Package ‘MicrobiotaProcess’

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Type Package

Title A comprehensive R package for managing and analyzing microbiome and other ecological data within the tidy framework

Version 1.14.1

Description MicrobiotaProcess is an R package for analysis, visualization and biomarker discovery of microbial datasets. It introduces MPSE class, this make it more interoperable with the existing computing ecosystem. Moreover, it introduces a tidy microbiome data structure paradigm and analysis grammar. It provides a wide variety of microbiome data analysis procedures under the unified and common framework (tidy-like framework).

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alphasample-class

Description
alphasample class

Slots
alpha  data.frame contained alpha metrics of samples
sampleda associated sample information

as.MPSE

as.MPSE method

Description
convert the .data object to MPSE object

Usage
as.MPSE(.data, ...)
as.mpse(.data, ...)

Arguments
.data one type of tbl_mpse, phyloseq, biom, SummarizedExperiment or TreeSumma-
riizedExperiment class
... additional parameters, meaningless now.

Value
MPSE object

Author(s)
Shuangbin Xu
as.phyloseq

convert to phyloseq object.

Description
convert to phyloseq object.

Usage
as.phyloseq(x, .abundance, ...)
as.phyloseq(x, .abundance, ...)

## S3 method for class 'MPSE'
as.phyloseq(x, .abundance, ...)

## S3 method for class 'tbl_mpse'
as.phyloseq(x, .abundance, ...)

Arguments
x object, tbl_mpse object, which the result of as_tibble for phyloseq object.
.abundance the column name to be as the abundance of otu table, default is Abundance.
... additional params

Value
phyloseq object.

as.treedata.taxonomyTable

Description
convert taxonomyTable to treedata

Usage
## S3 method for class 'taxonomyTable'
as.treedata(tree, include.rownames = FALSE, ...)

as.treedata
build_tree

Arguments
- tree: object, This is for taxonomyTable class, so it should be a taxonomyTable.
- include.rownames: logical, whether to set the rownames of taxonomyTable to tip labels, default is FALSE.
- ...: additional parameters.

Examples
```r
## Not run:
data(test_otu_data)
test_otu_data %<>% as.phyloseq()
tree <- as.treedata(phyloseq::tax_table(test_otu_data), include.rownames = TRUE)
## End(Not run)
```

Description
The function can be used to building tree.

Usage
```r
build_tree(seqs, ...)
```

Arguments
- seqs: DNAStringSet or DNAbin, the object of R.
- ...: additional parameters, see also AlignSeqs.

Value
- the phylo class of tree.

Author(s)
Shuangbin Xu
Examples

```r
## Not run:
seqtabfile <- system.file("extdata", "seqtab.nochim.rds", package="MicrobiotaProcess")
seqtab <- readRDS(seqtabfile)
refseq <- colnames(seqtab)
names(refseq) <- paste0("OTU_", seq_len(length(refseq)))
refseq <- Biostrings::DNAStringSet(refseq)
tree <- build_tree(refseq)
or

## End(Not run)
```

---

**convert_to_treedata**  
*convert dataframe contained hierarchical relationship or other classes to treedata class*

**Description**

convert dataframe contained hierarchical relationship or other classes to treedata class

**Usage**

`convert_to_treedata(data, type = "species", include.rownames = FALSE, ...)`

**Arguments**

- **data**  
data.frame, such like the tax_table of phyloseq.
- **type**  
character, the type of datasets, default is "species", if the dataset is not about species, # such as dataset of kegg function, you should set it to "others".
- **include.rownames**  
logical, whether to set the row names as the tip labels, default is FALSE.
- **...**  
additional parameters.

**Value**

treedata class.

**Author(s)**

Shuangbin Xu
data-hmp_aerobiosis_small

Examples

```r
## Not run:
data(hmp_aerobiosis_small)
head(taxda)
treedat <- convert_to_treedata(taxda, include.rownames = FALSE)
## End(Not run)
```

Data (Small subset of the HMP 16S dataset)

Description

Contained three datasets, featureda, sampleda, taxda. featureda contained 55 samples (nrow) and 1091 features (ncol). sampleda contained 55 samples from 6 body sites of 10 subjects. taxda contained 699 taxonomy by 6 rank. This datasets were built from the LEfSe.

http://huttenhower.sph.harvard.edu/webfm_send/129

Examples

```r
data(hmp_aerobiosis_small)
```

data-kostic2012crc

(Data) Genomic analysis identifies association of Fusobacterium with colorectal carcinoma (2012)

Description

This dataset was from a study on colorectal cancer, published in Genome Research (2012). This dataset had been removed samples with less than 500 reads, contained 91 Control and 86 Tumors. And it belongs to MPSE class, contained otu_table and sample_data.

Examples

```r
data(kostic2012crc)
```

data-test_otu_data

(Data) simulated dataset.

Description

This dataset was simulated. And it also was MPSE class, contained otu_table and sample_data.

Examples

```r
data(test_otu_data)
```
diffAnalysisClass-class

diffAnalysisClass class

Description

diffAnalysisClass class

Slots

originalD original feature data.frame.
sampledA associated sample information.
taxA the data.frame contained taxonomy.
result data.frame contained the results of first, second test and LDA or rf
kwres the results of first test, contained feature names, pvalue and fdr.
secondvars the results of second test, contained features names, gfc (TRUE representation the relevant feantures is enriched in relevant factorNames), Freq(the number of TRUE or FALSE), factorNames.
mlres the results of LDA or randomForest,
someparams, some arguments will be used in other functions diff_analysis

diff_analysis

Differential expression analysis

Description

Differential expression analysis

Usage

diff_analysis(obj, ...)

## S3 method for class 'data.frame'
diff_analysis(
  obj,
sampledA,
classgroup,
subclass = NULL,
taxA = NULL,
alltax = TRUE,
include.rownames = FALSE,
standard_method = NULL,
mlfun = "lda",


ratio = 0.7,
firstcomfun = "kruskal.test",
padjust = "fdr",
filtermod = "pvalue",
firstalpha = 0.05,
strictmod = TRUE,
fcfun = "generalizedFC",
secondcomfun = "wilcox.test",
clmin = 5,
clwilc = TRUE,
secondalpha = 0.05,
subclmin = 3,
subclwilc = TRUE,
ldascore = 2,
normalization = 1e+06,
bootnums = 30,
cl = 0.95,
type = "species",
...
)

## S3 method for class 'phyloseq'
diff_analysis(obj, ...)

Arguments

- **obj**: object, a phyloseq class contained otu_table, sample_data, taxda, or data.frame, nrow sample * ncol features.
- **...**: additional parameters.
- **sampleda**: data.frame, nrow sample * ncol factor, the sample names of sampleda and data should be the same.
- **classgroup**: character, the factor name in sampledata.
- **subclass**: character, the factor name in sampledata, default is NULL, meaning no subclass compare.
- **taxda**: data.frame, the classification of the feature in data. default is NULL.
- **alltax**: logical, whether to set all classification (taxonomy) as features when taxda is not NULL, default is TRUE.
- **include.rownames**: logical, whether to consider the OTU of obj as (all taxonomy) features, when taxda is not NULL, default is FALSE.
- **standard_method**: character, the method of standardization, see also `decostand`, default is NULL, it represents that the relative abundance of taxonomy will be used. If count was set, it represents the count reads of taxonomy will be used.
- **mlfun**: character, the method for calculating the effect size of features, choose "lda" or "rf", default is "lda".
ratio numeric, range from 0 to 1, the proportion of samples for calculating the effect size of features, default is 0.7.

firstcomfun character, the method for first test, "oneway.test" for normal distributions, suggested choosing "kruskal.test" for uneven distributions, default is "kruskal.test", or you can use lm, glm, or glm.nb (for negative binomial distribution), or 'kruskal_test', 'oneway_test' of 'coin'.

padjust character, the correction method, default is "fdr".

filtermod character, the method to filter, default is "pvalue".

firstalpha numeric, the alpha value for the first test, default is 0.05.

strictmod logical, whether to performed in one-against-one, default is TRUE (strict).

fcfun character, default is "generalizedFC", it can't be set another at the present time.

secondcomfun character, the method for one-against-one, default is "wilcox.test" for uneven distributions, or 'wilcox_test' of 'coin', or you can also use 'lm', 'glm', 'glm.nb'(for negative binomial distribution in 'MASS').

cmin integer, the minimum number of samples per classgroup for performing test, default is 5.

cwilc logical, whether to perform test of per classgroup, default is TRUE.

secondalpha numeric, the alpha value for the second test, default is 0.05.

subclmin integer, the minimum number of samples per subclass for performing test, default is 3.

subclwilc logical, whether to perform test of per subclass, default is TRUE, meaning more strict.

ldascore numeric, the threshold on the absolute value of the logarithmic LDA score, default is 2.

normalization integer, set the normalization value, set a big number if to get more meaningful values for the LDA score, or you can set NULL for no normalization, default is 1000000.

bootnums integer, set the number of bootstrap iteration for lda or rf, default is 30.

.ci numeric, the confidence interval of effect size (LDA or MDA), default is 0.95.

type character, the type of datasets, default is "species", if the dataset is not about species, such as dataset of kegg function, you should set it to "others".

Value
diff_analysis class.

Author(s)
Shuangbin Xu
Examples

```r
## Not run:
data(kostic2012crc)
kostic2012crc %<>% as.phyloseq()
head(phyloseq::sample_data(kostic2012crc),3)
kostic2012crc <- phyloseq::rarefy_even_depth(kostic2012crc,rngseed=1024)
table(phyloseq::sample_data(kostic2012crc)$DIAGNOSIS)
set.seed(1024)
diffres <- diff_analysis(kostic2012crc, classgroup="DIAGNOSIS",
mlfun="lda", filtermod="fdr",
firstcomfun = "kruskal.test",
firstalpha=0.05, strictmod=TRUE,
secondcomfun = "wilcox.test",
subclmin=3, subclwilc=TRUE,
secondalpha=0.01, ldascore=3)

## End(Not run)
```

### drop_taxa

Dropping Species with Few abundance and Few Occurrences

**Description**

Drop species or features from the feature data frame or phyloseq that occur fewer than or equal to a threshold number of occurrences and fewer abundance than to a threshold abundance.

**Usage**

```r
drop_taxa(obj, ...)
```

**Arguments**

- `obj` object, phyloseq or a dataframe of species (n_sample, n_feature).
- `...` additional parameters.
- `minocc` numeric, the threshold number of occurrences to be dropped, if < 1.0, it will be the threshold ratios of occurrences, default is 0.
- `minabu` numeric, the threshold abundance, if fewer than the threshold will be dropped, default is 0.

**Value**

dataframe of new features.
Author(s)

Shuangbin Xu

Examples

```r
## Not run:
otudafile <- system.file("extdata", "otu_tax_table.txt", 
  package="MicrobiotaProcess")
otuda <- read.table(otudafile, sep="\t",
  header=TRUE, row.names=1,
  check.names=FALSE, skip=1,
  comment.char="")
otuda <- otuda[sapply(otuda, is.numeric)]
otuda <- data.frame(t(otuda), check.names=FALSE)
dim(otuda)

## End(Not run)
```

---

**dr_extract**

Extracting the internal tbl_df attribute of tibble.

**Description**

Extracting the internal tbl_df attribute of tibble.

**Usage**

```r
dr_extract(name, .f = NULL)
```

**Arguments**

- `name` character the name of internal tbl_df attribute.
- `.f` a function (if any, default is NULL) that pre-operate the data

**Value**

tbl_df object

**Author(s)**

Shuangbin Xu
### Examples

```r
## Not run:
library(vegan)
data(varespec, varechem)
mpse <- MPSE(assays=list(Abundance=t(varespec)), colData=varechem)
tbl <-
mpse %>%
  mp_cal_nmds(.abundance=Abundance, action="add") %>%
  mp_envfit(.ord=NMDS, .env=colnames(varechem), action="only")
tbl
tbl %>% attributes %>% names
# This function is useful to extract the data to display with ggplot2
# you can also refer to the examples of mp_envfit.
dr_extract(name=NMDS_ENVFIT_tb)(tbl)
# add .f function
dr_extract(name=NMDS_ENVFIT_tb,
  .f=td_filter(pvals<=0.05 & label!="Humdepth"))(tbl)
## End(Not run)
```

---

**extract_binary_offspring**

*extract the binary offspring of the specified internal nodes*

### Description

extract the binary offspring of the specified internal nodes

### Usage

```
extract_binary_offspring(.data, .node, type = "tips", ...)
```

### Arguments

- `.data` phyl or treedata object
- `.node` the internal nodes
- `type` the type of binary offspring, options are 'tips' (default), 'all', 'internal'.
- `...` additional parameter, meaningless now.
generalizedFC  

**Description**

calculate the mean difference in a set of predefined quantiles of the logarithmic

**Usage**

generalizedFC(x, ...)

## Default S3 method:
generalizedFC(x, y, base = 10, steps = 0.05, pseudo = 1e-05, ...)

## S3 method for class 'formula'
generalizedFC(x, data, subset, na.action, ...)

**Arguments**

- `x` numeric vector, numeric vector of data values or formula, example `Ozone ~ Month`. Ozone is a numeric variable giving the data values ‘Month’ a factor giving the corresponding groups.
- `...` additional arguments.
- `y` numeric vector, numeric vector of data values
- `base` a positive or complex number, the base with respect to which logarithms are computed, default is 10.
- `steps` positive numeric, increment of the sequence, default is 0.05.
- `pseudo` positive numeric, avoid the zero for logarithmic, default is 0.00001.
- `data` data.frame, an optional matrix or data frame, containing the variables in the formula.
- `subset` (similar: see `wilcox.test`) an optional vector specifying a subset of observations to be used.
- `na.action` a function which indicates what should happen when the data, contain ’NA’s. Defaults to ’getOption("na.action")’.

**Value**

list contained gfc, the mean and median of different group.

**Author(s)**

Shuangbin Xu
get_alltaxadf

Examples

```r
set.seed(1024)
data <- data.frame(A=rnorm(1:10,mean=5),
                   B=rnorm(2:11, mean=6),
                   group=c(rep("case",5),rep("control",5)))
generalizedFC(B ~ group,data=data)
generalizedFC(x=c(1,2,3,4,5),y=c(3,4,5,6,7))
```

get_alltaxadf  get the table of abundance of all level taxonomy

Description

This function was designed to get the abundance of all level taxonomy, the input can be phyloseq object or data.frame.

Usage

```
generalizedFC(x, y, ...,
              lambda = NULL,
              method = NULL,
              type = "species",
              include.rownames = FALSE,
              ...)  
```

Arguments

- `obj` object, phyloseq or data.frame
- `...` additional parameters, see also `decostand`
- `method` character, the normalization method, see also `decostand`, default is NULL, the relative abundance will be return, if it set 'count', the count table will be return.
include.rownames

  logical whether to calculate the original feature data, default is FALSE.

taxda
data.frame, the taxonomy table.
taxa_are_rows

  logical, if the obj is data.frame, and the features are rownames, the taxa_are_rows
  should be set TRUE, default FALSE, meaning the features are colnames.

Value

  the all taxonomy abundance table

Author(s)

  Shuangbin Xu

Examples

  ## Not run:
  data(test_otu_data)
  alltaxatab <- get_alltaxadf(test_otu_data)
  head(alltaxatab[,1:10])

  ## End(Not run)

get_alphaindex

alpha index

calculate the alpha index (Obseve, Chao1, Shannon, Simpson) of sample with diversity

Usage

  get_alphaindex(obj, ...)

  ## S4 method for signature 'matrix'
  get_alphaindex(obj, mindepth, sampleda, force = FALSE, ...)

  ## S4 method for signature 'data.frame'
  get_alphaindex(obj, ...)

  ## S4 method for signature 'integer'
  get_alphaindex(obj, ...)

  ## S4 method for signature 'numeric'
  get_alphaindex(obj, ...)

  ## S4 method for signature 'phyloseq'
  get_alphaindex(obj, ...)

---

get_alphaindex

alpha index

calculate the alpha index (Obseve, Chao1, Shannon, Simpson) of sample with diversity

Usage

  get_alphaindex(obj, ...)

  ## S4 method for signature 'matrix'
  get_alphaindex(obj, mindepth, sampleda, force = FALSE, ...)

  ## S4 method for signature 'data.frame'
  get_alphaindex(obj, ...)

  ## S4 method for signature 'integer'
  get_alphaindex(obj, ...)

  ## S4 method for signature 'numeric'
  get_alphaindex(obj, ...)

  ## S4 method for signature 'phyloseq'
  get_alphaindex(obj, ...)

---
get_clust

Hierarchical cluster analysis for the samples

Description

Hierarchical cluster analysis for the samples

Arguments

- **obj**: object, data.frame of (nrow sample * ncol taxonomy(feature)) or phyloseq.
- **...**: additional arguments.
- **mindepth**: numeric, Subsample size for rarefying community.
- **sampledata**: data.frame, sample information, row sample * column factors.
- **force**: logical whether calculate the alpha index even the count of otu is not rarefied, default is FALSE. If it is TRUE, meaning the rarefaction is not be performed automatically.

Value

data.frame contained alpha Index.

Author(s)

Shuangbin Xu

Examples

```r
## Not run:
otudafile <- system.file("extdata", "otu_tax_table.txt", package="MicrobiotaProcess")
otuda <- read.table(otudafile, sep="\t", header=TRUE, row.names=1, check.names=FALSE, skip=1, comment.char="")
set.seed(1024)
alphatab <- get_alphaindex(otuda)
head(as.data.frame(alphatab))
data(test_otu_data)
class(test_otu_data)
test_otu_data %<>% as.phyloseq()
class(test_otu_data)
set.seed(1024)
alphatab2 <- get_alphaindex(test_otu_data)
head(as.data.frame(alphatab2))
## End(Not run)
```
Usage

get_clust(obj, ...)

## S3 method for class 'dist'
get_clust(obj, distmethod, sampleda = NULL, hclustmethod = "average", ...)

## S3 method for class 'data.frame'
get_clust(
  obj,
  distmethod = "euclidean",
  taxa_are_rows = FALSE,
  sampleda = NULL,
  tree = NULL,
  method = "hellinger",
  hclustmethod = "average",
  ...
)

## S3 method for class 'phyloseq'
get_clust(
  obj,
  distmethod = "euclidean",
  method = "hellinger",
  hclustmethod = "average",
  ...
)

Arguments

obj            phyloseq, phyloseq class or dist class, or data.frame, data.frame, default is nrow 
samples * ncol features.
...             additional parameters.
distmethod     character, the method of dist, when the obj is data.frame or phyloseq default is
                "euclidean". see also get_dist.
sampled        data.frame, nrow sample * ncol factor. default is NULL.
hclustmethod   character, the method of hierarchical cluster, default is average.
taxa_are_rows   logical, if the features of data.frame(obj) is in column, it should set FALSE.
tree            phylo, the phylo class, see also as.phylo.
method          character, the standardization methods for community ecologists, see also decostand

Value

treedata object.

Author(s)

Shuangbin Xu
Examples

```r
## Not run:
library(phyloseq)
data(GlobalPatterns)
subGlobal <- subset_samples(GlobalPatterns,
    SampleType %in% c("Feces", "Mock", "Ocean", "Skin"))
hcsample <- get_clust(subGlobal, distmethod="jaccard",
    method="hellinger", hclustmethod="average")

## End(Not run)
```

get ordination coordinates.

Description

get ordination coordinates.

Usage

```r
## S3 method for class 'pcoa'
get_coord(obj, pc)
get_coord(obj, pc)

## S3 method for class 'prcomp'
get_coord(obj, pc)
```

Arguments

- `obj`: object, prcomp class or pcoa class
- `pc`: integer vector, the component index.

Value

ordplotClass object.

Examples

```r
## Not run:
require(graphics)
data(USArrests)
pcares <- prcomp(USArrests, scale = TRUE)
coordtab <- get_coord(pcares, pc=c(1, 2))
coordtab2 <- get_coord(pcares, pc=c(2, 3))

## End(Not run)
```
get_count

calculate the count or relative abundance of replicate element with a specific column

Description

Calculate the count or relative abundance of replicate element with a specific columns.

Usage

get_count(data, featurelist, ...)

get_ratio(data, featurelist, ...)

Arguments

data: dataframe; a dataframe contained one character column and others is numeric, if featurelist is NULL. Or a numeric dataframe, if featurelist is not NULL, all columns should be numeric.

featurelist: dataframe; a dataframe contained one character column, default is NULL.

...: additional parameters.

Value

mean of data.frame by featurelist

Author(s)

Shuangbin Xu

Examples

## Not run:
otudafile <- system.file("extdata", "otu_tax_table.txt", package="MicrobiotaProcess")
samplefile <- system.file("extdata", "sample_info.txt", package="MicrobiotaProcess")

otuda <- read.table(otudafile, sep="\t", header=TRUE, row.names=1, check.names=FALSE, skip=1, comment.char="")
sampled <- read.table(samplefile, sep="\t", header=TRUE, row.names=1)
taxdf <- otuda[!sapply(otuda, is.numeric)]
taxdf <- split.str.to.list(taxdf)

phycount <- get_count(otuda, taxdf[,2,drop=FALSE])
phyratios <- get_ratio(otuda, taxdf[,2,drop=FALSE])

## End(Not run)
get_dist

calculate distance

Description
calculate distance

Usage
get_dist(obj, ...)

## S3 method for class 'data.frame'
get_dist(
  obj,
  distmethod = "euclidean",
  taxa_are_rows = FALSE,
  sampled = NULL,
  tree = NULL,
  method = "hellinger",
  ...
)

## S3 method for class 'phyloseq'
get_dist(obj, distmethod = "euclidean", method = "hellinger", ...)

Arguments

obj phyloseq, phyloseq class or data.frame nrow sample * ncol feature.

..., additional parameters.
distmethod character, default is "euclidean", see also distanceMethodList
taxa_are_rows logical, default is FALSE.
sampled data.frame, nrow sample * ncol factors.
tree object, the phylo class, see also as.phylo.
method character, default is hellinger, see also decostand

Value
distance class contained distmethod and originalD attr

See Also
distance
get_mean_median

get the mean and median of specific feature.

## Description
get the mean and median of specific feature.

## Usage
get_mean_median(datameta, feature, subclass)

## Arguments
- `datameta`: data.frame, nrow sample * ncol feature + factor.
- `feature`: character vector, the feature contained in datameta.
- `subclass`: character, factor name.

