add

Value

ggplot

See Also

[myRDA](#)

stamp_plot	<i>Stamp style plot</i>
------------	-------------------------

Description

Stamp style plot

Usage

```
stamp_plot(otutab, group_df, set_order = NULL, pal = NULL)
```

Arguments

otutab	otutab
group_df	a dataframe with rowname same to dist and one group column
set_order	set order of factor levels
pal	palette

Value

ggplot

Examples

```
data(otutab, package = "pcutils")
if (requireNamespace("ggfun")) stamp_plot(otutab[1:10, 1:12], metadata["Group"])
```

suijisenlin	<i>RandomForest</i>
-------------	---------------------

Description

RandomForest

Usage

```
suijisenlin(otutab, group_df, topN = 10)
```

Arguments

otutab	otutab
group_df	a dataframe with rowname same to dist and one group column
topN	default: 10

Value

diff

Examples

```
if (requireNamespace("randomForest")) {
  data(otutab, package = "pcutils")
  suijisenlin(otutab, metadata["Group"]) -> rf_res
}
```

summary.pc_otu	<i>Summary pc_otu</i>
----------------	-----------------------

Description

Summary pc_otu

Usage

```
## S3 method for class 'pc_otu'  
summary(object, ...)
```

Arguments

object	pc_otu
...	add

Value

No value

Examples

```
data("pc_tax1")  
summary(pc_tax1)
```

taxonkit_filter	<i>Filter TaxIDs based on Taxonomic Ranks</i>
-----------------	---

Description

This function uses the "taxonkit filter" command to filter TaxIDs based on taxonomic ranks.

Usage

```
taxonkit_filter(  
  file_path,  
  black_list = NULL,  
  discard_noranks = FALSE,  
  discard_root = FALSE,  
  equal_to = NULL,  
  higher_than = NULL,  
  lower_than = NULL,  
  rank_file = NULL,  
  root_taxid = NULL,
```

```

    save_predictable_norank = FALSE,
    taxid_field = NULL,
    text = FALSE,
    data_dir = NULL
  )

```

Arguments

<code>file_path</code>	The path to the input file containing TaxIDs. Or file text (text=TRUE)
<code>black_list</code>	A character vector specifying the ranks to discard.
<code>discard_noranks</code>	Logical value indicating whether to discard all ranks without order (default is FALSE).
<code>discard_root</code>	Logical value indicating whether to discard the root taxid (default is FALSE).
<code>equal_to</code>	A character vector specifying the ranks for which TaxIDs should be equal to.
<code>higher_than</code>	The rank above which the TaxIDs should be (exclusive).
<code>lower_than</code>	The rank below which the TaxIDs should be (exclusive).
<code>rank_file</code>	The path to a user-defined ordered taxonomic ranks file.
<code>root_taxid</code>	The root taxid (default is 1).
<code>save_predictable_norank</code>	Logical value indicating whether to save some special ranks without order when using lower_than (default is FALSE).
<code>taxid_field</code>	The field index of the taxid in the input file (default is 1).
<code>text</code>	logical
<code>data_dir</code>	directory containing nodes.dmp and names.dmp (default "/Users/asa/.taxonkit")

Value

A character vector containing the output of the "taxonkit filter" command.

See Also

Other Rtaxonkit: [check_taxonkit\(\)](#), [download_taxonkit_dataset\(\)](#), [install_taxonkit\(\)](#), [name_or_id2df\(\)](#), [taxonkit_lca\(\)](#), [taxonkit_lineage\(\)](#), [taxonkit_list\(\)](#), [taxonkit_name2taxid\(\)](#), [taxonkit_reformat\(\)](#)

Examples

```

## Not run:
taxids2 <- system.file("extdata/taxids2.txt", package = "pctax")
taxonkit_filter(taxids2, lower_than = "genus")

## End(Not run)

```

taxonkit_lca

Compute Lowest Common Ancestor (LCA) of TaxIDs

Description

This function uses the "taxonkit lca" command to compute the Lowest Common Ancestor (LCA) of TaxIDs.