## Value
featureMeanMedian object, contained the abundance of feature, and the mean and median of feature by subclass.

## Author(s)
Shuangbin Xu

## Examples
```r
## Not run:
data(hmp_aerobiosis_small)
head(sampleda)
featureda <- merge(featureda, sampleda, by=0)
rownames(featureda) <- as.vector(featureda$Row.names)
featureda$Row.names <- NULL
feameamed <- get_mean_median(datameta=featureda,
    feature="p__Actinobacteria",
    subclass="body_site")
fplot <- ggdifftaxbar(feameamed, featurename="p__Actinobacteria",
    classgroup="oxygen_availability", subclass="body_site")
## End(Not run)
```
get_NRI_NTI calculating related phylogenetic alpha metric

Description

calculating related phylogenetic alpha metric

Usage

get_NRI_NTI(obj, ...)

## S4 method for signature 'matrix'
get_NRI_NTI(
  obj,
  mindepth,
  sampleda,
  tree,
  metric = c("PAE", "NRI", "NTI", "PD", "HAED", "EAED", "IAC", "all"),
  abundance.weighted = FALSE,
  force = FALSE,
  seed = 123,
  ...
)

## S4 method for signature 'data.frame'
get_NRI_NTI(obj, mindepth, sampleda, tree, abundance.weighted = TRUE, ...)

## S4 method for signature 'phyloseq'
get_NRI_NTI(obj, mindepth, abundance.weighted = TRUE, ...)

Arguments

<table>
<thead>
<tr>
<th>Argument</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>obj</td>
<td>object, data.frame of (nrow sample * ncol taxonomy(feature)) or phyloseq.</td>
</tr>
<tr>
<td>...</td>
<td>additional arguments, meaningless now.</td>
</tr>
<tr>
<td>mindepth</td>
<td>numeric, Subsample size for rarefying community.</td>
</tr>
<tr>
<td>sampleda</td>
<td>data.frame, sample information, row sample * column factors.</td>
</tr>
<tr>
<td>tree</td>
<td>tree object, it can be phylo object or treedata object.</td>
</tr>
<tr>
<td>metric</td>
<td>the related phylogenetic metric, options is 'NRI', 'NTI', 'PD', 'PAE', 'HAED',</td>
</tr>
<tr>
<td></td>
<td>'EAED', 'IAC', 'all', default is 'PAE', meaning all the metrics ('NRI', 'NTI',</td>
</tr>
<tr>
<td></td>
<td>'PD', 'PAE', 'HAED', 'EAED', 'IAC').</td>
</tr>
<tr>
<td>abundance.weighted</td>
<td>logical, whether calculate mean nearest taxon distances for each species weighted</td>
</tr>
<tr>
<td>force</td>
<td>logical whether calculate the index even the count of otu is not rarefied, default</td>
</tr>
<tr>
<td></td>
<td>is FALSE. If it is TRUE, meaning the rarefaction is not be performed automatically.</td>
</tr>
</tbody>
</table>
get_pca

seed integer a random seed to make the result reproducible, default is 123.

Value

alphasample object contained NRT and NTI.

Author(s)

Shuangbin Xu

geta principal components analysis

Description

Performs a principal components analysis

Usage

geta(obj, ...)

## S3 method for class 'data.frame'
geta(obj, sampleda = NULL, method = "hellinger", ...)

## S3 method for class 'phyloseq'
geta(obj, method = "hellinger", ...)

Arguments

obj phyloseq, phyloseq class or data.frame shape of data.frame is nrow sample * ncol feature.

... additional parameters, seeprcomp.

sampleda data.frame, nrow sample * ncol factors.

method character, the standardization methods for community ecologists. see decostand.

Value

pcasample class, contained prcomp class and sample information.

Examples

## Not run:
library(phyloseq)
data(GlobalPatterns)
subGlobal <- subset_samples(GlobalPatterns,
    SampleType %in% c("Feces", "Mock", "Ocean", "Skin"))
pcares <- get_pca(subGlobal, method="hellinger")
pcaplot <- ggordpoint(pcares, biplot=TRUE,
get_pcoa

speciesannot=TRUE,
factorNames=c("SampleType"), ellipse=TRUE)

## End(Not run)

get_pcoa performs principal coordinate analysis (PCoA)

Description
performs principal coordinate analysis (PCoA)

Usage
get_pcoa(obj, ...)

## S3 method for class 'data.frame'
get_pcoa(
  obj,
  distmethod = "euclidean",
  taxa_are_rows = FALSE,
  sampleda = NULL,
  tree = NULL,
  method = "hellinger",
  ...
)

## S3 method for class 'dist'
get_pcoa(
  obj,
  distmethod,
  data = NULL,
  sampleda = NULL,
  method = "hellinger",
  ...
)

## S3 method for class 'phyloseq'
get_pcoa(obj, distmethod = "euclidean", ...)

Arguments

obj phyloseq, the phyloseq class or dist class.
..., additional parameter, see also get_dist.
distmethod character, the method of distance, see also distance
taxa_are_rows logical, if feature of data is column, it should be set FALSE.
get_pvalue

Methods for computation of the p-value

Description

Methods for computation of the p-value

Usage

get_pvalue(obj)

## S3 method for class 'htest'
get_pvalue(obj)

## S3 method for class 'lme'
get_pvalue(obj)
## S3 method for class 'negbin'
get_pvalue(obj)

## S3 method for class 'ScalarIndependenceTest'
get_pvalue(obj)

## S3 method for class 'QuadTypeIndependenceTest'
get_pvalue(obj)

## S3 method for class 'lm'
get_pvalue(obj)

## S3 method for class 'glm'
get_pvalue(obj)

### Arguments

**obj**
object, such as htest, lm, negbin ScalarIndependenceTest class.

### Value

pvalue.

### Author(s)

Shuangbin Xu

### Examples

```r
library(nlme)
lmeres <- lme(distance ~ Sex, data=Orthodont)
pvalue <- get_pvalue(lmeres)
```

---

### Description

generate the result of rare curve.

### Usage

```r
get_rarecurve(obj, ...)
```

```r
## S4 method for signature 'data.frame'
get_rarecurve(obj, sampleda, factorLevels = NULL, chunks = 400)
```

```r
## S4 method for signature 'phyloseq'
get_rarecurve(obj, ...)
```
get_sampledflist

**Arguments**

- **obj** phyloseq class or data.frame shape of data.frame (nrow sample * ncol feature)
- **...** additional parameters.
- **sampleda** data.frame, (nrow sample * ncol factor)
- **factorLevels** list, the levels of the factors, default is NULL, if you want to order the levels of factor, you can set this.
- **chunks** integer, the number of subsample in a sample, default is 400.

**Details**

This function is designed to calculate the rare curve result of otu table the result can be visualized by 'ggrarecurve'.

**Value**

rarecurve class, which can be visualized by ggrarecurve

**Author(s)**

Shuangbin Xu

**Examples**

```r
## Not run:
data(test_otu_data)
test_otu_data %<>% as.phyloseq()
set.seed(1024)
res <- get_rarecurve(test_otu_data, chunks=200)
p <- ggrarecurve(obj=res,
                 indexNames=c("Observe","Chao1","ACE"),
                 shadow=FALSE,
                 factorNames="group")
## End(Not run)
```

---

**get_sampledflist** *Generate random data list from a original data.*

**Description**

Generate random data list from a original data.

**Usage**

```r
get_sampledflist(dalist, bootnums = 30, ratio = 0.7, makerownames = FALSE)
```
Arguments

dalist: list, a list contained multi data.frame.
bootnums: integer, the number of bootstrap iteration, default is 30.
ratio: numeric, the ratios of each data.frame to keep.
makerownames: logical, whether build row.names, default is FALSE.

Value

the list contained the data.frame generated by bootstrap iteration.

Author(s)

Shuangbin Xu

Examples

## Not run:
data(iris)
irislist <- split(iris, iris$Species)
set.seed(1024)
irislist <- get_sampledflist(irislist)
## End(Not run)

get_taxadf

get the data of specified taxonomy

Description

get the data of specified taxonomy

Usage

get_taxadf(obj, ...)

## S4 method for signature 'phyloseq'
get_taxadf(obj, taxlevel = 2, type = "species", ...)

## S4 method for signature 'data.frame'
get_taxadf(
  obj,
  taxa,
  taxa_are_rows,
  taxlevel,
  sampleda = NULL,
  type = "species",
  ...
)

---
Arguments

obj: phyloseq, phyloseq class or data.frame the shape of data.frame (nrow sample * column feature taxa_are_rows set FALSE, nrow feature * ncol sample, taxa_are_rows set TRUE).

..., additional parameters.
taxlevel: character, the column names of taxda that you want to get. when the input is phyloseq class, you can use 1 to 7.
type: character, the type of datasets, default is "species", if the dataset is not about species, such as dataset of kegg function, you should set it to "others".
taxda: data.frame, the classifies of feature contained in obj(data.frame).
taxa_are_rows: logical, if the column of data.frame are features, it should be set FALSE.
sampled: data.frame, the sample information.

Value

phyloseq class contained tax data.frame and sample information.

Author(s)

Shuangbin Xu

Examples

```r
## Not run:
library(ggplot2)
data(test_otu_data)
test_otu_data %<>% as.phyloseq()
phytax <- get_taxadf(test_otu_data, taxlevel=2)
phytax
head(phyloseq::otu_table(phytax))
phybar <- ggbartax(phytax) +
  xlab(NULL) + ylab("relative abundance (%)")
## End(Not run)
```

get_upset

generate the dataset for upset of UpSetR

Description

generate the dataset for upset of UpSetR
Usage

callupset(obj, ...)  

## S4 method for signature 'data.frame'  
callupset(obj, sampleda, factorNames, threshold = 0)  

## S4 method for signature 'phyloseq'  
callupset(obj, ...)  

Arguments

- **obj**: object, phyloseq or data.frame, if it is data.frame, the shape of it should be row sample * columns features.
- **...**: additional parameters.
- **sampleda**: data.frame, if the obj is data.frame, the sampleda should be provided.
- **factorNames**: character, the column names of factor in sampleda.
- **threshold**: integer, default is 0.

Value

a data.frame for the input of ‘upset’ of ‘UpSetR’.

Author(s)

Shuangbin Xu

Examples

```r  
## Not run:  
data(test_otu_data)  
test_otu_data %<>% as.phyloseq()  
upsetda <- get_upset(test_otu_data, factorNames="group")  
otudafile <- system.file("extdata", "otu_tax_table.txt",  
package="MicrobiotaProcess")  
samplefile <- system.file("extdata", "sample_info.txt",  
package="MicrobiotaProcess")  
otuda <- read.table(otudafile, sep="\t", header=TRUE,  
row.names=1, check.names=FALSE,  
skip=1, comment.char="")  
sampleda <- read.table(samplefile, sep="\t",  
header=TRUE, row.names=1)  
head(sampleda)  
notuda <- notuda[sapply(notuda, is.numeric)]  
otuda <- data.frame(t(notuda), check.names=FALSE)  
head(notuda[1:5, 1:5])  
upsetda2 <- get_upset(obj=notuda, sampleda=sampleda,  
factorNames="group")  
#Then you can use `upset` of `UpSetR` to visualize the results.  
library(UpSetR)  
```
get_varct.pcoa

get the contribution of variables

Description
get the contribution of variables

Usage

## S3 method for class 'pcoa'
get_varct(obj, ...)

get_varct(obj, ...)

## S3 method for class 'prcomp'
get_varct(obj, ...)

## S3 method for class 'pcasample'
get_varct(obj, ...)

Arguments

obj prcomp class or pcasample class
...
additional parameters.

Value
the VarContrib class, contained the contribution and coordinate of features.

Examples

## Not run:
library(phyloseq)
data(GlobalPatterns)
subGlobal <- subset_samples(GlobalPatterns,
    SampleType %in% c("Feces", "Mock", "Ocean", "Skin"))
pcares <- get_pca(subGlobal, method="hellinger")
varres <- get_varct(pcares)

## End(Not run)
**get_vennlist**

generate a vennlist for VennDiagram

Description

generate a vennlist for VennDiagram

Usage

```r
get_vennlist(obj, ...)
```

## S4 method for signature 'phyloseq'
```r
get_vennlist(obj, factorNames, ...)
```

## S4 method for signature 'data.frame'
```r
get_vennlist(obj, sampleinfo = NULL, factorNames = NULL, ...)
```

Arguments

- `obj` phyloseq, phyloseq class or data.frame a dataframe contained one character column and the others are numeric. or all columns should be numeric if sampleinfo isn’t NULL.
- `...` additional parameters
- `factorNames` character, a column name of sampleinfo, when sampleinfo isn’t NULL, factorNames shouldn’t be NULL, default is NULL, when the input is phyloseq, the factorNames should be provided.
- `sampleinfo` dataframe; a sample information, default is NULL.

Value

return a list for VennDiagram.

Author(s)

Shuangbin Xu

Examples

```r
## Not run:
data(test_otu_data)
test_otu_data %<>% as.phyloseq()
vennlist <- get_vennlist(test_otu_data,
                         factorNames="group")
vennlist
library(VennDiagram)
venn.diagram(vennlist, height=5,
             width=5, filename = ".:/test_venn.pdf",
```

alpha = 0.85, fontfamily = "serif",
fontface = "bold",cex = 1.2,
cat.cex = 1.2, cat.default.pos = "outer",
cat.dist = c(0.22,0.22,0.12,0.12),
margin = 0.1, lwd = 3,
lty = 'dotted',
imagetype = "pdf")

## End(Not run)
ggbartax  taxonomy barplot

Description

taxonomy barplot

Usage

ggbartax(obj, ...)
ggbartaxa(obj, ...)

## S3 method for class 'phyloseq'
ggbartax(obj, ...)

## S3 method for class 'data.frame'
ggbartax(
  obj,
  mapping = NULL,
  position = "stack",
  stat = "identity",
  width = 0.7,
  topn = 30,
  count = FALSE,
  sampled = NULL,
  factorLevels = NULL,
  sampleLevels = NULL,
  facetNames = NULL,
  plotgroup = FALSE,
  groupfun = mean,
  ...
)

Arguments

obj phylseq, phyloseq class or data.frame, (nrow sample * ncol feature (factor)) or the data.frame for geom_bar.
additional parameters, see ggplot

mapping set of aesthetic mapping of ggplot2, default is NULL, if the data is the data.frame for geom_bar, the mapping should be set.

position character, default is ‘stack’.

stat character, default is ‘identity’.

width numeric, the width of bar, default is 0.7.

topn integer, the top number of abundance taxonomy(feature).

count logical, whether show the relative abundance.

sampled data.frame, (nrow sample * ncol factor), the sample information, if the data doesn’t contain the information.

factorLevels vector or list, the levels of the factors (contained names e.g. list(group=c("B","A","C")) or c(group=c("B","A","C")), adjust the order of facet, default is NULL, if you want to order the levels of factor, you can set this.

sampleLevels vector, adjust the order of x axis e.g. c("sample2", "sample4", "sample3"), default is NULL.

facetNames character, default is NULL.

plotgroup logical, whether calculate the mean or median etc for each group, default is FALSE.

groupfun character, how to calculate for feature in each group, the default is ‘mean’, this will plot the mean of feature in each group.

Value

barplot of tax

Author(s)

Shuangbin Xu

Examples

```r
## Not run:
library(ggplot2)
data(test_otu_data)
test_otu_data %<>% as.phyloseq()
.otubar <- ggbartax(test_otu_data) +
xlab(NULL) + ylab("relative abundance(%)")
```

```r
## End(Not run)
```
ggbox  

A box or violin plot with significance test

Description

A box or violin plot with significance test

Usage

ggbox(obj, factorNames, ...)

## S4 method for signature 'data.frame'

ggbox(
  obj,
  sampleda,
  factorNames,
  indexNames,
  geom = "boxplot",
  factorLevels = NULL,
  compare = TRUE,
  testmethod = "wilcox.test",
  signifmap = FALSE,
  p_textsize = 2,
  step_increase = 0.1,
  boxwidth = 0.2,
  facetnrow = 1,
  controlgroup = NULL,
  comparelist = NULL,
  ...
)

## S4 method for signature 'alphasample'

ggbox(obj, factorNames, ...)

Arguments

obj  
object, alphasample or data.frame (row sample x column features).

factorNames  
character, the names of factor contained in sampleda.

...  
additional arguments, see also stat_signif.

sampleda  
data.frame, sample information if obj is data.frame, the sampleda should be provided.

indexNames  
character, the vector character, should be the names of features contained object.

gem  
character, "boxplot" or "violin", default is "boxplot".

factorLevels  
list, the levels of the factors, default is NULL, if you want to order the levels of factor, you can set this.
ggbox

compare logical, whether test the features among groups, default is TRUE.
testmethod character, the method of test, default is ‘wilcox.test’. see also stat_signif.
signifmap logical, whether the pvalue are directly written a annotation or asterisks are used instead, default is (pvalue) FALSE. see also stat_signif.
p_textsize numeric, the size of text of pvalue or asterisks, default is 2.
step_increase numeric, see also stat_signif, default is 0.1.
boxwidth numeric, the width of boxplot when the geom is ‘violin’, default is 0.2.
facettrown integer, the nrow of facet, default is 1.
controlgroup character, the names of control group, if it was set, the other groups will compare to it, default is NULL.
comparelist list, the list of vector, default is NULL.

Value

a 'ggplot' plot object, a box or violine plot.

Author(s)

Shuangbin Xu

Examples

```r
## Not run:
library(magrittr)
otudafile <- system.file("extdata", "otu_tax_table.txt", package="MicrobiotaProcess")

otuda <- read.table(otudafile, sep="\t",
 header=TRUE, row.names=1,
 check.names=FALSE, skip=1,
 comment.char="")
samplefile <- system.file("extdata", "sample_info.txt", package="MicrobiotaProcess")
sampleda <- read.table(samplefile, sep="\t", header=TRUE, row.names=1)

alphaobj1 <- get_alphaindex(otuda, sampleda=sampleda)
p1 <- ggbox(alphaobj1, factorNames="group")
data(test_otu_data)
test_otu_data %>% as.phyloseq()
set.seed(1024)
alphaobj2 <- get_alphaindex(test_otu_data)
class(alphaobj2)
head(as.data.frame(alphaobj2))
p2 <- ggbox(alphaobj2, factorNames="group")

# set factor levels.
p3 <- ggbox(obj=alphaobj2, factorNames="group",
```
factorLevels=list(group=c("M", "N", "B", "D")))
# set control group.
p4 <- ggbox(obj=alphaobj2, factorNames="group", controlgroup="B")
set comparelist
p5 <- ggbox(obj=alphaobj2, factorNames="group",
            comparelist=list(c("B", "D"), c("B", "M"), c("B", "N")))

## End(Not run)

ggclust

plot the result of hierarchical cluster analysis for the samples

Description

plot the result of hierarchical cluster analysis for the samples

Usage

ggclust(obj, ...)

## S3 method for class 'treedata'
ggclust(  
  obj,  
  layout = "rectangular",  
  factorNames = NULL,  
  factorLevels = NULL,  
  pointsize = 2,  
  fontsize = 2.6,  
  hjust = -0.1,  
  ...  
)

Arguments

obj R object, treedata object.
...
additional params, see also geom_tipmap
layout character, the layout of tree, see also ggtree.
factorNames character, default is NULL.
factorLevels list, default is NULL.
pointsize numeric, the size of point, default is 2.
fontsize numeric, the size of text of tiplabel, default is 2.6.
hjust numeric, default is -0.1

Value

the figures of hierarchical cluster.
Author(s)
Shuangbin Xu

Examples
```r
## Not run:
library(phyloseq)
library(ggtree)
library(ggplot2)
data(GlobalPatterns)
subGlobal <- subset_samples(GlobalPatterns, SampleType %in% c("Feces", "Mock", "Ocean", "Skin"))
hcsample <- get_clust(subGlobal, distmethod="jaccard", method="hellinger", hclustmethod="average")
hc_p <- ggclust(hcsample, layout = "rectangular", pointsize=1, fontsize=0,
   factorNames=c("SampleType")) + theme_tree2(legend.position="right", plot.title = element_text(face="bold", lineheight=25,hjust=0.5))
## End(Not run)
```

Description
boxplot for the result of `diff_analysis`

Usage
`ggdiffbox(obj, ...)`

```r
## S4 method for signature 'diffAnalysisClass'

ggdiffbox(
   obj,
   geom = "boxplot",
   box_notch = TRUE,
   box_width = 0.05,
   dodge_width = 0.6,
   addLDA = TRUE,
   factorLevels = NULL,
   featurelist = NULL,
   removeUnknown = TRUE,
   colorlist = NULL,
   l_xlabtext = NULL,
   ...
)
```
Arguments

obj object, diffAnalysisClass class.

..., additional arguments.

geom character, "boxplot" or "violin", default is "boxplot".

box_notch logical, see also 'notch' of geom_boxplot, default is TRUE.

box_width numeric, the width of boxplot, default is 0.05

dodge_width numeric, the width of dodge of boxplot, default is 0.6.

addLDA logical, whether add the plot to visualize the result of LDA, default is TRUE.

factorLevels list, the levels of the factors, default is NULL, if you want to order the levels of factor, you can set this.

featurelist vector, the character vector, the sub feature of originalD in diffAnalysisClass, default is NULL.

removeUnknown logical, whether remove the unknown taxonomy, default is TRUE.

colorlist character, the color vector, default is NULL.

l_xlabtext character, the x axis text of left panel, default is NULL.

Value

a 'ggplot' plot object, a box or violine plot for the result of diffAnalysisClass.

Author(s)

Shuangbin Xu

Examples

```r
# Not run:
data(kostic2012crc)
kostic2012crc %<>% as.phyloseq()
head(phyloseq::sample_data(kostic2012crc),3)
kostic2012crc <- phyloseq::rarefy_even_depth(kostic2012crc, rngseed=1024)
table(phyloseq::sample_data(kostic2012crc)$DIAGNOSIS)
set.seed(1024)
diffres <- diff_analysis(kostic2012crc, classgroup="DIAGNOSIS", mlfun="lda", filtermod="fdr", firstcomfun = "kruskal.test", firstalpha=0.05, strictmod=TRUE, secondcomfun = "wilcox.test", subclmin=3, subclwilc=TRUE, secondalpha=0.01, ldascore=3)
library(ggplot2)
p <- ggdiffbox(diffres, box_notch=FALSE, l_xlabtext="relative abundance")
# set factor levels
p2 <- ggdiffbox(diffres, box_notch=FALSE, l_xlabtext="relative abundance", factorLevels=list(DIAGNOSIS=c("Tumor", "Healthy")))
```

```
## End(Not run)
```
ggdiffclade

plot the clade tree with highlight

Description
plot results of different analysis or data.frame, contained hierarchical relationship or other classes, such as the tax_data of phyloseq.

Usage

ggdiffclade(obj, ...)