Usage

```
taxonkit_lca(
  file_path,
  buffer_size = "1M",
  separator = " ",
  skip_deleted = FALSE,
  skip_unfound = FALSE,
  taxids_field = NULL,
  text = FALSE,
  data_dir = NULL
)
```

Arguments

file_path	The path to the input file containing TaxIDs. Or file text (text=TRUE)
buffer_size	The size of the line buffer (supported units: K, M, G).
separator	The separator for TaxIDs.
skip_deleted	Whether to skip deleted TaxIDs and compute with the remaining ones.
skip_unfound	Whether to skip unfound TaxIDs and compute with the remaining ones.
taxids_field	The field index of TaxIDs. Input data should be tab-separated (default 1).
text	logical
data_dir	directory containing nodes.dmp and names.dmp (default "/Users/asa/taxonkit")

Value

A character vector containing the computed LCAs.

See Also

Other Rtaxonkit: [check_taxonkit\(\)](#), [download_taxonkit_dataset\(\)](#), [install_taxonkit\(\)](#), [name_or_id2df\(\)](#), [taxonkit_filter\(\)](#), [taxonkit_lineage\(\)](#), [taxonkit_list\(\)](#), [taxonkit_name2taxid\(\)](#), [taxonkit_reformat\(\)](#)

Examples

```
## Not run:
taxonkit_lca("239934, 239935, 349741", text = TRUE, separator = ", ")

## End(Not run)
```

taxonkit_lineage	<i>Retrieve Taxonomic Lineage using taxonkit</i>
------------------	--

Description

Retrieve Taxonomic Lineage using taxonkit

Usage

```
taxonkit_lineage(
  file_path,
  delimiter = ";",
  no_lineage = FALSE,
  show_lineage_ranks = FALSE,
  show_lineage_taxids = FALSE,
  show_name = FALSE,
  show_rank = FALSE,
  show_status_code = FALSE,
  taxid_field = 1,
  text = FALSE,
  data_dir = NULL
)
```

Arguments

file_path	The path to the input file with taxonomic IDs. Or file text (text=TRUE)
delimiter	The field delimiter in the lineage (default ";").
no_lineage	Logical, indicating whether to exclude lineage information (default: FALSE).
show_lineage_ranks	Logical, indicating whether to append ranks of all levels in the lineage (default: FALSE).
show_lineage_taxids	Logical, indicating whether to append lineage consisting of taxids (default: FALSE).
show_name	Logical, indicating whether to append scientific name (default: FALSE).
show_rank	Logical, indicating whether to append rank of taxids (default: FALSE).
show_status_code	Logical, indicating whether to show status code before lineage (default: FALSE).
taxid_field	The field index of taxid. Input data should be tab-separated (default: 1).
text	logical,
data_dir	directory containing nodes.dmp and names.dmp (default "/Users/asa/.taxonkit")

Value

A character vector containing the taxonomic lineage information.

See Also

Other Rtaxonkit: [check_taxonkit\(\)](#), [download_taxonkit_dataset\(\)](#), [install_taxonkit\(\)](#), [name_or_id2df\(\)](#), [taxonkit_filter\(\)](#), [taxonkit_lca\(\)](#), [taxonkit_list\(\)](#), [taxonkit_name2taxid\(\)](#), [taxonkit_reformat\(\)](#)

Examples

```
## Not run:
taxonkit_lineage("9606\n63221", show_name = TRUE, show_rank = TRUE, text = TRUE)

## End(Not run)
```

taxonkit_list	<i>Taxonkit list</i>
---------------	----------------------

Description

This function uses Taxonkit to perform the "list" operation, which retrieves information about taxa based on their TaxIDs.