## S3 method for class 'data.frame'

ggdiffclade(
  obj,
  nodedf,
  factorName,
  size,
  layout = "radial",
  linewd = 0.6,
  bg.tree.color = "#bed0d1",
  bg.point.color = "#bed0d1",
  bg.point.stroke = 0.2,
  bg.point.fill = "white",
  skpointsize = 2,
  hilight.size = 0.2,
  alpha = 0.4,
  taxlevel = 5,
  cladetext = 2.5,
  tip.annot = TRUE,
  as.tiplab = TRUE,
  factorLevels = NULL,
  xlim = 12,
  removeUnknown = FALSE,
  reduce = FALSE,
  type = "species",
  ...
)

## S3 method for class 'diffAnalysisClass'

ggdiffclade(obj, size, removeUnknown = TRUE, ...)

Arguments

obj
  object, diffAnalysisClass, the results of diff_analysis see also diff_analysis, or data.frame, contained hierarchical relationship or other classes.

..., additional parameters.
nodedf data.frame, contained the tax and the factor information and/or pvalue.

factorName character, the names of factor in nodedf.

size the column name for mapping the size of points, default is 'pvalue'.

layout character, the layout of ggtree, but only 'rectangular', 'roundrect', 'ellipse', 'radial', 'slanted', 'inward_circular' and 'circular' in here, default is 'radial'.

linewd numeric, the size of segment of ggtree, default is 0.6.

bg.tree.color character, the line color of tree, default is '#bed0d1'.

bg.point.color character, the color of margin of background node points of tree, default is '#bed0d1'.

bg.point.stroke numeric, the margin thickness of point of background nodes of tree, default is 0.2.

bg.point.fill character, the point fill (since point shape is 21) of background nodes of tree, default is 'white'.

skpointsize numeric, the point size of skeleton of tree, default is 2.

hilight.size numeric, the margin thickness of high light clade, default is 0.2.

alpha numeric, the alpha of clade, default is 0.4.

taxlevel positive integer, the full text of clade, default is 5.

cladetext numeric, the size of text of clade, default is 2.

tip.annot logical whether to replace the differential tip labels with shorthand, default is TRUE.

as.tiplab logical, whether to display the differential tip labels with 'geom_tiplab' of 'ggtree', default is TRUE, if it is FALSE, it will use 'geom_text_repel' of 'ggrepel'.

factorLevels list, the levels of the factors, default is NULL, if you want to order the levels of factor, you can set this.

xlim numeric, the x limits, only works for 'inward_circular' layout, default is 12.

removeUnknown logical, whether do not show unknown taxonomy, default is TRUE.

reduce logical, whether remove the unassigned taxonomy, which will remove the clade of unassigned taxonomy, but the result of 'diff_analysis' should remove the unknown taxonomy, default is FALSE.

type character, the type of datasets, default is "species", if the dataset is not about species, such as dataset of kegg function, you should set it to "others".

Value

figures of tax clade show the significant different feature.

Author(s)

Shuangbin Xu
Examples

```r
## Not run:
data(kostic2012crc)
kostic2012crc %<>% as.phyloseq()
head(phyloseq::sample_data(kostic2012crc), 3)
kostic2012crc <- phyloseq::rarefy_even_depth(kostic2012crc,
  rngseed = 1024)
table(phyloseq::sample_data(kostic2012crc)$DIAGNOSIS)
set.seed(1024)
diffres <- diff_analysis(kostic2012crc, classgroup = "DIAGNOSIS",
  mlfun = "lda", filtermod = "fdr",
  firstcomfun = "kruskal.test",
  firstalpha = 0.05, strictmod = TRUE,
  secondcomfun = "wilcox.test",
  subclmin = 3, subclwilc = TRUE,
  secondalpha = 0.01, ldascore = 3)
library(ggplot2)
diffcladeplot <- ggdiffclade(diffres, alpha = 0.3, linewd = 0.2,
  skpointsize = 0.4,
  taxlevel = 5) +
  scale_fill_diff_cladogram(
    values = c("#00AED7",
               "#FD9347"
  )
) +
  scale_size_continuous(range = c(1, 3))
## End(Not run)
```

ggdiffcladeplot

---

**Description**

significantly discriminative feature barplot

**Usage**

```r
ggdifftaxbar(obj, ...)
ggdiffbartaxa(obj, ...)
```

## S4 method for signature 'diffAnalysisClass'

```r
ggdifftaxbar(
  obj,
  filepath = NULL,
  output = "biomarker_barplot",
  removeUnknown = TRUE,
  figwidth = 6,
```
figheight = 3,
ylabel = "relative abundance",
format = "pdf",
dpi = 300,
...)

## S3 method for class 'featureMeanMedian'
ggdifftaxbar(
  obj,
  featurename,
  classgroup,
  subclass,
  xtextsize = 3,
  factorLevels = NULL,
  coloslist = NULL,
  ylabel = "relative abundance",
  ...
)

Arguments

obj | object, diffAnalysisClass see also diff_analysis or feMeanMedian class, see also get_mean_median.
...
... | additional arguments.
filepath         | character, default is NULL, meaning current path.
output          | character, the output dir name, default is "biomarker_barplot".
removeUnknown       | logical, whether do not show unknown taxonomy, default is TRUE.
figwidth         | numeric, the width of figures, default is 6.
figheight        | numeric, the height of figures, default is 3.
ylabel            | character, the label of y, default is 'relative abundance'.
format            | character, the format of figure, default is pdf, png, tiff also be supported.
dpi               | numeric, the dpi of output, default is 300.
featurename      | character, the feature name, contained at the objet.
classgroup       | character, factor name.
subclass         | character, factor name.
xtextsize        | numeric, the size of axis x label, default is 3.
factorLevels     | list, the levels of the factors, default is NULL, if you want to order the levels of factor, you can set this.
coloslist        | vector, color vector, if the input is phyloseq, you should use this to adjust the color, not scale_color_manual.

Value

the figures of features show the distributions in samples.
## Examples

```r
## Not run:
data(kostic2012crc)
kostic2012crc %<>% as.phyloseq()
head(phyloseq::sample_data(kostic2012crc),3)
kostic2012crc <- phyloseq::rarefy_even_depth(kostic2012crc, rngseed=1024)
table(phyloseq::sample_data(kostic2012crc)$DIAGNOSIS)
set.seed(1024)
diffres <- diff_analysis(kostic2012crc, classgroup="DIAGNOSIS",
                        mfun="lda", filtermod="fdr",
                        firstcomfun = "kruskal.test",
                        firstalpha=0.05, strictmod=TRUE,
                        secondcomfun = "wilcox.test",
                        subclmin=3, subclwilc=TRUE,
                        secondalpha=0.01, ldascore=3)
ggdifftaxbar(diffres, output="biomarker_barplot")
## End(Not run)
```

---

**ggeffectsize**

visualization of effect size by the Linear Discriminant Analysis or randomForest

### Usage

```r
ggeffectsize(obj, ...)
```

## S3 method for class 'data.frame'

```r
ggeffectsize(
  obj,
  factorName,
  effectsizename,
  factorLevels = NULL,
  linecolor = "grey50",
  linewidth = 0.4,
  lineheight = 0.2,
  pointsize = 1.5,
  setFacet = TRUE,
  ...
)```

### Description

visualization of effect size by the Linear Discriminant Analysis or randomForest

Author(s)

Shuangbin Xu
## S3 method for class 'diffAnalysisClass'
ggeffectsize(obj, removeUnknown = TRUE, setFacet = TRUE, ...)

**Arguments**

- `obj` object, `diffAnalysisClass` see `diff_analysis`, or data.frame, contained effect size and the group information.
- `...` additional arguments.
- `factorName` character, the column name contained group information in data.frame.
- `effectSizeName` character, the column name contained effect size information.
- `factorLevels` list, the levels of the factors, default is NULL, if you want to order the levels of factor, you can set this.
- `lineColor` character, the color of horizontal error bars, default is grey50.
- `lineWidth` numeric, the width of horizontal error bars, default is 0.4.
- `lineHeight` numeric, the height of horizontal error bars, default is 0.2.
- `pointSize` numeric, the size of points, default is 1.5.
- `setFacet` logical, whether use facet to plot, default is TRUE.
- `removeUnknown` logical, whether do not show unknown taxonomy, default is TRUE.

**Value**

The figures of effect size show the LDA or MDA (MeanDecreaseAccuracy).

**Author(s)**

Shuangbin Xu

**Examples**

```r
## Not run:
data(kostic2012crc)
kostic2012crc %<>% as.phyloseq()
head(phyloseq::sample_data(kostic2012crc),3)
kostic2012crc <- phyloseq::rarefy_even_depth(kostic2012crc,rngseed=1024)
table(phyloseq::sample_data(kostic2012crc)$DIAGNOSIS)
set.seed(1024)
diffres <- diff_analysis(kostic2012crc, classgroup="DIAGNOSIS",
                        mlfun="lda", filtermod="fdr",
                        firstcomfun = "kruskal.test",
                        firstalpha=0.05, strictmod=TRUE,
                        secondcomfun = "wilcox.test",
                        subclmin=3, subclwilc=TRUE,
                        secondalpha=0.01, ldascore=3)
library(ggplot2)
effectplot <- ggeffectsize(diffres) +
              scale_color_manual(values=c('#00AED7',
```

ggordpoint

ordination plotter based on ggplot2.

Description

ordination plotter based on ggplot2.

Usage

ggordpoint(obj, ...)

## Default S3 method:

## End(Not run)

'"FD9347",
"#C1E168")+
theme_bw() +
theme(strip.background=element_rect(fill=NA),
panel.spacing = unit(0.2, "mm"),
panel.grid=element_blank(),
strip.text.y=element_blank())

## End(Not run)

ggordpoint ordination plotter based on ggplot2.

Description

ordination plotter based on ggplot2.

Usage

ggordpoint(obj, ...)

## Default S3 method:

## End(Not run)

'"FD9347",
"#C1E168")+
theme_bw() +
theme(strip.background=element_rect(fill=NA),
panel.spacing = unit(0.2, "mm"),
panel.grid=element_blank(),
strip.text.y=element_blank())

## End(Not run)
## S3 method for class 'pcasample'
ggordpoint(obj, ...)

Arguments

- `obj`: prcomp class or pcasample class.
- `...`: additional parameters, see `geom_text_repel`.
- `pc`: integer vector, the component index.
- `mapping`: set of aesthetic mapping of ggplot2, default is NULL when you want to set it by yourself, only alpha can be setted, and the first element of factorNames has been setted to map 'fill', and the second element of factorNames has been setted to map 'starshape', you can add 'scale_starshape_manual' of 'ggstar' to set the shapes.
- `sampleda`: data.frame, nrow sample * ncol factors, default is NULL.
- `factorNames`: vector, the names of factors contained sampleda.
- `factorLevels`: list, the levels of the factors, default is NULL, if you want to order the levels of factor, you can set this.
- `poinsize`: numeric, the size of point, default is 2.
- `linesize`: numeric, the line size of segment, default is 0.3.
- `arrowsize`: numeric, the size of arrow, default is 1.5.
- `arrowlinecolour`: character, the color of segment, default is grey.
- `ellipse`: logical, whether add confidence ellipse to ordinary plot, default is FALSE.
- `showsample`: logical, whether show the labels of sample, default is FALSE.
- `ellipse_pro`: numeric, confidence value for the ellipse, default is 0.9.
- `ellipse_alpha`: numeric, the alpha of ellipse, default is 0.2.
- `ellipse_linewd`: numeric, the width of ellipse line, default is 0.5.
- `ellipse_lty`: integer, the type of ellipse line, default is 3
- `biplot`: logical, whether plot the species, default is FALSE.
- `topn`: integer or vector, the number species have top important contribution, default is 5.
- `settheme`: logical, whether set the theme for the plot, default is TRUE.
- `speciesannot`: logical, whether plot the species, default is FALSE.
- `fontsize`: numeric, the size of text, default is 2.5.
- `labelfactor`: character, the factor want to be show in label, default is NULL.
- `stroke`: numeric, the line size of points, default is 0.1.
- `fontface`: character, the font face, default is "bold.italic".
- `fontfamily`: character, the font family, default is "sans".
- `textlinesize`: numeric, the segment size in `geom_text_repel`.
**Value**

point figures of PCA or PCoA.

**Author(s)**

Shuangbin Xu

**Examples**

```r
## Not run:
library(phyloseq)
data(GlobalPatterns)
subGlobal <- subset_samples(GlobalPatterns, SampleType %in% c("Feces", "Mock", "Ocean", "Skin"))
pcares <- get_pca(subGlobal, method="hellinger")
pcaplot <- ggordpoint(pcares, biplot=TRUE, speciesannot=TRUE, factorNames=c("SampleType"), ellipse=TRUE)
## End(Not run)
```

---

**ggrarecurve**  
*Rarefaction alpha index*

**Description**

Rarefaction alpha index

**Usage**

```r
ggrarecurve(obj, ...)
```

- **S3 method for class 'phyloseq'**
  ```r
ggrarecurve(obj, chunks = 400, factorLevels = NULL, ...)
  ```

- **S3 method for class 'data.frame'**
  ```r
ggrarecurve(obj, sampleda, factorLevels, chunks = 400, ...)
  ```

- **S3 method for class 'rarecurve'**
  ```r
ggrarecurve(
    obj,
    indexNames = "Observe",
    linesize = 0.5,
    facetnrow = 1,
    shadow = TRUE,
    factorNames,
    se = FALSE,
    method = "lm",
  )
  ```
formula = y ~ log(x),

Arguments

- **obj**: phyloseq, phyloseq class or data.frame shape of data.frame (nrow sample * ncol feature ( + factor)).
- **...**: additional parameters, see also ggplot2[ggplot].
- **chunks**: integer, the number of subsample in a sample, default is 400.
- **factorLevels**: list, the levels of the factors, default is NULL, if you want to order the levels of factor, you can set this.
- **sampled**: data.frame, (nrow sample * ncol factor)
- **indexNames**: character, default is "Observe", only for "Observe", "Chao1", "ACE".
- **linesize**: integer, default is 0.5.
- **facetnrow**: integer, the nrow of facet, default is 1.
- **shadow**: logical, whether merge samples with group (factorNames) and display the ribbon of group, default is TRUE.
- **factorNames**: character, default is missing.
- **se**: logical, default is FALSE.
- **method**: character, default is lm.
- **formula**: formula, default is ‘y ~ log(x)’

Value

figure of rarefaction curves

Author(s)

Shuangbin Xu

Examples

```r
## Not run:
data(test_otu_data)
test_otu_data %<>% as.phyloseq()
library(ggplot2)
prare <- ggrarecurve(test_otu_data,
  indexNames=c("Observe","Chao1","ACE"),
  shadow=FALSE,
  factorNames="group"
) +
  theme(legend.spacing.y=unit(0.02,"cm"),
        legend.text=element_text(size=6))

## End(Not run)
```
Import function to load the feature table and taxonomy table of dada2

Description

the function can import the output of dada2, and generated the phyloseq obj contained the argument class.

Usage

import_dada2(seqtab, taxatab = NULL, reftree = NULL, sampleda = NULL, ...)

mp_import_dada2(seqtab, taxatab = NULL, reftree = NULL, sampleda = NULL, ...)

Arguments

seqtab matrix, feature table, the output of removeBimeraDenovo.
taxatab matrix, a taxonomic table, the output of assignTaxonomy, or the output of addSpecies.
reftree phylo, treedata or character, the treedata or phylo class of tree, or the tree file.
sampleda data.frame or character, the data.frame of sample information, or the file of sample information, nrow samples X ncol factors.
..., additional parameters.

Value

phyloseq class contained the argument class.

Author(s)

Shuangbin Xu

Examples

seqtabfile <- system.file("extdata", "seqtab.nochim.rds", package="MicrobiotaProcess")
taxafile <- system.file("extdata", "taxa_tab.rds", package="MicrobiotaProcess")
seqtab <- readRDS(seqtabfile)
taxa <- readRDS(taxafile)
sampled <- system.file("extdata", "mouse.time.dada2.txt", package="MicrobiotaProcess")
mpse <- mp_import_dada2(seqtab=seqtab, taxatab=taxa, sampleda=sampled)
mpse
Import function to load the output of qiime2.

Description

The function was designed to import the output of qiime2 and convert them to phyloseq class.

Usage

import_qiime2(
  otuqza,
  taxaqza = NULL,
  mapfilename = NULL,
  refseqqza = NULL,
  treeqza = NULL,
  parallel = FALSE,
  ...
)

mp_import_qiime2(
  otuqza,
  taxaqza = NULL,
  mapfilename = NULL,
  refseqqza = NULL,
  treeqza = NULL,
  parallel = FALSE,
  ...
)

Arguments

- **otuqza**: character, the file contained otu table, the output of qiime2.
- **taxaqza**: character, the file contained taxonomy, the output of qiime2, default is NULL.
- **mapfilename**: character, the file contained sample information, the tsv format, default is NULL.
- **refseqqza**: character, the file contained reference sequences or the XStringSet object, default is NULL.
- **treeqza**: character, the file contained the tree file or treedata object, which is the result parsed by functions of treeio, default is NULL.
- **parallel**: logical, whether parsing the column of taxonomy multi-parallel, default is FALSE.
- **...**: additional parameters.

Value

MPSE-class or phyloseq-class contained the argument class.
Author(s)

Shuangbin Xu

Examples

```r
otuqzafile <- system.file("extdata", "table.qza",
    package="MicrobiotaProcess")
taxaqzafile <- system.file("extdata", "taxa.qza",
    package="MicrobiotaProcess")
mapfile <- system.file("extdata", "metadata_qza.txt",
    package="MicrobiotaProcess")
mpse <- mp_import_qiime2(otuqza=otuqzafile, taxaqza=taxaqzafile,
    mapfilename=mapfile)
mpse
```

(mouse.time.mpse) An example data

Description

This is a MPSE object example data.

MPSE

Construct a MPSE object

Usage

```r
MPSE(
    assays,
    colData = NULL,
    otutree = NULL,
    taxatree = NULL,
    refseq = NULL,
    ...
)
```
Arguments

assays  A 'list' or 'SimpleList' of matrix-like elements All elements of the list must have
the same dimensions, we also recommend they have names, e.g. list(\texttt{Abundance}=xx1,
Rare\texttt{Abundance}=xx2).
colData  An optional DataFrame describing the samples.
otutree  A treedata object of tidytree package, the result parsed by the functions of treeio.
taxatree  A treedata object of tidytree package, the result parsed by the functions of treeio.
refseq  A XStringSet object of Biostrings package, the result parsed by the readDNAs-
stringSet or readAAStringSet of Biostrings.
...  additional parameters, see also the usage of \texttt{SummarizedExperiment}.

Value

MPSE object

Examples

```r
set.seed(123)
x <- matrix(abs(round(rnorm(100, sd=4), 0)), 10)
x <- data.frame(x)
rownames(x) <- paste0("row", seq_len(10))
mpse <- MPSE(assays=xx)
mpse
```

---

MPSE-accessors  \textit{MPSE} accessors

Description

MPSE accessors

Usage

```
## S4 method for signature 'MPSE,ANY,ANY,ANY'
x[i, j, ..., drop = TRUE]

## S4 replacement method for signature 'MPSE,DataFrame'
colData(x, ...) <- value

## S4 replacement method for signature 'MPSE,NULL'
colData(x, ...) <- value

tax_table(object)

## S4 method for signature 'MPSE'
tax_table(object)
```
## S4 method for signature 'tbl_mpse'
tax_table(object)

## S4 method for signature 'grouped_df_mpse'
tax_table(object)

otutree(x, ...)

## S4 method for signature 'MPSE'

otutree(x, ...)

## S4 method for signature 'tbl_mpse'

otutree(x, ...)

## S4 method for signature 'MPSE'

otutree(x, ...)

otutree(x, ...) <- value

## S4 replacement method for signature 'MPSE,treedata'

otutree(x, ...) <- value

## S4 replacement method for signature 'MPSE,phylo'

otutree(x, ...) <- value

## S4 replacement method for signature 'MPSE,NULL'

otutree(x, ...) <- value

## S4 replacement method for signature 'tbl_mpse,treedata'

otutree(x, ...) <- value

## S4 replacement method for signature 'grouped_df_mpse,treedata'

otutree(x, ...) <- value

## S4 replacement method for signature 'tbl_mpse,NULL'

otutree(x, ...) <- value

## S4 replacement method for signature 'grouped_df_mpse,NULL'

otutree(x, ...) <- value

taxatree(x, ...)

## S4 method for signature 'MPSE'

taxatree(x, ...)

## S4 method for signature 'tbl_mpse'

taxatree(x, ...)
## S4 method for signature 'grouped_df_mpse'
taxatree(x, ...)
taxatree(x, ...) <- value

## S4 replacement method for signature 'MPSE,treedata'
taxatree(x, ...) <- value

## S4 replacement method for signature 'MPSE,NULL'
taxatree(x, ...) <- value

## S4 replacement method for signature 'tbl_mpse,treedata'
taxatree(x, ...) <- value

## S4 replacement method for signature 'tbl_mpse,NULL'
taxatree(x, ...) <- value

## S4 replacement method for signature 'grouped_df_mpse,treedata'
taxatree(x, ...) <- value

## S4 replacement method for signature 'grouped_df_mpse,NULL'
taxatree(x, ...) <- value
taxonomy(x, ...) <- value

## S4 replacement method for signature 'MPSE,data.frame'
taxonomy(x, ...) <- value

## S4 replacement method for signature 'MPSE,matrix'
taxonomy(x, ...) <- value

## S4 replacement method for signature 'MPSE,taxonomyTable'
taxonomy(x, ...) <- value

## S4 replacement method for signature 'MPSE,NULL'
taxonomy(x, ...) <- value

refsequence(x, ...)

## S4 method for signature 'MPSE'
refsequence(x, ...)

refsequence(x, ...) <- value

## S4 replacement method for signature 'MPSE,XStringSet'
refsequence(x, ...) <- value
## S4 replacement method for signature 'MPSE,NULL'
refsequence(x, ...) <- value

## S4 replacement method for signature 'MPSE'
rownames(x) <- value

### Arguments

- **x**: MPSE object
- **i, j, ...**: Indices specifying elements to extract or replace. Indices are 'numeric' or 'character' vectors or empty (missing) or NULL. Numeric values are coerced to integer as by 'as.integer' (and hence truncated towards zero). Character vectors will be matched to the 'names' of the object (or for matrices/arrays, the 'dimnames')
- **drop**: logical If 'TRUE' the result is coerced to the lowest possible dimension (see the examples). This only works for extracting elements, not for the replacement.
- **value**: XStringSet object or NULL
- **object**: parameter of tax_table, R object, MPSE class in here.

### Value

taxonomyTable class

### Description

MPSE class

### Slots

- **otutree**: A treedata object of tidytree package or NULL.
- **taxatree**: A treedata object of tidytree package or NULL.
- **refseq**: A XStringSet object of Biostrings package or NULL.
- **...**: Other slots from SummarizedExperiment
Permutational Multivariate Analysis of Variance Using Distance Matrices for MPSE or tbl_mpse object

Description

Permutational Multivariate Analysis of Variance Using Distance Matrices for MPSE or tbl_mpse object

Usage

```r
mp_adonis(
  .data,  # data frame
  .abundance,  # character string specifying the abundance column
  .formula,  # formula
  distmethod = "bray",  # distance method
  action = "get",  # action to perform
  permutations = 999,  # number of permutations
  seed = 123,  # seed for the random number generator
  ...  # additional arguments
)
```

## S4 method for signature 'MPSE'
```r
mp_adonis(
  .data,  # data frame
  .abundance,  # character string specifying the abundance column
  .formula,  # formula
  distmethod = "bray",  # distance method
  action = "get",  # action to perform
  permutations = 999,  # number of permutations
  seed = 123,  # seed for the random number generator
  ...  # additional arguments
)
```

## S4 method for signature 'tbl_mpse'
```r
mp_adonis(
  .data,  # data frame
  .abundance,  # character string specifying the abundance column
  .formula,  # formula
  distmethod = "bray",  # distance method
  action = "get",  # action to perform
  permutations = 999,  # number of permutations
  seed = 123,  # seed for the random number generator
  ...  # additional arguments
)
```

## S4 method for signature 'grouped_df_mpse'
```r
mp_adonis(
  .data,  # data frame
  .abundance,  # character string specifying the abundance column
  .formula,  # formula
  distmethod = "bray",  # distance method
  action = "get",  # action to perform
  permutations = 999,  # number of permutations
  seed = 123,  # seed for the random number generator
  ...  # additional arguments
)
```
\textbf{mp\_adonis}

\begin{verbatim}
mp\_adonis(
  .data,
  .abundance,
  .formula,
  distmethod = "bray",
  action = "get",
  permutations = 999,
  seed = 123,
  ...
)
\end{verbatim}

\textbf{Arguments}

.\textbf{data} \hspace{1cm} \text{MPSE or tbl\_mpse object}

.\textbf{abundance} \hspace{1cm} \text{the name of abundance to be calculated.}

.\textbf{formula} \hspace{1cm} \text{Model formula right hand side gives the continuous variables or factors, and keep left empty, such as ~ group, it is required.}

.\textbf{distmethod} \hspace{1cm} \text{character the method to calculate pairwise distances, default is 'bray'.}

.\textbf{action} \hspace{1cm} \text{character "add" joins the cca result to the object, "only" return a non-redundant tibble with the cca result. "get" return 'cca' object can be analyzed using the related vegan function.}

.\textbf{permutations} \hspace{1cm} \text{the number of permutations required, default is 999.}

.\textbf{seed} \hspace{1cm} \text{a random seed to make the adonis analysis reproducible, default is 123.}

.\textbf{...} \hspace{1cm} \text{additional parameters see also 'adonis2' of vegan.}

\textbf{Value}

\text{update object according action argument}

\textbf{Author(s)}

Shuangbin Xu

\textbf{Examples}

data(mouse\_time\_mpse)
mouse\_time\_mpse $\%\%$
  mp\_decostand(
    .abundance=Abundance,
    method="hellinger") $\%\%$
  mp\_adonis(.abundance=hellinger,
    .formula=time,
    distmethod="bray",
    permutations=999, # for more robust, set it to 9999.
    action="get")
mp_aggregate

aggregate the assays with the specific group of sample and fun.

Description

aggregate the assays with the specific group of sample and fun.

Usage

mp_aggregate(.data, .abundance, .group, fun = sum, keep_colData = TRUE, ...)

## S4 method for signature 'MPSE'
mp_aggregate(.data, .abundance, .group, fun = sum, keep_colData = TRUE, ...)

Arguments

- .data: MPSE object, required
- .abundance: the column names of abundance, default is Abundance.
- .group: the column names of sample meta-data, required
- fun: a function to compute the summary statistics, default is sum.
- keep_colData: logical whether to keep the sample meta-data with .group as row names, default is TRUE.
- ...: additional parameters, see also aggregate.

Value

a new object with .group as column names in assays

Examples

## Not run:
data(mouse.time.mpse)
newmpse <- mouse.time.mpse %>%
  mp_aggregate(.group = time)
newmpse

## End(Not run)
**mp_aggregate_clade**

calculate the mean/median (relative) abundance of internal nodes according to their children tips.

**Description**

calculate the mean/median (relative) abundance of internal nodes according to their children tips.

**Usage**

```r
mp_aggregate_clade(
  .data,
  .abundance = NULL,
  force = FALSE,
  relative = TRUE,
  aggregate_fun = c("mean", "median", "geometric.mean"),
  action = "get",
  ...
)
```

## S4 method for signature 'MPSE'

```r
mp_aggregate_clade(
  .data,
  .abundance = NULL,
  force = FALSE,
  relative = TRUE,
  aggregate_fun = c("mean", "median", "geometric.mean"),
  action = "get",
  ...
)
```

## S4 method for signature 'tbl_mpse'

```r
mp_aggregate_clade(
  .data,
  .abundance = NULL,
  force = FALSE,
  relative = TRUE,
  aggregate_fun = c("mean", "median", "geometric.mean"),
  action = "get",
  ...
)
```

## S4 method for signature 'grouped_df_mpse'

```r
mp_aggregate_clade(
  .data,
  .abundance = NULL,
  force = FALSE,
  relative = TRUE,
  aggregate_fun = c("mean", "median", "geometric.mean"),
  action = "get",
  ...
)
```
relative = TRUE,
aaggregate_fun = c("mean", "median", "geometric.mean"),
action = "get",
...
)

Arguments

.data MPSE object which must contain otutree slot, required
.abundance the column names of abundance.
force logical whether calculate the (relative) abundance forcibly when the abundance
is not be rarefied, default is FALSE.
relative logical whether calculate the relative abundance.
aggregate_fun function the method to calculate the (relative) abundance of internal nodes ac-
cording to their children tips, default is 'mean', other options are 'median', 'ge-
ometric.mean'.
action character, "add" joins the new information to the otutree slot if it exists (default).
In addition, "only" return a non-redundant tibble with the just new information.
"get" return a new 'mpse', which the features is the internal nodes.
... additional parameters, meaningless now.

Value

a object according to 'action' argument.

Examples

## Not run:
suppressPackageStartupMessages(library(curatedMetagenomicData))
xx <- curatedMetagenomicData('ZellerG_2014.relative_abundance', dryrun=F)
xx[[1]] %>% as.mpse -> mpse
otu.tree <- mpse %>%
  mp_aggregate_clade(
    .abundance = Abundance,
    force = TRUE,
    relative = FALSE,
    action = 'get' # other option is 'add' or 'only'.
  )
  otu.tree

## End(Not run)
Analysis of Similarities (ANOSIM) with MPSE or tbl_mpse object

Description

Analysis of Similarities (ANOSIM) with MPSE or tbl_mpse object

Usage

mp_anosim(
  .data,
  .abundance,
  .group,
  distmethod = "bray",
  action = "add",
  permutations = 999,
  seed = 123,
  ...
)

## S4 method for signature 'MPSE'
mp_anosim(
  .data,
  .abundance,
  .group,
  distmethod = "bray",
  action = "add",
  permutations = 999,
  seed = 123,
  ...
)

## S4 method for signature 'tbl_mpse'
mp_anosim(
  .data,
  .abundance,
  .group,
  distmethod = "bray",
  action = "add",
  permutations = 999,
  seed = 123,
  ...
)

## S4 method for signature 'grouped_df_mpse'
mp_anosim(
  .data,
mp_anosim

Arguments

.data MPSE or tbl_mpse object
.abundance the name of abundance to be calculated.
.group The name of the column of the sample group information.
distmethod character the method to calculate pairwise distances, default is 'bray'.
.action character "add" joins the ANOSIM result to internal attribute of the object, "only" and "get" return 'anosim' object can be analyzed using the related vegan function.
.permutations the number of permutations required, default is 999.
.seed a random seed to make the ANOSIM analysis reproducible, default is 123.
... additional parameters see also 'anosim' of vegan.

Value

update object according action argument

Author(s)

Shuangbin Xu

Examples

data(mouse.time.mpse)
mouse.time.mpse %<>%
m_p_decostand(.abundance=Abundance)
# action = "get" will return a anosim object
mouse.time.mpse %>%
m_p_anosim(.abundance=hellinger, .group=time, action="get")
# action = "only" will return a tbl_df that can be as the input of ggplot2.
library(ggplot2)
tbl <- mouse.time.mpse %>%
m_p_anosim(.abundance=hellinger, .group=time,
.permutations=999, # for more robust, set it to 9999
.action="only")
tbl
tbl %>%
ggplot(aes(x=class, y=rank, fill=class)) +
geom_boxplot(notch=TRUE, varwidth = TRUE)
Calculating the balance score of internal nodes (clade) according to the geometric.mean/mean/median abundance of their binary children tips.

**Description**

Calculating the balance score of internal nodes (clade) according to the geometric.mean/mean/median abundance of their binary children tips.

**Usage**

```r
mp_balance_clade(
  .data,
  .abundance = NULL,
  force = FALSE,
  relative = TRUE,
  balance_fun = c("geometric.mean", "mean", "median"),
  pseudonum = 0.001,
  action = "get",
  ...
)
```

```r
## S4 method for signature 'MPSE'
mp_balance_clade(
  .data,
  .abundance = NULL,
  force = FALSE,
  relative = TRUE,
  balance_fun = c("geometric.mean", "mean", "median"),
  pseudonum = 0.001,
  action = "get",
  ...
)
```

```r
## S4 method for signature 'tbl_mpse'
mp_balance_clade(
  .data,
  .abundance = NULL,
  force = FALSE,
  relative = TRUE,
  balance_fun = c("geometric.mean", "mean", "median"),
  pseudonum = 0.001,
  action = "get",
  ...
)
```
## S4 method for signature 'grouped_df_mpse'

```r
mp_balance_clade(
  .data,
  .abundance = NULL,
  force = FALSE,
  relative = TRUE,
  balance_fun = c("geometric.mean", "mean", "median"),
  pseudonum = 0.001,
  action = "get",
  ...
)
```

### Arguments

- `.data` MPSE object which must contain otutree slot, required
- `.abundance` the column names of abundance.
- `force` logical whether calculate the (relative) abundance forcibly when the abundance is not be rarefied, default is FALSE.
- `relative` logical whether calculate the relative abundance.
- `balance_fun` function the method to calculate the (relative) abundance of internal nodes according to their children tips, default is 'geometric.mean', other options are 'mean' and 'median'.
- `pseudonum` numeric add a pseudo numeric to avoid the error of division in calculation, default is 0.001.
- `action` character, "add" joins the new information to the otutree slot if it exists (default). In addition, "only" return a non-redundant tibble with the just new information, "get" return a new 'MPSE' object, and the 'OTU' column is the internal nodes and 'Abundance' column is the balance scores.
- `...` additional parameters, meaningless now.

### Value

a object according to 'action' argument.

### References


Justin D Silverman, Alex D Washburne, Sayan Mukherjee, Lawrence A David. A phylogenetic transform enhances analysis of compositional microbiota data. eLife 2017;6:e21887. https://doi.org/10.7554/eLife.21887.001

### Examples

```r
## Not run:
suppressPackageStartupMessages(library(curatedMetagenomicData))
```
mp_cal_abundance

Calculate the (relative) abundance of each taxonomy class for each sample or group.

**Description**

Calculate the (relative) abundance of each taxonomy class for each sample or group.

**Usage**

```r
mp_cal_abundance(
  .data,
  .abundance = NULL,
  .group = NULL,
```

```r
xx <- curatedMetagenomicData('ZellerG_2014.relative_abundance', dryrun=F)
xx[[1]] %>% as.mpse -> mpse
mpse.balance.clade <- mpse %>%
  mp_balance_clade(
    .abundance = Abundance,
    force = TRUE,
    relative = FALSE,
    action = 'get',
    pseudonum = .01
  )
mpse.balance.clade

# Performing the Euclidean distance or PCA.
mpse.balance.clade %>%
  mp_cal_dist(.abundance = Abundance, distmethod = 'euclidean') %>%
  mp_plot_dist(.distmethod = 'euclidean', .group = disease, group.test = T)

mpse.balance.clade %>%
  mp_adonis(.abundance = Abundance, .formula=~disease, distmethod = 'euclidean', permutation = 9999)

mpse.balance.clade %>%
  mp_cal_pca(.abundance = Abundance) %>%
  mp_plot_ord(.group = disease)

# Detecting the signal balance nodes.
mpse.balance.clade %>% mp_diff_analysis(
  .abundance = Abundance,
  force = TRUE,
  relative = FALSE,
  .group = disease,
  fc.method = 'compare_mean'
)
```


Arguments

<table>
<thead>
<tr>
<th>Argument</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td><code>.data</code></td>
<td>MPSE or tbl_mpse object</td>
</tr>
<tr>
<td><code>.abundance</code></td>
<td>the name of otu abundance to be calculated</td>
</tr>
<tr>
<td><code>.group</code></td>
<td>the name of group to be calculated.</td>
</tr>
<tr>
<td><code>relative</code></td>
<td>logical whether calculate the relative abundance.</td>
</tr>
<tr>
<td><code>action</code></td>
<td>character, &quot;add&quot; joins the new information to the taxatree and otutree if they exists (default). In addition, All taxonomy class will be added the taxatree,</td>
</tr>
</tbody>
</table>
and OTU (tip) information will be added to the otutree.”only” return a non-redundant tibble with the just new information. ”get” return ‘taxatree’ slot which is a treedata object.

force logical whether calculate the relative abundance forcibly when the abundance is not be rarefied, default is FALSE.

... additional parameters.

Value
update object or tibble according the 'action'

Author(s)
Shuangbin Xu

See Also
[mp_plot_abundance()] and [mp_extract_abundance()]

Examples

data(mouse.time.mpse)
mouse.time.mpse %<>% mp_rrarefy()
mouse.time.mpse %<>% mp_cal_abundance(.abundance=RareAbundance, action="add") %>% mp_cal_abundance(.abundance=RareAbundance, .group=time, action="add")
mouse.time.mpse
library(ggplot2)
f <- mouse.time.mpse %>%
  mp_plot_abundance(
    .abundance=RelRareAbundanceBySample,
    .group = time,
    taxa.class = "Phylum",
    topn = 20,
    geom = "heatmap",
    feature.dist = "bray",
    feature.hclust = "average"
  ) %>%
  set_scale_theme(
    x = scale_fill_manual(values=c("orange", "deepskyblue")),
    aes_var = time
  )
  f
pl <- mouse.time.mpse %>%
  mp_plot_abundance(.abundance=RelRareAbundanceBySample,
    .group=time, taxa.class="Phylum",
    topn=20, order.by.feature = "p__Firmicutes",
    width = 4/5
  )
p2 <- mouse.time.mpse %>%
  mp_plot_abundance(.abundance = RareAbundance,
    .group = time,
    taxa.class = Phylum,
    topn = 20,
    relative = FALSE,
    force = TRUE,
    order.by.feature = TRUE
  )

p1 / p2

# Or you can also extract the result and visualize it with ggplot2 and ggplot2-extension
## Not run:
tbl <- mouse.time.mpse %>%
  mp_extract_abundance(taxa.class="Class", topn=10)
tbl
library(ggplot2)
library(ggalluvial)
library(dplyr)
tbl %<><%
tidyrr::unnest(cols=RareAbundanceBySample)
tbl
p <- ggplot(data=tbl,
  mapping=aes(x=Sample,
    y=RelRareAbundanceBySample,
    alluvium=label,
    fill=label)
) +
  geom_flow(stat="alluvium", lode.guidance = "frontback", color = "darkgray") +
  geom_stratum(stat="alluvium") +
  labs(x=NULL, y="Relative Abundance (%)") +
  scale_fill_brewer(name="Class", type = "qual", palette = "Paired") +
  facet_grid(cols=vars(time), scales="free_x", space="free") +
  theme(axis.text.x=element_text(angle=-45, hjust=0))
p
## End(Not run)

---

**mp_cal_alpha**

*calculate the alpha index with MPSE or tbl_mpse*

**Description**

calculate the alpha index with MPSE or tbl_mpse

**Usage**

```r
mp_cal_alpha(
  .data,
  .abundance = NULL,
  action = c("add", "only", "get"),
```

```r
```
### mp_cal_alpha

```r
force = FALSE, ...
}
```

```r
## S4 method for signature 'MPSE'
mp_cal_alpha(.data, .abundance = NULL, action = "add", force = FALSE, ...)
```

```r
## S4 method for signature 'tbl_mpse'
mp_cal_alpha(.data, .abundance = NULL, action = "add", force = FALSE, ...)
```

```r
## S4 method for signature 'grouped_df_mpse'
mp_cal_alpha(.data, .abundance = NULL, action = "add", force = FALSE, ...)
```

**Arguments**

- `.data` MPSE or tbl_mpse object
- `.abundance` The column name of OTU abundance column to be calculate
- `.action` character it has three options, "add" joins the new information to the input tbl (default), "only" return a non-redundant tibble with the just new information, ang 'get' return a 'alphasample' object.
- `.force` logical whether calculate the alpha index even the `.abundance` is not rarefied, default is FALSE.
- `...` additional arguments

**Value**

update object or other (refer to action)

**Author(s)**

Shuangbin Xu

**See Also**

[mp_plot_alpha()]

**Examples**

```r
data(mouse.time.mpse)
mpse <- mouse.time.mpse %>%
  mp_rrarefy() %>%
  mp_cal_alpha(.abundance=RareAbundance)
mpse
p <- mpse %>% mp_plot_alpha(.group=time, .alpha=c(Observe, Shannon, Pielou))
p
```

# Or you can extract the result and visualize it with ggplot2 and ggplot2-extensions
## Not run:
tbl <- mpse %>%
  mp_extract_sample
mp_cal_cca

**Description**

[Partial] [Constrained] Correspondence Analysis with MPSE or tbl_mpse object

**Usage**

```r
mp_cal_cca(.data, .abundance, .formula = NULL, .dim = 3, action = "add", ...)
```

## S4 method for signature 'MPSE'
```r
mp_cal_cca(.data, .abundance, .formula = NULL, .dim = 3, action = "add", ...)
```

## S4 method for signature 'tbl_mpse'
```r
mp_cal_cca(.data, .abundance, .formula = NULL, .dim = 3, action = "add", ...)
```

## S4 method for signature 'grouped_df_mpse'
```r
mp_cal_cca(.data, .abundance, .formula = NULL, .dim = 3, action = "add", ...)
```

**Arguments**

- `.data` MPSE or tbl_mpse object
- `.abundance` the name of abundance to be calculated.
- `.formula` Model formula right hand side gives the constraining variables, and conditioning variables can be given within a special function `Condition()` and keep left empty, such as ~ A + B or ~ A + Condition(B), default is NULL.
- `.dim` integer The number of dimensions to be returned, default is 3.
mp_cal_clust

Hierarchical cluster analysis for the samples with MPSE or tbl_mpse object

Description
Hierarchical cluster analysis for the samples with MPSE or tbl_mpse object

Usage

mp_cal_clust(
  .data,
  .abundance,
  distmethod = "bray",
  hclustmethod = "average",
  action = "get",
  ...
)

## S4 method for signature 'MPSE'
mp_cal_clust(
  .data,
.abundance,
distmethod = "bray",
hclustmethod = "average",
action = "get",
...
)

## S4 method for signature 'tbl_mpse'
mp_cal_clust(
  .data,
  .abundance,
distmethod = "bray",
hclustmethod = "average",
action = "get",
...
)

## S4 method for signature 'grouped_df_mpse'
mp_cal_clust(
  .data,
  .abundance,
distmethod = "bray",
hclustmethod = "average",
action = "get",
...
)

Arguments

.data the MPSE or tbl_mpse object
.abundance the name of abundance to be calculated.
distmethod the method of distance.
hclustmethod the method of hierarchical cluster
action a character "add" will return a MPSE object with the cluster result as a attributes, and it can be extracted with 'object "only" or "get" will return 'treedata' object, default is 'get'.
... additional parameters

Value

update object with the action argument, the treedata object contained hierarchical cluster analysis of sample, it can be visualized with 'ggtree' directly.

Author(s)

Shuangbin Xu
mp_cal_dca

Examples

```r
library(ggtree)
library(ggplot2)
data(mouse.time.mpse)
res <- mouse.time.mpse %>%
  mp_decostand(.abundance=Abundance) %>%
  mp_cal_clust(.abundance=hellinger, distmethod="bray")
res
res %>%
ggtree() +
geom_tippoint(aes(color=time))
```

mp_cal_dca  
**Detrended Correspondence Analysis with MPSE or tbl_mpse object**

Description

Detrended Correspondence Analysis with MPSE or tbl_mpse object

Usage

```r
mp_cal_dca(.data, .abundance, .dim = 3, action = "add", origin = TRUE, ...)
```

## S4 method for signature 'MPSE'

```r
mp_cal_dca(.data, .abundance, .dim = 3, action = "add", origin = TRUE, ...)
```

## S4 method for signature 'tbl_mpse'

```r
mp_cal_dca(.data, .abundance, .dim = 3, action = "add", origin = TRUE, ...)
```

## S4 method for signature 'grouped_df_mpse'

```r
mp_cal_dca(.data, .abundance, .dim = 3, action = "add", origin = TRUE, ...)
```

Arguments

- `.data`  
MPSE or tbl_mpse object

- `.abundance`  
the name of abundance to be calculated.

- `.dim`  
integer The number of dimensions to be returned, default is 3.

- `action`  
character "add" joins the 'decorana' result to the object, "only" return a non-redundant tibble with the 'decorana' result. "get" return 'decorana' object can be processed with related vegan function.

- `origin`  
logical Use true origin even in detrended correspondence analysis. default is TRUE.

- `...`  
additional parameters see also 'vegan::decorana'

Value

update object or tbl according to the action.
mp_cal_dist  

*Calculate the distances between the samples or features with specified abundance.*

**Description**

Calculate the distances between the samples or features with specified abundance.

**Usage**

```r
mp_cal_dist(
  .data,
  .abundance,
  .env = NULL,
  distmethod = "bray",
  action = "add",
  scale = FALSE,
  cal.feature.dist = FALSE,
  ...
)
```

```r
## S4 method for signature 'MPSE'
mp_cal_dist(
  .data,
  .abundance,
  .env = NULL,
  distmethod = "bray",
  action = "add",
  scale = FALSE,
  cal.feature.dist = FALSE,
  ...
)
```