Usage

```
taxonkit_list(
  ids,
  indent = "  ",
  json = FALSE,
  show_name = FALSE,
  show_rank = FALSE,
  data_dir = NULL
)
```

Arguments

ids	A character vector of TaxIDs to retrieve information for.
indent	The indentation string to use for pretty-printing the output. Default is " ".
json	Logical value indicating whether to output the result in JSON format. Default is FALSE.
show_name	Logical value indicating whether to show the scientific names of taxa. Default is FALSE.
show_rank	Logical value indicating whether to show the ranks of taxa. Default is FALSE.
data_dir	directory containing nodes.dmp and names.dmp (default "/Users/asa/.taxonkit")

Value

The output of the Taxonkit list operation.

See Also

Other Rtaxonkit: [check_taxonkit\(\)](#), [download_taxonkit_dataset\(\)](#), [install_taxonkit\(\)](#), [name_or_id2df\(\)](#), [taxonkit_filter\(\)](#), [taxonkit_lca\(\)](#), [taxonkit_lineage\(\)](#), [taxonkit_name2taxid\(\)](#), [taxonkit_reformat\(\)](#)

Examples

```
## Not run:
taxonkit_list(ids = c(9605), indent = "-", show_name = TRUE, show_rank = TRUE)

## End(Not run)
```

taxonkit_name2taxid	<i>Convert Taxonomic Names to TaxIDs</i>
---------------------	--

Description

This function uses the "taxonkit taxonkit_name2taxid" command to convert taxonomic names to corresponding taxonomic IDs (TaxIDs).

Usage

```
taxonkit_name2taxid(
  file_path,
  name_field = NULL,
  sci_name = FALSE,
  show_rank = FALSE,
  text = FALSE,
  data_dir = NULL
)
```

Arguments

file_path	The path to the input file containing taxonomic names. Or file text (text=TRUE)
name_field	The field index of the taxonomic name in the input file (default is 1).
sci_name	Logical value indicating whether to search only for scientific names (default is FALSE).
show_rank	Logical value indicating whether to show the taxonomic rank in the output (default is FALSE).
text	Logical
data_dir	directory containing nodes.dmp and names.dmp (default "/Users/asa/.taxonkit")

Value

A character vector containing the output of the "taxonkit_name2taxid" command.

See Also

Other Rtaxonkit: [check_taxonkit\(\)](#), [download_taxonkit_dataset\(\)](#), [install_taxonkit\(\)](#), [name_or_id2df\(\)](#), [taxonkit_filter\(\)](#), [taxonkit_lca\(\)](#), [taxonkit_lineage\(\)](#), [taxonkit_list\(\)](#), [taxonkit_reformat\(\)](#)

Examples

```
## Not run:
names <- system.file("extdata/name.txt", package = "pctax")
taxonkit_name2taxid(names, name_field = 1, sci_name = FALSE, show_rank = FALSE)
"Homo sapiens" %>% taxonkit_name2taxid(text = TRUE)

## End(Not run)
```

taxonkit_reformat	<i>Reformat Taxonomic Lineage using taxonkit</i>
-------------------	--

Description

Reformat Taxonomic Lineage using taxonkit

Usage

```
taxonkit_reformat(
  file_path,
  delimiter = NULL,
  add_prefix = FALSE,
  prefix_kingdom = "K__",
  prefix_phylum = "p__",
  prefix_class = "c__",
  prefix_order = "o__",
  prefix_family = "f__",
  prefix_genus = "g__",
  prefix_species = "s__",
  prefix_subspecies = "t__",
  prefix_strain = "T__",
  fill_miss_rank = FALSE,
  format_string = "",
  miss_rank_repl_prefix = "unclassified ",
  miss_rank_repl = "",
  miss_taxid_repl = "",
  output_ambiguous_result = FALSE,
  lineage_field = 2,
```

```

    taxid_field = NULL,
    pseudo_strain = FALSE,
    trim = FALSE,
    text = FALSE,
    data_dir = NULL
)