```r
## S4 method for signature 'tbl_mpse'
mp_cal_dist(
  .data,
  .abundance,
  .env = NULL,
  distmethod = "bray",
  action = "add",
  scale = FALSE,
  cal.feature.dist = FALSE,
  ...
)
```

```r
## S4 method for signature 'grouped_df_mpse'
mp_cal_dist(
```
Arguments

.data | MPSE or tbl_mpse object
.abundance | the name of otu abundance to be calculated
.env | the column names of continuous environment factors, default is NULL.
distmethod | character the method to calculate distance. option is "manhattan", "euclidean", "canberra", "bray", "kulczynski", "jaccard", "gower", "altdGower", "morisita", "horn", "mountford", "raup", "binomial", "choa", "cao", "mahalanobis", "chisq", "chord", "aitchison", "robust.aitchison" (implemented in vegdist of vegan), and "w", "-1", "c", "wb", "t", "l", "e", "i", "me", "ij", "sor", "m", ".-2", "co", "cc", "g", ".-3", ".1", ".19", "hk", "rlb", "sim", "gl", "z" (implemented in betadiver of vegan), "maximum", "binary", "minkowski" (implemented in dist of stats), "unifrac", "weighted unifrac" (implemented in phyloseq), "cor", "abs_cor", "cosangle", "abs_cosangle" (implemented in hopach), or other customized distance function.
.action | character, "add" joins the distance data to the object, "only" return a non-redundant tibble with the distance information. "get" return 'dist' object.
scale | logical whether scale the metric of environment (.env is provided) before the distance was calculated, default is FALSE. The environment matrix can be processed when it was joined to the MPSE or tbl_mpse object.
cal.feature.dist | logical whether to calculate the distance between the features. default is FALSE, meaning calculate the distance between the samples.
... | additional parameters.

some dot arguments if distmethod is unifrac or weighted unifrac:

- weighted logical, whether to use weighted-UniFrac calculation, which considers the relative abundance of taxa, default is FALSE, meaning unweighted-UniFrac, which only considers presence/absence of taxa.
- normalized logical, whether normaized the branch length of tree to the range between 0 and 1 when the weighted=TRUE.
- parallel logical, whether to execute the calculation in parallel, default is FALSE.

Value

update object or tibble according the 'action'
Author(s)

Shuangbin Xu

See Also

[mp_extract_dist()] and [mp_plot_dist()]

Examples

data(mouse.time.mpse)
mouse.time.mpse %<>% 
  mp_decostand(.abundance=Abundance) %>%  
  mp_cal_dist(.abundance=hellinger, distmethod="bray")
mouse.time.mpse
p1 <- mouse.time.mpse %>% 
  mp_plot_dist(.distmethod = bray)
p2 <- mouse.time.mpse %>% 
  mp_plot_dist(.distmethod = bray, .group = time, group.test = TRUE)
p3 <- mouse.time.mpse %>% 
  mp_plot_dist(.distmethod = bray, .group = time)
  
  # adjust the legend of heatmap of distance between the samples.
  # the p3 is a aplot object, we define set_scale_theme to adjust the
  # character (color, size or legend size) of figure with specified
  # 'aes_var' according to legend title.
library(ggplot2)
p3 %>%
  set_scale_theme(  
    x = scale_size_continuous(  
      range = c(0.1, 4),  
      guide = guide_legend(keywidth = 0.5, keyheight = 1)),  
    aes_var = bray  
  ) %>%
  set_scale_theme(  
    x = scale_colour_gradient(  
      guide = guide_legend(keywidth = 0.5, keyheight = 1)),  
    aes_var = bray  
  ) %>%
  set_scale_theme(  
    x = scale_fill_manual(values = c("orangered", "deepskyblue"),  
      guide = guide_legend(keywidth = 0.5, keyheight = 0.5, label.theme = element_text(size=6))),  
    aes_var = time) %>%
  set_scale_theme(  
    x = theme(axis.text=element_text(size=6), panel.background=element_blank()),  
    aes_var = bray  
  )

## Not run:
# Visualization manual
library(ggplot2)
tbl <- mouse.time.mpse %>%
  mp_extract_dist(distmethod="bray", .group=time)
tbl
tbl %>%
**mp_cal_divergence**

mp_cal_divergence

```
ggplot(aes(x=GroupsComparison, y=bray)) +
geom_boxplot(aes(fill=GroupsComparison)) +
geom_jitter(width=0.1) +
xlab(NULL) +
theme(legend.position="none")
```

## End(Not run)

---

**mp_cal_divergence**  calculate the divergence with MPSE or tbl_mpse

**Description**

calculate the divergence with MPSE or tbl_mpse

**Usage**

```r
mp_cal_divergence(
  .data,
  .abundance,
  .name = "divergence",
  reference = "mean",
  distFUN = vegan::vegdist,
  method = "bray",
  action = "add",
  ...
)
```

## S4 method for signature 'MPSE'

```r
mp_cal_divergence(
  .data,
  .abundance,
  .name = "divergence",
  reference = "mean",
  distFUN = vegan::vegdist,
  method = "bray",
  action = "add",
  ...
)
```

## S4 method for signature 'tbl_mpse'

```r
mp_cal_divergence(
  .data,
  .abundance,
  .name = "divergence",
  reference = "mean",
  distFUN = vegan::vegdist,
```
method = "bray",
action = "add",
...
)

## S4 method for signature 'grouped_df_mpse'
mp_cal_divergence(
  .data,
  .abundance,
  .name = "divergence",
  reference = "mean",
  distFUN = vegan::vegdist,
  method = "bray",
  action = "add",
  ...
)

Arguments

.data MPSE or tbl_mpse object
.abundance The column name of OTU abundance column to be calculate.
.name the colname name of the divergence results, default is 'divergence'.
reference a no-empty character, either 'median' or 'mean' or the sample name, or a numeric vector which has length equal to the number of features, default is 'mean'.
distFUN the function to calculate the distance between the reference and samples, default is 'vegan::vegdist'.
method the method to calculate the distance, which will pass to the function that is specified in 'distFUN', default is 'bray'.
action character it has three options, "add" joins the new information to the input tbl (default), "only" return a non-redundant tibble with the just new information, ang 'get' return a 'alphasample' object.
...
additional arguments, see also the arguments of 'distFUN' function.

Value

update object or other (refer to action)

Author(s)

Shuangbin Xu

See Also

[mp_plot_alpha()]
Examples

## Not run:
# example(mp_cal_divergence, run.dontrun = TRUE) to run the example.

```r
data(mouse.time.mpse)
mouse.time.mpse %>%
  mp_cal_divergence(
    .abundance = Abundance,
    .name = 'divergence.mean',
    distFUN = vegan::vegdist,
    method = 'bray'
  ) %>%
  mp_plot_alpha(
    .alpha = divergence.mean,
    .group = time,
  )
```

## End(Not run)

---

**mp_cal_nmds**

Nonmetric Multidimensional Scaling Analysis with MPSE or tbl_mpse object

**Description**

Nonmetric Multidimensional Scaling Analysis with MPSE or tbl_mpse object

**Usage**

```r
mp_cal_nmds(
  .data, 
  .abundance, 
  distmethod = "bray", 
  .dim = 2, 
  action = "add", 
  seed = 123, 
  ... 
)
```

## S4 method for signature 'MPSE'

```r
mp_cal_nmds(
  .data, 
  .abundance, 
  distmethod = "bray", 
  .dim = 2, 
  action = "add", 
  seed = 123, 
  ... 
)
```
## S4 method for signature 'tbl_mpse'
mp_cal_nmds(
  .data, 
  .abundance, 
  distmethod = "bray", 
  .dim = 2, 
  action = "add", 
  seed = 123, 
  ...
)

## S4 method for signature 'grouped_df_mpse'
mp_cal_nmds(
  .data, 
  .abundance, 
  distmethod = "bray", 
  .dim = 2, 
  action = "add", 
  seed = 123, 
  ...
)

### Arguments
- `.data` MPSE or tbl_mpse object
- `.abundance` the name of abundance to be calculated.
- `distmethod` character the method to calculate distance.
- `.dim` integer The number of dimensions to be returned, default is 2.
- `action` character "add" joins the NMDS result to the object, "only" return a non-redundant tibble with the NMDS result. "get" return 'metaMDS' object can be analyzed with related 'vegan' function.
- `seed` a random seed to make this analysis reproducible, default is 123.
- `...` additional parameters see also 'mp_cal_dist'.

### Value
- update object or tbl according to the action.

### Author(s)
Shuangbin Xu

### Examples
```r
data(mouse.time.mpse)
mpse <- mouse.time.mpse %>%
  mp_decostand(.abundance=Abundance) %>%
```
mp_cal_pca

Principal Components Analysis with MPSE or tbl_mpse object

Description

Principal Components Analysis with MPSE or tbl_mpse object

Usage

mp_cal_pca(.data, .abundance, .dim = 3, action = "add", ...)

## S4 method for signature 'MPSE'
mp_cal_pca(.data, .abundance, .dim = 3, action = "add", ...)

## S4 method for signature 'tbl_mpse'
mp_cal_pca(.data, .abundance, .dim = 3, action = "add", ...)

## S4 method for signature 'grouped_df_mpse'
mp_cal_pca(.data, .abundance, .dim = 3, action = "add", ...)
Arguments
.data MPSE or tbl_mpse object
.abundance the name of abundance to be calculated.
.dim integer The number of dimensions to be returned, default is 3.
.action character "add" joins the pca result to the object, "only" return a non-redundant tibble with the pca result. "get" return 'prcomp' object.
... additional parameters see also 'prcomp'

Value
update object or tbl according to the action.

Author(s)
Shuangbin Xu

Examples
data(mouse.time.mpse)
library(ggplot2)
mpse <- mouse.time.mpse %>%
  mp_decostand(.abundance=Abundance) %>%
  mp_cal_pca(.abundance=hellinger, action="add")
mpse
p1 <- mpse %>% mp_plot_ord(.ord=pca, .group=time, ellipse=TRUE)
p2 <- mpse %>% mp_plot_ord(.ord=pca, .group=time, .color=time, ellipse=TRUE)
p1 + scale_fill_manual(values=c("#00AED7", "#009E73"))
p2 + scale_fill_manual(values=c("#00AED7", "#009E73")) +
  scale_color_manual(values=c("#00AED7", "#009E73"))
## Not run:
# action = "only" to extract the non-redundant tibble to visualize
tbl <- mouse.time.mpse %>%
  mp_decostand(.abundance=Abundance) %>%
  mp_cal_pca(.abundance=hellinger, action="only")
tbl
x <- names(tbl)[grepl("PC1 ", names(tbl))] %>% as.symbol()
y <- names(tbl)[grepl("PC2 ", names(tbl))] %>% as.symbol()
ggplot(tbl) +
  geom_point(aes(x=!!x, y=!!y, color=time))
## End(Not run)
Principal Coordinate Analysis with MPSE or tbl_mpse object

Description

Principal Coordinate Analysis with MPSE or tbl_mpse object

Usage

mp_cal_pcoa(
  .data,
  .abundance,
  distmethod = "bray",
  .dim = 3,
  action = "add",
  ...
)

## S4 method for signature 'MPSE'
mp_cal_pcoa(
  .data,
  .abundance,
  distmethod = "bray",
  .dim = 3,
  action = "add",
  ...
)

## S4 method for signature 'tbl_mpse'
mp_cal_pcoa(
  .data,
  .abundance,
  distmethod = "bray",
  .dim = 3,
  action = "add",
  ...
)

## S4 method for signature 'grouped_df_mpse'
mp_cal_pcoa(
  .data,
  .abundance,
  distmethod = "bray",
  .dim = 3,
  action = "add",
  ...
)

Arguments

- `.data` MPSE or tbl_mpse object
- `.abundance` the name of abundance to be calculated.
- `distmethod` character the method to calculate distance.
- `.dim` integer The number of dimensions to be returned, default is 3.
- `action` character "add" joins the pca result to the object and the 'pcoa' object also was add to the internal attributes of the object, "only" return a non-redundant tibble with the pca result. "get" return 'pcoa' object.
- `...` additional parameters see also 'mp_cal_dist'.

Value

update object or tbl according to the action.

Author(s)

Shuangbin Xu

Examples

data(mouse.time.mpse)
mpse <- mouse.time.mpse %>%
  mp_decostand(.abundance=Abundance)
mpse
mpse %<>% mp_cal_pcoa(.abundance=hellinger, distmethod="bray", action="add")
library(ggplot2)
p <- mpse %>%
  mp_plot_ord(.ord=pcoa, .group=time, .color=time, ellipse=TRUE)
p <- p +
  scale_fill_manual(values=c("#00AED7", ", #009E73")) +
  scale_color_manual(values=c("#00AED7", ", #009E73"))
## Not run:
# Or run with action='only' and return tbl_df to visualize manual.
mouse.time.mpse %>%
  mp_decostand(.abundance=Abundance) %>%
  mp_cal_pcoa(.abundance=hellinger, distmethod="bray", .dim=2, action="only") -> tbl
x <- names(tbl)[grep("PCo1 ", names(tbl))] %>% as.symbol()
y <- names(tbl)[grep("PCo2 ", names(tbl))] %>% as.symbol()
library(ggplot2)
tbl %>%
ggplot(aes(x=x!x, y=!y, color=time)) +
  stat_ellipse(aes(fill=time), geom="polygon", alpha=0.5) +
  geom_point() +
  geom_vline(xintercept=0, color="grey20", linetype=2) +
  geom_hline(yintercept=0, color="grey20", linetype=2) +
  theme_bw() +
  theme(panel.grid=element_blank())
## End(Not run)
Calculating related phylogenetic alpha metric with MPSE or tbl_mpse object

**Usage**

```r
mp_cal_pd_metric(
  .data,
  .abundance,
  action = "add",
  metric = c("PAE", "NRI", "NTI", "PD", "HAED", "EAED", "all"),
  abundance.weighted = FALSE,
  force = FALSE,
  seed = 123,
  ...
)
```

## S4 method for signature 'MPSE'

```r
mp_cal_pd_metric(
  .data,
  .abundance,
  action = "add",
  metric = c("PAE", "NRI", "NTI", "PD", "HAED", "EAED", "IAC", "all"),
  abundance.weighted = FALSE,
  force = FALSE,
  seed = 123,
  ...
)
```

## S4 method for signature 'tbl_mpse'

```r
mp_cal_pd_metric(
  .data,
  .abundance,
  action = "add",
  metric = c("PAE", "NRI", "NTI", "PD", "HAED", "EAED", "all"),
  abundance.weighted = TRUE,
  force = FALSE,
  seed = 123,
  ...
)
```

## S4 method for signature 'grouped_df_mpse'

```r
mp_cal_pd_metric(
```
.data,
.abundance,
action = "add",
metric = c("PAE", "NRI", "NTI", "PD", "HAED", "EAED", "all"),
abundance.weighted = TRUE,
force = FALSE,
seed = 123,
...)

Arguments

.data object, MPSE or tbl_mpse object
.abundance The column name of OTU abundance column to be calculate.
action character it has three options, "add" joins the new information to the input tbl (default), "only" return a non-redundant tibble with the just new information, ang 'get' return a 'alphasample' object.
metric the related phylogenetic metric, options is 'NRI', 'NTI', 'PD', 'PAE', 'HAED', 'EAED', 'IAC', 'all', default is 'PAE', 'all' meaning all the metrics ('NRI', 'NTI', 'PD', 'PAE', 'HAED', 'EAED', 'IAC').
abundance.weighted logical, whether calculate mean nearest taxon distances for each species weighted by species abundance, default is TRUE.
force logical whether calculate the alpha index even the '.abundance' is not rarefied, default is FALSE.
seed integer a random seed to make the result reproducible, default is 123.
... additional arguments see also "ses.mpd" and "ses.mntd" of "picante".

Value

update object.

Author(s)

Shuangbin Xu

References


Examples

```r
## Not run:
suppressPackageStartupMessages(library(curatedMetagenomicData))
xx <- curatedMetagenomicData('ZellerG_2014.relative_abundance', dryrun=F)
xx[[1]] %>% as.mpse -> mpse
mpse %<>%
  mp_cal_pd_metric(
    .abundance = Abundance,
    force = TRUE,
    metric = 'PAE'
  )
mpse %>%
  mp_plot_alpha(
    .alpha = PAE,
    .group = disease
  )

## End(Not run)
```

## mp_cal_rarecurve

Calculating the different alpha diversities index with different depth

Description

Calculating the different alpha diversities index with different depth

Usage

```r
mp_cal_rarecurve(
  .data, 
  .abundance = NULL, 
  action = "add", 
  chunks = 400, 
  seed = 123, 
  force = FALSE, 
  ...
)
```

```r
## S4 method for signature 'MPSE'
mp_cal_rarecurve(
  .data, 
  .abundance = NULL, 
  action = "add", 
  chunks = 400, 
  seed = 123, 
  force = FALSE, 
  ...
)
```
## S4 method for signature 'tbl_mpse'
mp_cal_rarecurve(
  .data,
  .abundance = NULL,
  action = "add",
  chunks = 400,
  seed = 123,
  force = FALSE,
  ...
)

## S4 method for signature 'grouped_df_mpse'
mp_cal_rarecurve(
  .data,
  .abundance = NULL,
  action = "add",
  chunks = 400,
  seed = 123,
  force = FALSE,
  ...
)

### Arguments

- **.data**: MPSE or tbl_mpse object
- **.abundance**: the name of otu abundance to be calculated.
- **action**: character it has three options, "add" joins the new information to the input tbl (default), "only" return a non-redundant tibble with the just new information, ang 'get' return a 'rarecurve' object.
- **chunks**: numeric the split number of each sample to calculate alpha diversity, default is 400. eg. A sample has total 40000 reads, if chunks is 400, it will be split to 100 sub-samples (100, 200, 300,..., 40000), then alpha diversity index was calculated based on the sub-samples.
- **seed**: a random seed to make the result reproducible, default is 123.
- **force**: logical whether calculate rarecurve forcibly when the `.abundance` is not be rarefied, default is FALSE
- **...**: additional parameters.

### Value

update rarecurve calss

### Author(s)

Shuangbin Xu
mp_cal_rda

See Also

[mp_plot_rarecurve()] and [mp_extract_rarecurve()]

Examples

```r
data(mouse.time.mpse)
mouse.time.mpse %>%
mp_rrarefy() -> mpse
mpse
# larger 'chunks' means more robust, but it will become slower.
mpse %>% mp_cal_rarecurve(.abundance=RareAbundance, chunks=100, action="add")
mpse
p1 <- mpse %>%
  mp_plot_rarecurve(.rare=RareAbundanceRarecurve, .alpha="Observe")
p2 <- mpse %>%
  mp_plot_rarecurve(.rare=RareAbundanceRarecurve, .alpha=c("Observe", "ACE"))
```

Description

[Partial] [Constrained] Redundancy Analysis with MPSE or tbl_mpse object

Usage

```r
mp_cal_rda(.data, .abundance, .formula = NULL, .dim = 3, action = "add", ...)

## S4 method for signature 'MPSE'
mp_cal_rda(.data, .abundance, .formula = NULL, .dim = 3, action = "add", ...)

## S4 method for signature 'tbl_mpse'
mp_cal_rda(.data, .abundance, .formula = NULL, .dim = 3, action = "add", ...)

## S4 method for signature 'grouped_df_mpse'
mp_cal_rda(.data, .abundance, .formula = NULL, .dim = 3, action = "add", ...)
```

Arguments

- `.data` MPSE or tbl_mpse object
- `.abundance` the name of abundance to be calculated.
- `.formula` Model formula right hand side gives the constraining variables, and conditioning variables can be given within a special function ’Condition’ and keep left empty, such as ~ A + B or ~ A + Condition(B), default is NULL.
- `.dim` integer The number of dimensions to be returned, default is 3.
Calculating the samples or groups for each OTU, the result can be visualized by 'ggupset'

Description

Calculating the samples or groups for each OTU, the result can be visualized by 'ggupset'

Usage

mp_cal_upset(
  .data,
  .group,  # Add this line to the usage example
  .abundance = NULL,
  action = "add",
  force = FALSE,
  ...
)

## S4 method for signature 'MPSE'
mp_cal_upset(
  .data,
Arguments

.data MPSE or tbl_mpse object
.group the name of group to be calculated. if it is no provided, the sample will be used.
.abundance the name of otu abundance to be calculated. if it is null, the rarefied abundance will be used.
.action character, "add" joins the new information to the tibble of tbl_mpse or rowData of MPSE. "only" and "get" return a non-redundant tibble with the just new information. which is a treeData object.
.force logical whether calculate the relative abundance forcibly when the abundance is not be rarefied, default is FALSE.
... additional parameters.

Value

update object or tibble according the 'action'

Author(s)

Shuangbin Xu
See Also

[mp_plot_upset()]

Examples

data(mouse.time.mpse)
mpse <- mouse.time.mpse %>%
  mp_rrarefy() %>%
  mp_cal_upset(.abundance=RareAbundance, .group=time, action="add")
mpse
library(ggplot2)
library(ggupset)
p <- mpse %>% mp_plot_upset(.group=time, .upset=ggupsetOftime)
p
# or set action="only"
## Not run:
tbl <- mouse.time.mpse %>%
  mp_rrarefy() %>%
  mp_cal_upset(.abundance=RareAbundance, .group=time, action="only")
tbl
p2 <- tbl %>%
  ggplot(aes(x=ggupsetOftime)) +
  geom_bar() +
  ggupset::scale_x_upset() +
  ggupset::theme_combmatrix(combmatrix.label.extra_spacing=30)
## End(Not run)

mp_cal_venn

Calculating the OTU for each sample or group, the result can be visualized by 'ggVennDiagram'

Description

Calculating the OTU for each sample or group, the result can be visualized by 'ggVennDiagram'

Usage

mp_cal_venn(
  .data,
  .group,
  .abundance = NULL,
  action = "add",
  force = FALSE,
  ...
)

## S4 method for signature 'MPSE'
mp_cal_venn

mp_cal_venn(.data, .group, .abundance = NULL, action = "add", force = FALSE, ...)

## S4 method for signature 'tbl_mpse'
mp_cal_venn(.data, .group, .abundance = NULL, action = "add", force = FALSE, ...)

## S4 method for signature 'grouped_df_mpse'
mp_cal_venn(.data, .group, .abundance = NULL, action = "add", force = FALSE, ...)

Arguments

.data MPSE or tbl_mpse object

.group the name of group to be calculated. if it is no provided, the sample will be used.

.abundance the name of otu abundance to be calculated. if it is null, the rarefied abundance will be used.

.action character, "add" joins the new information to the tibble of tbl_mpse or rowData of MPSE. "only" and "get" return a non-redundant tibble with the just new information.

.force logical whether calculate the relative abundance forcibly when the abundance is not be rarefied, default is FALSE.

... additional parameters.

Value

update object or tibble according the 'action'
mp_decostand

Author(s)
Shuangbin Xu

See Also
[mp_plot_venn()]

Examples