```

Arguments

<code>file_path</code>	The path to the input file with taxonomic lineages. Or file text (text=TRUE)
<code>delimiter</code>	The field delimiter in the input lineage (default ";").
<code>add_prefix</code>	Logical, indicating whether to add prefixes for all ranks (default: FALSE).
<code>prefix_kingdom</code>	The prefix for kingdom, used along with <code>–add-prefix</code> (default: "K__").
<code>prefix_phylum</code>	The prefix for phylum, used along with <code>–add-prefix</code> (default: "p__").
<code>prefix_class</code>	The prefix for class, used along with <code>–add-prefix</code> (default: "c__").
<code>prefix_order</code>	The prefix for order, used along with <code>–add-prefix</code> (default: "o__").
<code>prefix_family</code>	The prefix for family, used along with <code>–add-prefix</code> (default: "f__").
<code>prefix_genus</code>	The prefix for genus, used along with <code>–add-prefix</code> (default: "g__").
<code>prefix_species</code>	The prefix for species, used along with <code>–add-prefix</code> (default: "s__").
<code>prefix_subspecies</code>	The prefix for subspecies, used along with <code>–add-prefix</code> (default: "t__").
<code>prefix_strain</code>	The prefix for strain, used along with <code>–add-prefix</code> (default: "T__").
<code>fill_miss_rank</code>	Logical, indicating whether to fill missing rank with lineage information of the next higher rank (default: FALSE).
<code>format_string</code>	The output format string with placeholders for each rank.
<code>miss_rank_repl_prefix</code>	The prefix for estimated taxon level for missing rank (default: "unclassified").
<code>miss_rank_repl</code>	The replacement string for missing rank.
<code>miss_taxid_repl</code>	The replacement string for missing taxid.
<code>output_ambiguous_result</code>	Logical, indicating whether to output one of the ambiguous result (default: FALSE).
<code>lineage_field</code>	The field index of lineage. Input data should be tab-separated (default: 2).
<code>taxid_field</code>	The field index of taxid. Input data should be tab-separated. It overrides <code>–i/–lineage-field</code> .
<code>pseudo_strain</code>	Logical, indicating whether to use the node with lowest rank as strain name (default: FALSE).
<code>trim</code>	Logical, indicating whether to not fill missing rank lower than current rank (default: FALSE).
<code>text</code>	logical
<code>data_dir</code>	directory containing nodes.dmp and names.dmp (default "/Users/asa/.taxonkit")

Value

A character vector containing the reformatted taxonomic lineages.

See Also

Other Rtaxonkit: [check_taxonkit\(\)](#), [download_taxonkit_dataset\(\)](#), [install_taxonkit\(\)](#), [name_or_id2df\(\)](#), [taxonkit_filter\(\)](#), [taxonkit_lca\(\)](#), [taxonkit_lineage\(\)](#), [taxonkit_list\(\)](#), [taxonkit_name2taxid\(\)](#)

Examples

```
## Not run:
# Use taxid
taxids2 <- system.file("extdata/taxids2.txt", package = "pctax")
reformatted_lineages <- taxonkit_reformat(taxids2,
  add_prefix = TRUE, taxid_field = 1, fill_miss_rank = TRUE
)
reformatted_lineages
taxonomy <- strsplit2(reformatted_lineages, "\t")
taxonomy <- strsplit2(taxonomy$V2, ";")

# Use lineage result
taxonkit_lineage("9606\n63221", show_name = TRUE, show_rank = TRUE, text = TRUE) %>%
  taxonkit_reformat(text = TRUE)

## End(Not run)
```

tax_lca	<i>Calculate the lowest common ancestor (LCA) of a set of taxa</i>
---------	--

Description

Calculate the lowest common ancestor (LCA) of a set of taxa

Usage

```
tax_lca(df)
```

Arguments

df	a data frame with taxonomic information, with columns representing taxonomic levels
----	---