```r
data(mouse.time.mpse)
mouse.time.mpse %>%
mp_rrarefy() %>%
mp_cal_venn(.abundance=RareAbundance, .group=time, action="add") -> mpse
mpse
p <- mpse %>% mp_plot_venn(.venn = vennOftime, .group = time)
## Not run:
# visualized by manual
library(ggplot2)
mpse %>%
  mp_extract_sample() %>%
  select(time, vennOftime) %>%
  distinct() %>%
  pull(var=vennOftime, name=time) %>%
  ggVennDiagram::ggVennDiagram()
## End(Not run)
```

mp_decostand

This Function Provides Several Standardization Methods for Community Data

Description

This Function Provides Several Standardization Methods for Community Data

Usage

```r
mp_decostand(.data, .abundance = NULL, method = "hellinger", logbase = 2, ...)
```

## S4 method for signature 'data.frame'

```r
mp_decostand(.data, .abundance = NULL, method = "hellinger", logbase = 2, ...)
```

## S4 method for signature 'MPSE'

```r
mp_decostand(.data, .abundance = NULL, method = "hellinger", logbase = 2, ...)
```

## S4 method for signature 'tbl_mpse'

```r
mp_decostand(.data, .abundance = NULL, method = "hellinger", logbase = 2, ...)
```

## S4 method for signature 'grouped_df_mpse'

```r
mp_decostand(.data, .abundance = NULL, method = "hellinger", logbase = 2, ...)
```
Arguments

.data MPSE or tbl_mpse object
.abundance the names of otu abundance to be applied standardization.
.method character the name of standardization method, it can one of 'total', 'max', 'frequency', 'normalize', 'range', 'rank', 'rrank', 'standardize' 'pa', 'chi.square', 'hellinger' and 'log', see also decostand
.logbase numeric The logarithm base used in 'method=log', default is 2.
... additional parameters, see also decostand

Value

update object

Author(s)

Shuangbin Xu

Source

mp_decostand for data.frame object is a wrapper method of vegan::decostand from the vegan package

See Also

[mp_extract_assays()] and [mp_rrarefy()]
decostand

Examples

data(mouse.time.mpse)
mouse.time.mpse %>%
mp_decostand(.abundance=Abundance, method="hellinger")
Usage

mp_diff_analysis(
  .data,
  .abundance,
  .group,
  .sec.group = NULL,
  action = "add",
  tip.level = "OTU",
  force = FALSE,
  relative = TRUE,
  taxa.class = "all",
  first.test.method = "kruskal.test",
  first.test.alpha = 0.05,
  p.adjust = "fdr",
  filter.p = "fdr",
  strict = TRUE,
  fc.method = "generalizedFC",
  second.test.method = "wilcox.test",
  second.test.alpha = 0.05,
  cl.min = 5,
  cl.test = TRUE,
  subcl.min = 3,
  subcl.test = TRUE,
  ml.method = "lda",
  normalization = 1e+06,
  ldascore = 2,
  bootnums = 30,
  sample.prop.boot = 0.7,
  ci = 0.95,
  seed = 123,
  type = "species",
  ...
)

## S4 method for signature 'MPSE'
mp_diff_analysis(
  .data,
  .abundance,
  .group,
  .sec.group = NULL,
  action = "add",
  tip.level = "OTU",
  force = FALSE,
  relative = TRUE,
  taxa.class = "all",
  first.test.method = "kruskal.test",
  first.test.alpha = 0.05,
  p.adjust = "fdr",
  filter.p = "fdr",
  strict = TRUE,
  fc.method = "generalizedFC",
  second.test.method = "wilcox.test",
  second.test.alpha = 0.05,
  cl.min = 5,
  cl.test = TRUE,
  subcl.min = 3,
  subcl.test = TRUE,
  ml.method = "lda",
  normalization = 1e+06,
  ldascore = 2,
  bootnums = 30,
  sample.prop.boot = 0.7,
  ci = 0.95,
  seed = 123,
  type = "species",
  ...
mp_diff_analysis

filter.p = "fdr",
strict = TRUE,
fc.method = "generalizedFC",
second.test.method = "wilcox.test",
second.test.alpha = 0.05,
cl.min = 5,
cl.test = TRUE,
subcl.min = 3,
subcl.test = TRUE,
ml.method = "lda",
normalization = 1e+06,
ldascore = 2,
bootnums = 30,
sample.prop.boot = 0.7,
ci = 0.95,
seed = 123,
type = "species",
...

## S4 method for signature 'tbl_mpse'
mp_diff_analysis(  
.data,
.abundance,
.group,
.sec.group = NULL,
.action = "add",
tip.level = "OTU",
.force = FALSE,
.relative = TRUE,
taxa.class = "all",
.first.test.method = "kruskal.test",
.first.test.alpha = 0.05,
p.adjust = "fdr",
filter.p = "fdr",
strict = TRUE,
fc.method = "generalizedFC",
second.test.method = "wilcox.test",
second.test.alpha = 0.05,
cl.min = 5,
cl.test = TRUE,
subcl.min = 3,
subcl.test = TRUE,
ml.method = "lda",
.normalization = 1e+06,
ldascore = 2,
bootnums = 30,
sample.prop.boot = 0.7,
## mp_diff_analysis

```r
ci = 0.95,
seed = 123,
type = "species",
```

## Arguments

- `.data` MPSE or tbl_mpse object
- `.abundance` the name of abundance to be calculated
- `.group` the group name of the samples to be calculated.
- `.sec.group` the second group name of the samples to be calculated.
- `action` character, "add" joins the new information to the taxatree (if it exists) or `rowData` and return MPSE object,"only" return a non-redundant tibble with the result of
different analysis. "get" return 'diffAnalysisClass' object.

tip.level character the taxa level to be as tip level
force logical whether to calculate the relative abundance forcibly when the abundance
is not be rarefied, default is FALSE.
relative logical whether calculate the relative abundance.
taxa.class character if taxa class is not 'all', only the specified taxa class will be identified,
default is 'all'.

first.test.method
the method for first test, option is "kruskal.test", "oneway.test", "lm", "glm", or
"glm.nb", "kruskal_test", "oneway_test" of "coin" package. default is "kruskal.test".

first.test.alpha numeric the alpha value for the first test, default is 0.05.
p.adjust character the correction method, default is "fdr", see also p.adjust function de-
fault is fdr.

filter.p character the method to filter pvalue, default is fdr, meanings the features that
fdr <= .first.test.alpha will be kept, if it is set to pvalue, meanings the features
that pvalue <= .first.test.alpha will be kept.

strict logical whether to performed in one-against-one when .sec.group is provided,
default is TRUE (strict).

fc.method character the method to check which group has more abundance for the signifi-
cantly different features, default is "generalizedFC", options are generalizedFC, compar_median, compar_mean.

second.test.method
the method for one-against-one (the second test), default is "wilcox.test" other
option is one of 'wilcox_test' of 'coin'; 'glm'; 'glm.nb' of 'MASS'.

second.test.alpha numeric the alpha value for the second test, default is 0.05.

cl.min integer the minimum number of samples per group for performing test, default is 5.

cl.test logical whether to perform test (second test) between the groups (the number of
sample of the .group should be also larger that cl.min), default is TRUE.

subcl.min integer the minimum number of samples in each second groups for performing
test, default is 3.

subcl.test logical whether to perform test for between the second groups (the .sec.group
should be provided and the number sample of each .sec.group should be larger
than subcl.min, and strict is TRUE), default is TRUE.

ml.method the method for calculating the effect size of features, option is 'lda' or 'rf'.
default is 'lda'.

normalization integer set a big number if to get more meaningful values for the LDA score, or
you can set NULL for no normalization, default is 1000000.

ldascore numeric the threshold on the absolute value of the logarithmic LDA score, de-
default is 2.

bootnums integer, set the number of bootstrap iteration for lda or rf, default is 30.
sample.prop.boot
numeric range from 0 to 1, the proportion of samples for calculating the effect
size of features, default is 0.7.

**ci**
numeric, the confidence interval of effect size (LDA or MDA), default is 0.95.

**seed**
a random seed to make the analysis reproducible, default is 123.

**type**
character type="species" meaning the abundance matrix is from the species
abundance, other option is "others", default is "species".

Value
update object according to the action argument.

Author(s)
Shuangbin Xu

Examples
```r
data(mouse.time.mpse)
mouse.time.mpse %<>% mp_rrarefy()
mouse.time.mpse
mouse.time.mpse %<>% mp_diff_analysis(.abundance=RareAbundance,
            .group=time,
            first.test.alpha=0.01,
            action="add")
```

```r
library(ggplot2)
p <- mouse.time.mpse %>% mp_plot_diff_res()
p <- p +
scale_fill_manual(
aesthetics = "fill_new", # The fill aes was renamed to "fill_new" for the abundance dotplot layer
values = c("skyblue", "orange")
) +
scale_fill_manual(
values=c("skyblue", "orange") # The LDA barplot layer
)
```

```r
### and the fill aes for hight light layer of tree was renamed to 'fill_new_new'
p <- p +
scale_fill_manual(
aesthetics = "fill_new_new",
values = c("#E41A1C", "#377EB8", "#4DAF4A",
    "#984EA3", "#FF7F00",="#FFFF33",
    "#A65628", "#F781BF",="#999999")
)
```

```r
p
```
### Description

Differential internal and tip nodes (clades) analysis for MPSE or tbl_mpse object

### Usage

```r
mp_diff_clade(.data, .abundance, .group, .sec.group = NULL, action = "add", force = FALSE, relative = TRUE, first.test.method = "kruskal.test", first.test.alpha = 0.05, p.adjust = "fdr", filter.p = "fdr", strict = TRUE, fc.method = "generalizedFC", second.test.method = "wilcox.test", second.test.alpha = 0.05, cl.min = 5, cl.test = TRUE, subcl.min = 3, subcl.test = TRUE, ml.method = "lda", normalization = 1e+06, ldascore = 2, bootnums = 30,
```

mp_diff_clade

sample.prop.boot = 0.7,
ci = 0.95,
seed = 123,
type = "species",

## S4 method for signature 'MPSE'
mp_diff_clade(
  .data,
  .abundance,
  .group,
  .sec.group = NULL,
  action = "add",
  force = FALSE,
  relative = TRUE,
  first.test.method = "kruskal.test",
  first.test.alpha = 0.05,
  p.adjust = "fdr",
  filter.p = "fdr",
  strict = TRUE,
  fc.method = "generalizedFC",
  second.test.method = "wilcoxon.test",
  second.test.alpha = 0.05,
  cl.min = 5,
  cl.test = TRUE,
  subcl.min = 3,
  subcl.test = TRUE,
  ml.method = "lda",
  normalization = 1e+06,
  ldascore = 2,
  bootnums = 30,
  sample.prop.boot = 0.7,
  ci = 0.95,
  seed = 123,
  type = "species",

## S4 method for signature 'tbl_mpse'
mp_diff_clade(
  .data,
  .abundance,
  .group,
  .sec.group = NULL,
  action = "add",
  force = FALSE,
  relative = TRUE,
first.test.method = "kruskal.test",
first.test.alpha = 0.05,
p.adjust = "fdr",
filter.p = "fdr",
strict = TRUE,
fc.method = "generalizedFC",
second.test.method = "wilcox.test",
second.test.alpha = 0.05,
cl.min = 5,
cl.test = TRUE,
subcl.min = 3,
subcl.test = TRUE,
ml.method = "lda",
normalization = 1e+06,
ldascore = 2,
bootnums = 30,
sample.prop.boot = 0.7,
cl.test = TRUE,
subcl.min = 3,
subcl.test = TRUE,
ml.method = "lda",
normalization = 1e+06,
ldascore = 2,
bootnums = 30,

## S4 method for signature 'grouped_df_mpse'
mp_diff_clade(
  .data,
  .abundance,
  .group,
  .sec.group = NULL,
  action = "add",
  force = FALSE,
  relative = TRUE,
  first.test.method = "kruskal.test",
  first.test.alpha = 0.05,
p.adjust = "fdr",
filter.p = "fdr",
strict = TRUE,
fc.method = "generalizedFC",
second.test.method = "wilcox.test",
second.test.alpha = 0.05,
cl.min = 5,
cl.test = TRUE,
subcl.min = 3,
subcl.test = TRUE,
ml.method = "lda",
(normalization = 1e+06,
ldascore = 2,
bootnums = 30,
sample.prop.boot = 0.7,
ci = 0.95,
seed = 123,
type = "species",
...)

Arguments

.data MPSE or tbl_mpse object
.abundance the name of abundance to be calculated
.group the group name of the samples to be calculated.
.sec.group the second group name of the samples to be calculated.
.action character, "add" joins the new information to the taxatree (if it exists) and otutree (if it exists) or rowData and return MPSE object,"only" return a non-redundant tibble with the result of different analysis. "get" return 'diffAnalysisClass' object.
.force logical whether to calculate the relative abundance forcibly when the abundance is not be rarefied, default is FALSE.
.relative logical whether calculate the relative abundance, default is TRUE.
.first.test.method
.first.test.alpha numeric the alpha value for the first test, default is 0.05.
p.adjust character the correction method, default is "fdr", see also p.adjust function default is fdr.
.filter.p character the method to filter pvalue, default is fdr, meanings the features that fdr <= .first.test.alpha will be kept, if it is set to pvalue, meanings the features that pvalue <= .first.test.alpha will be kept.
.strict logical whether to performed in one-against-one when .sec.group is provided, default is TRUE (strict).
.fc.method character the method to check which group has more abundance for the significantly different features, default is "generalizedFC", options are generalizedFC, compare_median, compare_mean.
.second.test.method
.second.test.alpha numeric the alpha value for the second test, default is 0.05.
.cl.min integer the minimum number of samples per group for performing test, default is 5.
.cl.test logical whether to perform test (second test) between the groups (the number of sample of the .group should be also larger that cl.min), default is TRUE.
subcl.min integer the minimum number of samples in each second groups for performing test, default is 3.

subcl.test logical whether to perform test for between the second groups (the .sec.group should be provided and the number sample of each .sec.group should be larger than subcl.min, and strict is TRUE), default is TRUE.

ml.method the method for calculating the effect size of features, option is 'lda' or 'rf'. default is 'lda'.

normalization integer set a big number if to get more meaningful values for the LDA score, or you can set NULL for no normalization, default is 1000000.

ldascore numeric the threshold on the absolute value of the logarithmic LDA score, default is 2.

bootnums integer, set the number of bootstrap iteration for lda or rf, default is 30.

sample.prop.boot numeric range from 0 to 1, the proportion of samples for calculating the effect size of features, default is 0.7.

ci numeric, the confidence interval of effect size (LDA or MDA), default is 0.95.

seed a random seed to make the analysis reproducible, default is 123.

type character type="species" meaning the abundance matrix is from the species abundance, other option is "others", default is "species".

... additional parameters

Value

update object according to the action argument.

Author(s)

Shuangbin Xu

Examples

```r
## Not run:
suppressPackageStartupMessages(library(curatedMetagenomicData))
x <- curatedMetagenomicData('ZellerG_2014.relative_abundance', dryrun=F)
x[[1]] %>% as.mpsse -> mpse
mpse.agg.clade <- mpse %>%
  mp_aggregate.clade(
    .abundance = Abundance,
    force = TRUE,
    relative = FALSE,
    action = 'add' # other option is 'get' or 'only'.
  )
mpse.agg.clade %>% mp_diff.clade(
    .abundance = Abundance,
    force = TRUE,
    relative = FALSE,
    .group = disease,
    fc.method = "compare_mean"
```

mp_dmn

Fit Dirichlet-Multinomial models to MPSE or tbl_mpse

Description

Fit Dirichlet-Multinomial models to MPSE or tbl_mpse

Usage

mp_dmn(.data, .abundance, k = 1, seed = 123, mc.cores = 2, action = "get", ...)

## S4 method for signature 'MPSE'
mp_dmn(.data, .abundance, k = 1, seed = 123, mc.cores = 2, action = "get", ...)

## S4 method for signature 'tbl_mpse'
mp_dmn(.data, .abundance, k = 1, seed = 123, mc.cores = 2, action = "get", ...)

## S4 method for signature 'grouped_df_mpse'
mp_dmn(.data, .abundance, k = 1, seed = 123, mc.cores = 2, action = "get", ...)

Arguments

.data MPSE or tbl_mpse object
.abundance The column name of OTU abundance column to be calculated.
k the number of Dirichlet components to fit, default is 1.
seed random number seed to be reproducible, default is 123.
m.cores The number of cores to use, default is 2.
action character it has three options, 'get' return a 'list' contained DMN (default),
"add" joins the new information to the input (can be extracted with mp_extract_internal_attr(name='DMN')
"only" return a non-redundant tibble with the just new information a column
contained 'DMN'.

... additional parameters, see also the mclapply and dmn.

Value

update object or other (refer to action)
mp_dmngroup

Examples

```r
## Not run:
data(mouse.time.mpse)
res <- mouse.time.mpse %>%
  mp_dmn(.abundance = Abundance,
         k = seq_len(2),
         mc.cores = 4,
         action = 'get')
res
## End(Not run)
```

mp_dmngroup

Dirichlet-Multinomial generative classifiers to MPSE or tbl_mpse

Description

Dirichlet-Multinomial generative classifiers to MPSE or tbl_mpse

Usage

```r
mp_dmngroup(.data, .abundance, .group, k = 1, action = "get", ...)
```

## S4 method for signature 'MPSE'
```r
mp_dmngroup(.data, .abundance, .group, k = 1, action = "get", ...)
```

## S4 method for signature 'tbl_mpse'
```r
mp_dmngroup(.data, .abundance, .group, k = 1, action = "get", ...)
```

## S4 method for signature 'grouped_df_mpse'
```r
mp_dmngroup(.data, .abundance, .group, k = 1, action = "get", ...)
```

Arguments

- `.data` MPSE or tbl_mpse object
- `.abundance` The column name of OTU abundance column to be calculate.
- `.group` the column name of group variable.
- `k` the number of Dirichlet components to fit, default is 1.
- `action` character it has three options, 'get' return a 'list' contained DMN (default), "add" joins the new information to the input (can be extracted with mp_extract_internal_attr(name='DMNGroup')) "only" return a non-redundant tibble with the just new information a column contained 'DMNGroup'.
- `...` additional parameters, see also the mclapply and dmngroup.

Value

update object or others (refer to action argument)
Examples

## Not run:
data(mouse.time.mpse)
mouse.time.mpse %>%
  mp_dmngroup(
    .abundance = Abundance,
    .group = time,
    k = seq_len(2),
    action = 'get'
  )

## End(Not run)

---

**mp_envfit**

*Fits an Environmental Vector or Factor onto an Ordination With MPSE or tbl_mpse Object*

**Description**

Fits an Environmental Vector or Factor onto an Ordination With MPSE or tbl_mpse Object

**Usage**

```r
mp_envfit(
  .data,
  .ord,
  .env,
  .dim = 3,
  action = "only",
  permutations = 999,
  seed = 123,
  ...
)
```

## S4 method for signature 'MPSE'

```r
mp_envfit(
  .data,
  .ord,
  .env,
  .dim = 3,
  action = "only",
  permutations = 999,
  seed = 123,
  ...
)
```

## S4 method for signature 'tbl_mpse'
mp_envfit

mp_envfit(
  .data,
  .ord,
  .env,
  .dim = 3,
  action = "only",
  permutations = 999,
  seed = 123,
  ...
)

## S4 method for signature 'grouped_df_mpse'
mp_envfit(
  .data,
  .ord,
  .env,
  .dim = 3,
  action = "only",
  permutations = 999,
  seed = 123,
  ...
)

Arguments

.data MPSE or tbl_mpse object
.ord a name of ordination, option it is DCA, NMDS, RDA, CCA.
.env the names of columns of sample group or environment information.
.dim integer The number of dimensions to be returned, default is 3.
.action character "add" joins the envfit result to internal attributes of the object, "only" return a non-redundant tibble with the envfit result. "get" return 'envfit' object can be analyzed using the related vegan funtion.
.permutations the number of permutations required, default is 999.
.seed a random seed to make the analysis reproducible, default is 123.
... additional parameters see also 'vegan::envfit'

Value

update object according action

Author(s)

Shuangbin Xu
Examples

```r
library(vegan)
data(varespec, varechem)
mpse <- MPSE(assays=list(Abundance=t(varespec)), colData=varechem)
envformula <- paste("~", paste(colnames(varechem), collapse="+"))) %>% as.formula
mpse %<>%
   mp_cal_cca(.abundance=Abundance, .formula=envformula, action="add")
mpse2 <- mpse %>%
   mp_envfit(.ord=cca,
     .env=colnames(varechem),
     permutations=9999,
     action="add")
mpse2 %>% mp_plot_ord(.ord=cca, .group=Al, .size=Mn, show.shample=TRUE, show.envfit=TRUE)
## Not run:
tbl <- mpse %>%
   mp_envfit(.ord=CCA,
     .env=colnames(varechem),
     permutations=9999,
     action="only")
tbl
library(ggplot2)
library(ggrepel)
x <- names(tbl)[grepl("^CCA1 ", names(tbl))] %>% as.symbol()
y <- names(tbl)[grepl("^CCA2 ", names(tbl))] %>% as.symbol()
p <- tbl %>%
   ggplot(aes(x=x!!x, y=y!!y)) +
   geom_point(aes(color=Al, size=Mn)) +
   geom_segment(data=dr_extract(
     name="CCA_ENVFIT_tb",
     .f=td_filter(pvals<=0.05 & label="Humdepth")
   ),
   aes(x=0, y=0, xend=CCA1, yend=CCA2),
   arrow=arrow(length = unit(0.02, "npc"))
) +
   geom_text_repel(data=dr_extract(
     name="CCA_ENVFIT_tb",
     .f=td_filter(pvals<=0.05 & label="Humdepth")
   ),
   aes(x=CCA1, y=CCA2, label=label)
) +
   geom_vline(xintercept=0, color="grey20", linetype=2) +
   geom_hline(yintercept=0, color="grey20", linetype=2) +
   theme_bw() +
   theme(panel.grid=element_blank())
p
## End(Not run)
```

---

**mp_extract_abundance**

Extracting the abundance metric from MPSE or tbl_mpse object
Description

Extracting the abundance metric from the MPSE or tbl_mpse, the 'mp_cal_abundance' must have been run with action='add'.

Usage

mp_extract_abundance(x, taxa.class = "all", topn = NULL, rmun = FALSE, ...)

## S4 method for signature 'MPSE'
mp_extract_abundance(x, taxa.class = "all", topn = NULL, rmun = FALSE, ...)

## S4 method for signature 'tbl_mpse'
mp_extract_abundance(x, taxa.class = "all", topn = NULL, rmun = FALSE, ...)

## S4 method for signature 'grouped_df_mpse'
mp_extract_abundance(x, taxa.class = "all", topn = NULL, rmun = FALSE, ...)

Arguments

x                MPSE or tbl_mpse object
taxa.class          character the name of taxonomy class level what you want to extract
topn          integer the number of the top most abundant, default is NULL.
rmun        logical whether to remove the unknown taxa, such as "g__un_xxx", default is FALSE (the unknown taxa class will be considered as 'Others').
...              additional parameters

Author(s)

Shuangbin Xu

mp_extract_assays

extract the abundance matrix from MPSE object or tbl_mpse object

Description

extract the abundance matrix from MPSE object or tbl_mpse object

Usage

mp_extract_assays(x, .abundance, byRow = TRUE, ...)

## S4 method for signature 'MPSE'
mp_extract_assays(x, .abundance, byRow = TRUE, ...)

## S4 method for signature 'tbl_mpse'
mp_extract_assays(x, .abundance, byRow = TRUE, ...)

## S4 method for signature 'grouped_df_mpse'
mp_extract_assays(x, .abundance, byRow = TRUE, ...)
## S4 method for signature 'grouped_df_mpse'
mp_extract_assays(x, .abundance, byRow = TRUE, ...)

**Arguments**

- `x` MPSE or tbl_mpse object
- `.abundance` the name of abundance to be extracted.
- `byRow` logical if it is set TRUE, 'otu X sample' shape will return, else 'sample X otu' will return.
- `...` additional parameters.

**Value**

otu abundance a data.frame object

```r
cmp_extract_dist
## S4 method for signature 'MPSE'
mp_extract_dist(x, distmethod, type = "sample", .group = NULL, ...)
## S4 method for signature 'tbl_mpse'
mp_extract_dist(x, distmethod, type = "sample", .group = NULL, ...)
## S4 method for signature 'grouped_df_mpse'
mp_extract_dist(x, distmethod, type = "sample", .group = NULL, ...)
```

**Arguments**

- `x` MPSE object or tbl_mpse object
- `distmethod` character the method of calculated distance.
- `type` character, which type distance to be extracted, 'sample' represents the distance between the samples based on feature abundance matrix, 'feature' represents the distance between the features based on feature abundance matrix, 'env' represents the the distance between the samples based on continuous environment factors, default is 'sample'.
mp_extract_feature

Description
extract the feature (OTU) information in MPSE object

Usage
mp_extract_feature(x, addtaxa = FALSE, ...)

## S4 method for signature 'MPSE'
mp_extract_feature(x, addtaxa = FALSE, ...)

## S4 method for signature 'tbl_mpse'
mp_extract_feature(x, addtaxa = FALSE, ...)

## S4 method for signature 'grouped_df_mpse'
mp_extract_feature(x, addtaxa = FALSE, ...)

Arguments
x MPSE object
addtaxa logical whether adding the taxonomy information default is FALSE.
... additional arguments

Value
tbl_df contained feature (OTU) information.
**mp_extract_internal_attr**

*Extracting the PCA, PCoA, etc results from MPSE or tbl_mpse object*

**Description**

Extracting the PCA, PCoA, etc results from MPSE or tbl_mpse object

**Usage**

```r
cmp_extract_internal_attr(x, name, ...)
```

## S4 method for signature 'MPSE'
```r
cmp_extract_internal_attr(x, name, ...)
```

## S4 method for signature 'tbl_mpse'
```r
cmp_extract_internal_attr(x, name, ...)
```

## S4 method for signature 'grouped_df_mpse'
```r
cmp_extract_internal_attr(x, name, ...)
```

**Arguments**

- `x` MPSE or tbl_mpse object
- `name` character 'PCA' or 'PCoA'
- `...` additional parameters

**Value**

prcomp or pcoa etc object

---

**mp_extract_rarecurve**

*Extract the result of mp_cal_rarecurve with action="add" from MPSE or tbl_mpse object*

**Description**

Extract the result of mp_cal_rarecurve with action="add" from MPSE or tbl_mpse object
**mp_extract_refseq**

**Usage**

```r
mp_extract_rarecurve(x, .rarecurve, ...)
```

```r
## S4 method for signature 'MPSE'
mp_extract_rarecurve(x, .rarecurve, ...)
```

```r
## S4 method for signature 'tbl_mpse'
mp_extract_rarecurve(x, .rarecurve, ...)
```

```r
## S4 method for signature 'grouped_df_mpse'
mp_extract_rarecurve(x, .rarecurve, ...)
```

**Arguments**

- `x` MPSE object or tbl_mpse object
- `.rarecurve` the column name of rarecurve after run `mp_cal_rarecurve` with action="add".
- `...` additional parameter

**Value**

rarecurve object that be be visualized by ggrarecurve

---

**mp_extract_refseq**

*Extract the representative sequences from MPSE object*

**Description**

Extract the representative sequences from MPSE object

**Usage**

```r
mp_extract_refseq(x, ...)
```

```r
## S4 method for signature 'MPSE'
mp_extract_refseq(x, ...)
```

```r
## S4 method for signature 'tbl_mpse'
mp_extract_refseq(x, ...)
```

```r
## S4 method for signature 'grouped_df_mpse'
mp_extract_refseq(x, ...)
```

**Arguments**

- `x` MPSE object
- `...` additional parameters, meaningless now.
mp_extract_sample  

**extract the sample information in MPSE object**

**Description**

extract the sample information in MPSE object

**Usage**

```r
mp_extract_sample(x, ...)
```

## S4 method for signature 'MPSE'

```r
mp_extract_sample(x, ...)
```

## S4 method for signature 'tbl_mpse'

```r
mp_extract_sample(x, ...)
```

## S4 method for signature 'grouped_df_mpse'

```r
mp_extract_sample(x, ...)
```

**Arguments**

- x  
  MPSE object
- ...  
  additional arguments

**Value**

tbl_df contained sample information.

mp_extract_tree  

**extract the taxonomy tree in MPSE object**

**Description**

extract the taxonomy tree in MPSE object

**Usage**

```r
mp_extract_tree(x, type = "taxatree", tip.level = "OTU", ...)
```

## S4 method for signature 'MPSE'

```r
mp_extract_tree(x, type = "taxatree", tip.level = "OTU", ...)
```

## S4 method for signature 'tbl_mpse'

```r
mp_extract_tree(x, type = "taxatree", tip.level = "OTU", ...)
```
mp_filter_taxa

## S4 method for signature 'grouped_df_mpse'
mp_extract_tree(x, type = "taxatree", tip.level = "OTU", ...)

mp_extract_taxatree(x, tip.level = "OTU", ...)

mp_extract_otutree(x, ...)

Arguments

x MPSE object
type character taxatree or otutree
tip.level character This argument will keep the nodes belong to the tip.level as tip nodes when type is taxatree, default is OTU, which will return the taxa tree with OTU level as tips.
...
additional arguments

Value
taxatree treedata object

mp_filter_taxa Filter OTU (Features) By Abundance Level

Description

Filter OTU (Features) By Abundance Level

Usage

mp_filter_taxa(
  .data,
  .abundance = NULL,
  min.abun = 0,
  min.prop = 0.05,
  include.lowest = FALSE,
  ...
)

## S4 method for signature 'MPSE'
mp_filter_taxa(
  .data,
  .abundance = NULL,
  min.abun = 0,
  min.prop = 0.05,
  include.lowest = FALSE,
mp_filter_taxa

## S4 method for signature 'tbl_mpse'
mp_filter_taxa(
  .data,
  .abundance = NULL,
  min.abun = 0,
  min.prop = 0.05,
  include.lowest = FALSE,
  ...
)

## S4 method for signature 'grouped_df_mpse'
mp_filter_taxa(
  .data,
  .abundance = NULL,
  min.abun = 0,
  min.prop = 0.05,
  include.lowest = FALSE,
  ...
)

### Arguments

- **.data** MPSE or tbl_mpse or grouped_df_mpse object.
- **.abundance** the column names of abundance, default is NULL, meaning the 'Abundance' column.
- **min.abun** numeric minimum abundance required for each one sample default is 0 (.abundance=Abundance or NULL), meaning the abundance of OTU (Features) for each one sample should be >= 0.
- **min.prop** numeric minimum proportion of samples that contains the OTU (Features) when min.prop larger than 1, meaning the minimum number of samples that contains the OTU (Features).
- **include.lowest** logical whether include the lower boundary of min.abun default is FALSE (> min.abun), if it is TRUE, meaning (>= min.abun).
- **...** additional parameters, meaningless now.

### Author(s)

Shuangbin Xu

### Examples

data(mouse.time.mpse)
mouse.time.mpse %>% mp_filter_taxa(.abundance=Abundance, min.abun=1, min.prop=1)

# For tbl_mpse object.
mouse.time.mpse %>% as_tibble %>% mp_filter_taxa(.abundance=Abundance, min.abun=1, min.prop=1)
# This also can be done using group_by, filter of dplyr.
mouse.time.mpse %>%
dplyr::group_by(OTU) %>%
dplyr::filter(sum(Abundance>=1)>=1)

---

**mp_fortify**

**Description**

Fortify a model with data in MicrobiotaProcess

**Usage**

```r
mp_fortify(model, ...) 
```

**Arguments**

- `model` object
- `...` additional parameters

**Value**

data frame or tbl_df object

---

**mp_import_biom**

building MPSE object from biom-format file.

**Description**

building MPSE object from biom-format file.

**Usage**

```r
mp_import_biom( 
  biomfilename, 
  mapfilename = NULL, 
  otutree = NULL, 
  refseq = NULL, 
  ... 
)
```
**Arguments**

<table>
<thead>
<tr>
<th>Argument</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>biomfilename</td>
<td>character the biom-format file path.</td>
</tr>
<tr>
<td>mapfilename</td>
<td>character, the file contained sample information, the tsv format, default is NULL.</td>
</tr>
<tr>
<td>otutree</td>
<td>treedata, phylo or character, the file contained reference sequences, or treedata object, which is the result parsed by functions of treeio, default is NULL.</td>
</tr>
<tr>
<td>refseq</td>
<td>XStringSet or character, the file contained the representation sequence file or XStringSet class to store the representation sequence, default is NULL.</td>
</tr>
<tr>
<td>...</td>
<td>additional parameter, which is meaningless now.</td>
</tr>
</tbody>
</table>

**Value**

MPSE-class

---

**mp_import_humann_regroup**

*Import function to load the output of human_regroup_table in HUMAnN.*

---

**Description**

Import function to load the output of human_regroup_table in HUMAnN.

**Usage**

```r
mp_import_humann_regroup(
  profile,
  mapfilename = NULL,
  rm.unknown = TRUE,
  keep.contribute.abundance = FALSE,
  ...
)
```

**Arguments**

<table>
<thead>
<tr>
<th>Argument</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>profile</td>
<td>the output file (text format) of human_regroup_table in HUMAnN.</td>
</tr>
<tr>
<td>mapfilename</td>
<td>the sample information file or data.frame,</td>
</tr>
<tr>
<td>rm.unknown</td>
<td>logical whether remove the unmapped and ungrouped features.</td>
</tr>
<tr>
<td>keep.contribute.abundance</td>
<td>logical whether keep the abundance of contributed taxa, default is FALSE, it will consume more memory if it set to TRUE.</td>
</tr>
<tr>
<td>...</td>
<td>additional parameters, meaningless now.</td>
</tr>
</tbody>
</table>

**Author(s)**

Shuangbin Xu
mp_import_metaphlan

Import function to load the output of MetaPhlAn.

Description

Import function to load the output of MetaPhlAn.

Usage

mp_import_metaphlan(
  profile,
  mapfilename = NULL,
  treefile = NULL,
  linenum = NULL,
  ...
)

Arguments

- **profile**: the output file (text format) of MetaPhlAn.
- **mapfilename**: the sample information file or data.frame, default is NULL.
- **treefile**: the path of MetaPhlAn tree file (mpa_v30_CHOCOPhlan_201901_species_tree.nwk), default is NULL.
- **linenum**: a integer, sometimes the output file of MetaPhlAn (<3) contained the sample information in the first several lines. The linenum should be required. for example:

  group A A A A B B B B
  subgroup A1 A1 A2 A2 B1 B1 B2 B2
  subject S1 S2 S3 S4 S5 S6 S7 S8
  ...

  the linenum should be set to 3.

- **sampleid**: A1 A2 A3 A4 A5
  Bacteria 99 99 99 99 99
  ...

  The linenum should be set to 1.

  ... additional parameters, meaningless now.

Details

When the output abundance of MetaPhlAn is relative abundance, the force of mp_cal_abundance should be set to TRUE, and the relative of mp_cal_abundance should be set to FALSE. Because the abundance profile will be rarefied in the default (force=FALSE), which requires the integer (count) abundance, then the relative abundance will be calculated in the default (relative=TRUE).
Author(s)
Shuangbin Xu

Examples

```r
file1 <- system.file("extdata/MetaPhlAn", "metaphlan_test.txt", package="MicrobiotaProcess")
sample.file <- system.file("extdata/MetaPhlAn", "sample_test.txt", package="MicrobiotaProcess")
readLines(file1, n=3) %>% writeLines()
mpse1 <- mp_import_metaphlan(profile=file1, mapfilename=sample.file)
mpse1
```

mp_import_qiime

Import function to load the output of qiime.

Description

The function was designed to import the output of qiime and convert them to MPSE class.

Usage

```r
mp_import_qiime(
  otufilename,
  mapfilename = NULL,
  otutree = NULL,
  refseq = NULL,
  ...
)
```

Arguments

- `otufilename`: character, the file contained otu table, the ouput of qiime.
- `mapfilename`: character, the file contained sample information, the tsv format, default is NULL.
- `otutree`: treedata, phylo or character, the file contained reference sequences, or treedata object, which is the result parsed by functions of treeio, default is NULL.
- `refseq`: XStringSet or character, the file contained the representation sequence file or XStringSet class to store the representation sequence, default is NULL.
- `...`, additional parameters.

Value

MPSE-class.

Author(s)

Shuangbin Xu
**mp_mantel**

*Mantel and Partial Mantel Tests for MPSE or tbl_mpse Object*

**Description**

Mantel and Partial Mantel Tests for MPSE or tbl_mpse Object

**Usage**

```r
mp_mantel(
  .data,
  .abundance,
  .y.env,
  .z.env = NULL,
  distmethod = "bray",
  distmethod.y = "euclidean",
  distmethod.z = "euclidean",
  method = "pearson",
  permutations = 999,
  action = "get",
  seed = 123,
  scale.y = FALSE,
  scale.z = FALSE,
  ...
)
```

## S4 method for signature 'MPSE'

```r
mp_mantel(
  .data,
  .abundance,
  .y.env,
  .z.env = NULL,
  distmethod = "bray",
  distmethod.y = "euclidean",
  distmethod.z = "euclidean",
  method = "pearson",
  permutations = 999,
  action = "get",
  seed = 123,
  scale.y = FALSE,
  scale.z = FALSE,
  ...
)
```

## S4 method for signature 'tbl_mpse'

```r
mp_mantel(
  .data,
```
### Arguments

- **.data** MPSE or tbl_mpse object
- **.abundance** the name of otu abundance to be calculated
- **.y.env** the column names of continuous environment factors to perform Mantel statistic, it is required.
- **.z.env** the column names of continuous environment factors to perform Partial Mantel statistic based on this, default is NULL.
- **distmethod** character the method to calculate distance based on .abundance.
- **distmethod.y** character the method to calculate distance based on .y.env.
- **distmethod.z** character the method of calculated distance based on .z.env
- **method** character Correlation method, options is "pearson", "spearman" or "kendall"
- **permutations** the number of permutations required, default is 999.
Analysis of Multi Response Permutation Procedure (MRPP) with MPSE or tbl_mpse object

Description
Analysis of Multi Response Permutation Procedure (MRPP) with MPSE or tbl_mpse object

Usage

mp_mrpp(
  .data,  
  .abundance,  
  .group,  
  distmethod = "bray",  
  action = "add",  
  permutations = 999,  
  seed = 123,  
  ...
)
## S4 method for signature 'MPSE'

```
mp_mrpp(
  .data, 
  .abundance, 
  .group, 
  distmethod = "bray", 
  action = "add", 
  permutations = 999, 
  seed = 123, 
  ...
)
```

## S4 method for signature 'tbl_mpse'

```
mp_mrpp(
  .data, 
  .abundance, 
  .group, 
  distmethod = "bray", 
  action = "add", 
  permutations = 999, 
  seed = 123, 
  ...
)
```

## S4 method for signature 'grouped_df_mpse'

```
mp_mrpp(
  .data, 
  .abundance, 
  .group, 
  distmethod = "bray", 
  action = "add", 
  permutations = 999, 
  seed = 123, 
  ...
)
```

### Arguments

- `.data` (MPSE or tbl_mpse object)
- `.abundance` (the name of abundance to be calculated).
- `.group` (The name of the column of the sample group information).
- `.distmethod` (character the method to calculate pairwise distances, default is 'bray').
- `.action` (character "add" joins the ANOSIM result to internal attribute of the object, "only" return a tibble contained the statistic information of MRPP analysis, and "get" return 'mrpp' object can be analyzed using the related vegan function).
permutations  the number of permutations required, default is 999.
seed        a random seed to make the MRPP analysis reproducible, default is 123.
...         additional parameters see also ’mrpp’ of vegan.

Value
update object according action argument

Author(s)
Shuangbin

Examples

data(mouse.time.mpse)
mouse.time.mpse %>%
  mp_decostand(.abundance=Abundance) %>%
  mp_mrpp(.abundance=hellinger,
          .group=time,
          distmethod="bray",
          permutations=999, # for more robust, set it to 9999.
          action="get")

Description
plotting the abundance of taxa via specified taxonomy class

Usage

mp_plot_abundance(  
  .data,  
  .abundance = NULL,  
  .group = NULL,  
  taxa.class = NULL,  
  topn = 10,  
  relative = TRUE,  
  force = FALSE,  
  plot.group = FALSE,  
  geom = "flowbar",  
  feature.dist = "bray",  
  feature.hclust = "average",  
  sample.dist = "bray",  
  sample.hclust = "average",  
  .sec.group = NULL,  
  rmun = FALSE,
rm.zero = TRUE,
order.by.feature = FALSE,

)

## S4 method for signature 'MPSE'
mp_plot_abundance(
  .data,
  .abundance = NULL,
  .group = NULL,
  taxa.class = NULL,
  topn = 10,
  relative = TRUE,
  force = FALSE,
  plot.group = FALSE,
  geom = "flowbar",
  feature.dist = "bray",
  feature.hclust = "average",
  sample.dist = "bray",
  sample.hclust = "average",
  .sec.group = NULL,
  rmun = FALSE,
  rm.zero = TRUE,
  order.by.feature = FALSE,

)

## S4 method for signature 'tbl_mpse'
mp_plot_abundance(
  .data,
  .abundance = NULL,
  .group = NULL,
  taxa.class = NULL,
  topn = 10,
  relative = TRUE,
  force = FALSE,
  plot.group = FALSE,
  geom = "flowbar",
  feature.dist = "bray",
  feature.hclust = "average",
  sample.dist = "bray",
  sample.hclust = "average",
  .sec.group = NULL,
  rmun = FALSE,
  rm.zero = TRUE,
  order.by.feature = FALSE,

)
## S4 method for signature 'grouped_df_mpse'

```r
mp_plot_abundance(
  .data,
  .abundance = NULL,
  .group = NULL,
  taxa.class = NULL,
  topn = 10,
  relative = TRUE,
  force = FALSE,
  plot.group = FALSE,
  geom = "flowbar",
  feature.dist = "bray",
  feature.hclust = "average",
  sample.dist = "bray",
  sample.hclust = "average",
  .sec.group = NULL,
  rmun = FALSE,
  rm.zero = TRUE,
  order.by.feature = FALSE,
  ...
)
```

### Arguments

- `.data` MPSE object or tbl_mpse object
- `.abundance` the column name of abundance to be plotted.
- `.group` the column name of group to be calculated and plotted, default is NULL.
- `taxa.class` name of taxonomy class, default is NULL, meaning the Phylum class will be plotted.
- `topn` integer the number of the top most abundant, default is 10.
- `relative` logical whether calculate the relative abundance and plotted.
- `force` logical whether calculate the relative abundance forcibly when the abundance is not be rarefied, default is FALSE.
- `plot.group` logical whether plotting the abundance of specified taxa.class taxonomy with group not sample level, default is FALSE.
- `geom` character which type plot, options is 'flowbar' 'bar' and 'heatmap', default is 'flowbar'.
- `feature.dist` character the method to calculate the distance between the features, based on the `.abundance` of `taxa.class`, default is 'bray', options refer to the 'distmethod' of [mp_cal_dist()](#) (except unifrac related).
- `feature.hclust` character the agglomeration method for the features, default is 'average', options are 'single', 'complete', 'average', 'ward.D', 'ward.D2', 'centroid' 'median' and 'mcquitty'.

---

---
sample.dist  character the method to calculate the distance between the samples based on the '.abundance' of 'taxa.class', default is 'bray', options refer to the 'distmethod' of [mp_cal_dist()] (except unifrac related).

sample.hclust  character the agglomeration method for the samples, default is 'average', options are 'single', 'complete', 'average', 'ward.D', 'ward.D2', 'centroid' 'median' and 'mcquitty'.

.sec.group  the column name of second group to be plotted with nested facet, default is NULL, this argument will be deprecated in the next version.

rmun  logical whether to group the unknown taxa to Others category, such as "g__un_xxx", default is FALSE, meaning do not group them to Others category.

rm.zero  logical whether to display the zero abundance, which only work with geom='heatmap' default is TRUE.

order.by.feature  character adjust the order of axis x, default is FALSE, if it is NULL or TRUE, meaning the order of axis.x will be visualizing with the order of samples by highest abundance of features.

...  additional parameters, when the geom = "flowbar", it can specify the parameters of 'geom_stratum' of 'ggalluvial', when the geom = 'bar', it can specify the parameters of 'geom_bar' of 'ggplot2', when the geom = "heatmap", it can specify the parameter of 'geom_tile' of 'ggplot2'.

Author(s)

Shuangbin Xu

Examples

## Not run:
data(mouse.time.mpse)
mouse.time.mpse %<>% mp_rrarefy()  
mouse.time.mpse %<>% mp_cal_abundance(.abundance=rareabundance, action="add") %>%  
mouse.time.mpse %<>% mp_cal_abundance(.abundance=rareabundance, .group=time, action="add")
mouse.time.mpse  
p1 <- mouse.time.mpse %>%  
  mp_plot_abundance(.abundance=relrareabundancebysample,  
  .group=time,  
  taxa.class="Phylum",  
  topn=20)  
p2 <- mouse.time.mpse %>%  
  mp_plot_abundance(.abundance = abundance,  
  taxa.class = Phylum,  
  topn = 20,  
  relative = FALSE,  
  force = TRUE  
)  
p3 <- mouse.time.mpse %>%
mp_plot_alpha

mp_plot_abundance(.abundance = RareAbundance,
   .group = time,
   taxa.class = Phylum,
   topn = 20,
   relative = FALSE,
   force = TRUE
)

p4 <- mouse.time.mpse %>%
  mp_plot_abundance(.abundance = RareAbundance,
   .group = time,
   taxa.class = Phylum,
   topn = 20,
   relative = FALSE,
   force = TRUE,
   plot.group = TRUE
)

## End(Not run)

mp_plot_alpha

Plotting the alpha diversity between samples or groups.

Description

Plotting the alpha diversity between samples or groups.

Usage

mp_plot_alpha(
   .data,
   .group,
   .alpha = c("Observe", "Shannon"),
   test = "wilcox.test",
   comparisons = NULL,
   step_increase = 0.05,
   ...
)

## S4 method for signature 'MPSE'
mp_plot_alpha(
   .data,
   .group,
   .alpha = c("Observe", "Shannon"),
   test = "wilcox.test",
   comparisons = NULL,
   step_increase = 0.05,
   ...
)
## S4 method for signature 'tbl_mpse'
mp_plot_alpha(
  .data,
  .group,
  .alpha = c("Observe", "Shannon"),
  test = "wilcox.test",
  comparisons = NULL,
  step_increase = 0.05,
  ...
)

## S4 method for signature 'grouped_df_mpse'
mp_plot_alpha(
  .data,
  .group,
  .alpha = c("Observe", "Shannon"),
  test = "wilcox.test",
  comparisons = NULL,
  step_increase = 0.05,
  ...
)

### Arguments
- `.data` MPSE or tbl_mpse object
- `.group` the column name of sample group information
- `.alpha` the column name of alpha index after run mp_cal_alpha or mp_cal_pd_metric.
- `test` the name of the statistical test, default is 'wilcox.test'
- `comparisons` A list of length-2 vectors. The entries in the vector are either the names of 2 values on the x-axis or the 2 integers that correspond to the index of the columns of interest, default is NULL, meaning it will be calculated automatically with the names in the .group.
- `step_increase` numeric vector with the increase in fraction of total height for every additional comparison to minimize overlap, default is 0.05.
- `...` additional parameters, see also `geom_signif`

### Author(s)
Shuangbin Xu

### Examples
```r
## Not run:
data(mouse.time.mpse)
mpse <- mouse.time.mpse %>%
  mp_rrarefy() %>%
  mp_cal_alpha(.abundance=RareAbundance)
```
mp_plot_diff_boxplot