Value

character

Examples

```
df <- data.frame(
  A = c("a", "a", "a", "a"),
  B = c("x", "x", "y", "y"),
  C = c("1", "1", "2", "3"),
  stringsAsFactors = FALSE
)
tax_lca(df)
```

time_by_cm

*Time series analysis***Description**

Time series analysis

Usage

```
time_by_cm(otu_time, n_cluster = 6, min.std = 0)
```

Arguments

otu_time	otutab hebing by a time variable
n_cluster	number of clusters
min.std	min.std

Value

time_cm

Examples

```
if (interactive()) {
  data(otutab, package = "pcutils")
  otu_time <- pcutils::hebing(otutab, metadata$Group)
  time_by_cm(otu_time, n_cluster = 4) -> time_cm_res
  plot(time_cm_res)
}
```

volcano_p	<i>Volcano plot for difference analysis</i>
-----------	---

Description

Volcano plot for difference analysis

Usage

```
volcano_p(  
  res,  
  logfc = 1,  
  adjp = 0.05,  
  text = TRUE,  
  repel = TRUE,  
  mode = 1,  
  number = FALSE  
)
```

Arguments

res	result of diff_da which have colnames: tax, log2FoldChange, padj, compare, sig
logfc	log_fold_change threshold
adjp	adjust_p_value threshold
text	text, TRUE
repel	repel, TRUE
mode	1:normal; 2:multi_contrast
number	show the tax number

Value

ggplot

See Also

[diff_da](#)

z_diversity

*Calculate Zeta Diversity***Description**

This function calculates Zeta diversity for each group in the provided otutab.

This function plots the Zeta diversity results obtained from the z_diversity function.

Usage

```
z_diversity(otutab, group_df = NULL, zetadiv_params = list())

## S3 method for class 'zeta_res'
plot(x, lm_model = c("exp", "pl")[1], ribbon = FALSE, text = TRUE, ...)
```

Arguments

otutab	A matrix or data frame containing OTU (Operational Taxonomic Unit) counts.
group_df	A data frame containing group information.
zetadiv_params	Additional parameters to be passed to the Zeta.decline.mc function from the zetadiv package.
x	Zeta diversity results obtained from z_diversity function.
lm_model	The linear model to be used for fitting ('exp' or 'pl').
ribbon	Logical, whether to add a ribbon to the plot for standard deviation.
text	Logical, whether to add R-squared and p-value text annotations.
...	Additional arguments to be passed to ggplot2 functions.

Value

zeta_res
A ggplot object.

Examples

```
if (requireNamespace("zetadiv")) {
  data(otutab, package = "pcutils")
  zeta_result <- z_diversity(otutab, metadata["Group"], zetadiv_params = list(sam = 10))
  plot(zeta_result, lm_model = "exp", text = TRUE)
}
```

z_diversity_decay	<i>Calculate Zeta Diversity with Distance</i>
-------------------	---

Description

This function calculates Zeta diversity for each group in the provided otutab.

Usage

```
z_diversity_decay(otutab, xy_df, group_df = NULL, zetadiv_params = list())

## S3 method for class 'zeta_decay'
plot(x, ribbon = TRUE, ...)
```

Arguments

otutab	A matrix or data frame containing OTU (Operational Taxonomic Unit) counts.
xy_df	Site coordinates.
group_df	A data frame containing group information.
zetadiv_params	Additional parameters to be passed to the Zeta.ddecay function from the zetadiv package.
x	Zeta diversity results obtained from z_diversity_decay function.
ribbon	Logical, whether to add a ribbon to the plot for standard deviation.
...	Additional arguments to be passed to ggplot2 functions.

Value

zeta_decay
A ggplot object.

Examples

```
if (requireNamespace("zetadiv")) {
  data(otutab, package = "pcutils")
  zeta_decay_result <- z_diversity_decay(otutab, metadata[, c("lat", "long")],
    metadata["Group"],
    zetadiv_params = list(sam = 10)
  )
  plot(zeta_decay_result)
}
```

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