```r
mpse
p <- mpse %>%
  mp_plot_alpha(.group=time, .alpha=c(Observe, Shannon, Pielou))
p
## End(Not run)
```

---

**mp_plot_diff_boxplot** displaying the differential result contained abundance and LDA with boxplot (abundance) and error bar (LDA).

---

**Description**

displaying the differential result contained abundance and LDA with boxplot (abundance) and error bar (LDA).

**Usage**

```r
mp_plot_diff_boxplot(
  .data,
  .group,
  .size = 2,
  errorbar.xmin = NULL,
  errorbar.xmax = NULL,
  point.x = NULL,
  taxa.class = "all",
  group.abun = FALSE,
  removeUnknown = FALSE,
  ...
)
```

```r
## S4 method for signature 'MPSE'
mp_plot_diff_boxplot(
  .data,
  .group,
  .size = 2,
  errorbar.xmin = NULL,
  errorbar.xmax = NULL,
  point.x = NULL,
  taxa.class = "all",
  group.abun = FALSE,
  removeUnknown = FALSE,
  ...
)
```

```r
## S4 method for signature 'tbl_mpse'
mp_plot_diff_boxplot(
```
mp_plot_diff_boxplot

Arguments

.data MPSE or tbl_mpse after run mp_diff_analysis with 'action="add"'.
.group the column name for mapping the different color.
.size the column name for mapping the size of points or numeric, default is 2.
.errorbar.xmin the column name for 'xmin' mapping of error barplot layer, default is NULL.
.errorbar.xmax the column name for 'xmax' mapping of error barplot layer, default is NULL.
.point.x the column name for 'x' mapping of point layer (right panel), default is NULL.
taxa.class the taxonomy class features will be displayed, default is 'all'.
.group.abun logical whether plot the abundance in each group with bar plot, default is FALSE.
.removeUnknown logical whether mask the unknown taxonomy information but differential species, default is FALSE.

Examples

data(mouse.time.mpse)
mouse.time.mpse %<>% mp_rrarefy()
mouse.time.mpse
mouse.time.mpse %<>%
mp_diff_analysis(.abundance=RareAbundance, 
    .group=time, 
    first.test.alpha=0.01, 
    action="add")

library(ggplot2)
p1 <- mouse.time.mpse %>%
    mp_plot_diff_boxplot(.group = time) %>%
    set_diff_boxplot_color(
        values = c("deepskyblue", "orange"),
        guide = guide_legend(title=NULL)
    )
p1

p2 <- mouse.time.mpse %>%
    mp_plot_diff_boxplot(
        taxa.class = c(Genus, OTU),
        group.abun = TRUE,
        removeUnknown = TRUE,
    ) %>%
    set_diff_boxplot_color(
        values = c("deepskyblue", "orange"),
        guide = guide_legend(title=NULL)
    )
p2

mp_plot_diff_cladogram

Visualizing the result of mp_diff_analysis with cladogram.

**Description**

Visualizing the result of mp_diff_analysis with cladogram.

**Usage**

```r
mp_plot_diff_cladogram(
    .data, 
    .group, 
    .size = "pvalue", 
    taxa.class, 
    removeUnknown = FALSE, 
    layout = "radial", 
    hilight.alpha = 0.3, 
    hilight.size = 0.2, 
    bg.tree.size = 0.15, 
    bg.tree.color = 
        "#bed0d1", 
    bg.point.color = 
        "#bed0d1", 
    bg.point.fill = "white", 
    bg.point.stroke = 0.2, 
    bg.point.size = 2,
```
mp_plot_diff_cladogram

label.size = 2.6,
tip.annot = TRUE,
as.tiplab = TRUE,
...
)

Arguments

.data MPSE object or treedata which was from the taxatree slot after running the 'mp_diff_analysis'.
.group the column name for mapping the different color.
.size the column name for mapping the size of points, default is 'pvalue'.
taxa.class the taxonomy class name will be replaced shorthand, default is the one level above ‘OTU’.
.removeUnknown logical, whether mask the unknown taxonomy information but differential species, default is FALSE.
.layout character, the layout of tree, default is 'radial', see also the 'layout' of 'ggtree'.
hilight.alpha numeric, the transparency of high light clade, default is 0.3.
hilight.size numeric, the margin thickness of high light clade, default is 0.2.
bg.tree.size numeric, the line size (width) of tree, default is 0.15.
bg.tree.color character, the line color of tree, default is '#bed0d1'.
b.g.point.color character, the color of margin of background node points of tree, default is '#bed0d1'.
b.g.point.fill character, the point fill (since point shape is 21) of background nodes of tree, default is 'white'.
b.g.point.stroke numeric, the margin thickness of point of background nodes of tree, default is 0.2.
b.g.point.size numeric, the point size of background nodes of tree, default is 2.
label.size numeric, the label size of differential taxa, default is 2.6.
tip.annot logical whether to replace the differential tip labels with shorthand, default is TRUE.
as.tiplab logical, whether to display the differential tip labels with 'geom_tiplab' of 'ggtree', default is TRUE, if it is FALSE, it will use 'geom_text_repel' of 'ggrepel'.

Details

The color scale of differential group can be designed by 'scale_fill_diff_cladogram'
Examples

```r
## Not run:
data(mouse.time.mpse)
mouse.time.mpse %<>%
   mp_rrarefy()
mouse.time.mpse
mouse.time.mpse %<>%
   mp_diff_analysis(.abundance=RareAbundance,
                   .group=time,
                   first.test.alpha=0.01,
                   action="add")
# visualizing the differential taxa with cladogram
library(ggplot2)
f <- mouse.time.mpse %>%
   mp_plot_diff_cladogram(
     label.size = 2.5,
     hilight.alpha = .3,
     bg.tree.size = .5,
     bg.point.size = 2,
     bg.point.stroke = .25
   ) +
   scale_fill_diff_cladogram(
     values = c('skyblue', 'orange')
   ) +
   scale_size_continuous(range = c(1, 4))
f
## End(Not run)
```

### Description

`mp_plot_diff_manhattan`

```r
displaying the differential result contained abundance and LDA with manhattan plot.
```

### Usage

```r
mp_plot_diff_manhattan(
  .data,
  .group,
  .y = "fdr",
  .size = 2,
  taxa.class = "OTU",
  anno.taxa.class = NULL,
  removeUnknown = FALSE,
  ...
)```
## S4 method for signature 'MPSE'
mp_plot_diff_manhattan(
  .data,
  .group,
  .y = "fdr",
  .size = 2,
  taxa.class = "OTU",
  anno.taxa.class = NULL,
  removeUnknown = FALSE,
  ...
)

## S4 method for signature 'tbl_mpse'
mp_plot_diff_manhattan(
  .data,
  .group,
  .y = "fdr",
  .size = 2,
  taxa.class = "OTU",
  anno.taxa.class = NULL,
  removeUnknown = FALSE,
  ...
)

## S4 method for signature 'grouped_df_mpse'
mp_plot_diff_manhattan(
  .data,
  .group,
  .y = "fdr",
  .size = 2,
  taxa.class = "OTU",
  anno.taxa.class = NULL,
  removeUnknown = FALSE,
  ...
)

### Arguments

- `.data`: MPSE or tbl_mpse after run `mp_diff_analysis` with `action="add"`.
- `.group`: the column name for mapping the different color.
- `.y`: the column name for mapping the y axis, default is `"fdr"`.
- `.size`: the column name for mapping the size of points or numeric, default is 2.
- `.taxa.class`: the taxonomy class features will be displayed, default is `"OTU"`.
- `.anno.taxa.class`: the taxonomy class to annotate the sign taxa with color, default is `"Phylum"` if `"taxatree"` is not empty.
removeUnknown logical whether mask the unknown taxonomy information but differential species, default is FALSE.

... additional params, see also the 'geom_text_repel' and 'geom_point'.

Examples

data(mouse.time.mpse)
mouse.time.mpse %<>% mp_rrarefy()
mouse.time.mpse %<>% mp_diff_analysis(.abundance=RareAbundance,
.group=time,
.first.test.alpha=0.01,
.action="add")
p <- mouse.time.mpse %>%
  mp_plot_diff_manhattan(
    .group = Sign_time,
    .y = fdr,
    .size = 2,
    taxa.class = OTU,
    anno.taxa.class = Phylum,
  )

Description

The visualization of result of mp_diff_analysis

Usage

mp_plot_diff_res(
  .data, 
  .group, 
  layout = "radial", 
  tree.type = "taxatree", 
  taxa.class = NULL, 
  barplot.x = NULL, 
  point.size = NULL, 
  sample.num = 50, 
  tiplab.size = 2, 
  offset.abun = 0.04, 
  pwidth.abun = 0.8, 
  offset.effsize = 0.3, 
  pwidth.effsize = 0.5, 
  group.abun = FALSE,
tiplab.linetype = 3,
...

## S4 method for signature 'MPSE'
mp_plot_diff_res(
  .data,
  .group,
  layout = "radial",
  tree.type = "taxatree",
  .taxa.class = NULL,
  barplot.x = NULL,
  point.size = NULL,
  sample.num = 50,
  tiplab.size = 2,
  offset.abun = 0.04,
  pwidth.abun = 0.8,
  offset.effsize = 0.3,
  pwidth.effsize = 0.5,
  group.abun = FALSE,
  tiplab.linetype = 3,
  ...
)

## S4 method for signature 'tbl_mpse'
mp_plot_diff_res(
  .data,
  .group,
  layout = "radial",
  tree.type = "taxatree",
  .taxa.class = NULL,
  barplot.x = NULL,
  point.size = NULL,
  sample.num = 50,
  tiplab.size = 2,
  offset.abun = 0.04,
  pwidth.abun = 0.8,
  offset.effsize = 0.3,
  pwidth.effsize = 0.5,
  group.abun = FALSE,
  tiplab.linetype = 3,
  ...
)

## S4 method for signature 'grouped_df_mpse'
mp_plot_diff_res(
  .data,
  .group,
```r
layout = "radial",
.tree.type = "taxatree",
.taxa.class = NULL,
.barplot.x = NULL,
.point.size = NULL,
sample.num = 50,
tiplab.size = 2,
offset.abun = 0.04,
pwidth.abun = 0.8,
offset.effsize = 0.3,
pwidth.effsize = 0.5,
group.abun = FALSE,
tiplab.linetype = 3,
... )
```

Arguments

- `.data` MPSE or tbl_mpse after run mp_diff_analysis with action="add"
- `.group` the column name for mapping the different color, default is the column name has 'Sign_' prefix, which contains the enriched group name, but the insignificant should be NA.
- `layout` the type of tree layout, should be one of "rectangular", "roundrect", "ellipse", "circular", "slanted", "radial", "inward_circular".
- `tree.type` one of 'taxatree' and 'otutree', taxatree is the taxonomy class tree 'otutree' is the phylogenetic tree built with the representative sequences.
- `.taxa.class` character the name of taxonomy class level, default is NULL, meaning it will extract the phylum annotation automatically.
- `barplot.x` the column name of continuous value mapped to barplot, default is NULL, meaning the 'LDAmean' will be used internally.
- `point.size` the column name of continuous value mapped to the size of point in the tree, default is NULL, meaning the 'fdr' will be used internally.
- `sample.num` integer when it is smaller than the sample number of `.data`, the abundance of `.group` will replace the abundance of sample, default is 50.
- `tiplab.size` numeric the size of tiplab, default is 2.
- `offset.abun` numeric the gap (width) (relative width to tree) between the tree and abundance panel, default is 0.04.
- `pwidth.abun` numeric the panel width (relative width to tree) of abundance panel, default is 0.3.
- `offset.effsize` numeric the gap (width) (relative width to tree) between the tree and effect size panel, default is 0.3.
- `pwidth.effsize` numeric the panel width (relative width to tree) of effect size panel, default is 0.5.
- `group.abun` logical whether to display the relative abundance of group instead of sample, default is FALSE.
mp_plot_dist

Plotting the distance between the samples with heatmap or boxplot.

Description
Plotting the distance between the samples with heatmap or boxplot.

Usage

mp_plot_dist(
  .data,
  .distmethod,
  .group = NULL,
  group.test = FALSE,
  hclustmethod = "average",
  test = "wilcox.test",
  comparisons = NULL,
  step_increase = 0.1,
  ...
)

## S4 method for signature 'MPSE'
mp_plot_dist(
  .data,
  .distmethod,
  .group = NULL,
  group.test = FALSE,
  hclustmethod = "average",
  test = "wilcox.test",
  comparisons = NULL,
  step_increase = 0.1,
  ...
)

## S4 method for signature 'tbl_mpse'
mp_plot_dist(
  .data,
  .distmethod,
  .group = NULL,
  group.test = FALSE,
  hclustmethod = "average",
  test = "wilcox.test",
  comparisons = NULL,
## S4 method for signature 'grouped_df_mpse'

```r
mp_plot_dist(
  .data,
  .distmethod,
  .group = NULL,
  group.test = FALSE,
  hclustmethod = "average",
  test = "wilcox.test",
  comparisons = NULL,
  step_increase = 0.1,
  ...
)
```

### Arguments

- **.data**
  - the MPSE or tbl_mpse object after \([\text{mp_cal_dist}()]\) is performed with action="add"

- **.distmethod**
  - the column names of distance of samples, it will generate after \([\text{mp_cal_dist}()]\) is performed.

- **.group**
  - the column names of group, default is NULL, when it is not provided the heatmap of distance between samples will be returned. If it is provided and group.test is TRUE, the comparisons boxplot of distance between the group will be returned, but when group.test is FALSE, the heatmap of distance between samples with group information will be returned.

- **group.test**
  - logical default is FALSE, see the .group argument.

- **hclustmethod**
  - character the method of \([\text{hclust}()]\), default is 'average' (= UPGMA).

- **test**
  - the name of the statistical test, default is 'wilcox.test'

- **comparisons**
  - A list of length-2 vectors. The entries in the vector are either the names of 2 values on the x-axis or the 2 integers that correspond to the index of the columns of interest, default is NULL, meaning it will be calculated automatically with the names in the .group.

- **step_increase**
  - numeric vector with the increase in fraction of total height for every additional comparison to minimize overlap, default is 0.1.

- **...**
  - additional parameters, see also \([\text{geom_signif}()]\)

### Author(s)

Shuangbin Xu

### See Also

- \([\text{mp_cal_dist}()]\) and \([\text{mp_extract_dist}()]\)
### Examples

```r
## Not run:
data(mouse.time.mpse)
mouse.time.mpse %<>% mp_decostand(.abundance=Abundance)
mouse.time.mpse
mp_cal_dist(.abundance=hellinger, distmethod="bray")
mouse.time.mpse
p1 <- mouse.time.mpse %>%
  mp_plot_dist(.distmethod=bray)
p2 <- mouse.time.mpse %>%
  mp_plot_dist(.distmethod=bray, .group=time, group.test=TRUE)
p3 <- mouse.time.mpse %>%
  mp_plot_dist(.distmethod=bray, .group=time)

## End(Not run)
```

---

### mp_plot_ord

**Plotting the result of PCA, PCoA, CCA, RDA, NDMS or DCA**

**Description**

Plotting the result of PCA, PCoA, CCA, RDA, NDMS or DCA

**Usage**

```r
mp_plot_ord(
  .data,
  .ord,
  .dim = c(1, 2),
  .group = NULL,
  .starshape = 15,
  .size = 2,
  .alpha = 1,
  .color = "black",
  starstroke = 0.5,
  show.side = TRUE,
  show.adonis = FALSE,
  ellipse = FALSE,
  show.sample = FALSE,
  show.envfit = FALSE,
  p.adjust = NULL,
  filter.envfit = FALSE,
  ...
)
```

## S4 method for signature 'MPSE'

```r
mp_plot_ord(
```

---

### mp_plot_ord

**Plotting the result of PCA, PCoA, CCA, RDA, NDMS or DCA**

**Description**

Plotting the result of PCA, PCoA, CCA, RDA, NDMS or DCA

**Usage**

```r
mp_plot_ord(
  .data,
  .ord,
  .dim = c(1, 2),
  .group = NULL,
  .starshape = 15,
  .size = 2,
  .alpha = 1,
  .color = "black",
  starstroke = 0.5,
  show.side = TRUE,
  show.adonis = FALSE,
  ellipse = FALSE,
  show.sample = FALSE,
  show.envfit = FALSE,
  p.adjust = NULL,
  filter.envfit = FALSE,
  ...
)
```

## S4 method for signature 'MPSE'

```r
mp_plot_ord(
```
mp_plot_ord

.data,
.ord,
.dim = c(1, 2),
.group = NULL,
.starshape = 15,
.size = 2,
.alpha = 1,
.color = "black",
.starstroke = 0.5,
.show.side = TRUE,
.show.adonis = FALSE,
.ellipse = FALSE,
.show.sample = FALSE,
.show.envfit = FALSE,
.p.adjust = NULL,
.filter.envfit = FALSE,
...
)

## S4 method for signature 'tbl_mpse'
mp_plot_ord(
  .data,
  .ord,
  .dim = c(1, 2),
  .group = NULL,
  .starshape = 15,
  .size = 2,
  .alpha = 1,
  .color = "black",
  .starstroke = 0.5,
  .show.side = TRUE,
  .show.adonis = FALSE,
  .ellipse = FALSE,
  .show.sample = FALSE,
  .show.envfit = FALSE,
  .p.adjust = NULL,
  .filter.envfit = FALSE,
  ...
)

## S4 method for signature 'grouped_df_mpse'
mp_plot_ord(
  .data,
  .ord,
  .dim = c(1, 2),
  .group = NULL,
  .starshape = 15,
  .size = 2,
```r
.alpha = 1,
.color = "black",
starstroke = 0.5,
show.side = TRUE,
show.adonis = FALSE,
ellipse = FALSE,
show.sample = FALSE,
show.envfit = FALSE,
p.adjust = NULL,
filter.envfit = FALSE,
```

Arguments

- `.data` MPSE or tbl_mpse object, it is required.
- `.ord` a name of ordination (required), options are PCA, PCoA, DCA, NMDS, RDA, CCA, but the corresponding calculation methods (mp_cal_pca, mp_cal_pcoa, ...) should be done with action="add" before it.
- `.dim` integer which dimensions will be displayed, it should be a vector (length=2) default is c(1, 2). if the length is one the default will also be displayed.
- `.group` the column name of variable to be mapped to the color of points (fill character of geom_star) or one specified color code, default is NULL, meaning fill=NA, the points are hollow.
- `.starshape` the column name of variable to be mapped to the shapes of points (starshape character of geom_star) or one specified starshape of point of ggstar, default is NULL, meaning starshape=15 (circle point).
- `.size` the column name of variable to be mapped to the size of points (size character of geom_star) or one specified size of point of ggstar, default is NULL, meaning size=1.5, the size of points.
- `.alpha` the column name of variable to be mapped to the transparency of points (alpha character of geom_star) or one specified alpha of point of ggstar. default is NULL, meaning the alpha=1, the transparency of points.
- `.color` the column name of variable to be mapped to the color of line of points (color character of geom_star) or one specified starshape of point of ggstar, default is NULL, meaning the color is 'black'.
- `starstroke` numeric the width of edge of points, default is 0.5.
- `show.side` logical whether display the side boxplot with the specified .dim dimensions, default is TRUE.
- `show.adonis` logical whether display the result of mp_adonis with action='all', default is FALSE.
- `ellipse` logical, whether to plot ellipses, default is FALSE. (.group or .color variables according to the 'geom', the default geom is path, so .color can be mapped to the corresponding variable).
- `show.sample` logical, whether display the sample names of points, default is FALSE.
show.envfit    logical, whether display the result after run [mp_envfit()], default is FALSE.

p.adjust     a character method of p.adjust p.adjust, default is NULL, options are ‘fdr’, 'bonferroni', 'BH' etc.

filter.envfit logical or numeric, whether to remove the no significant environment factor after run [mp_envfit()], default is FALSE, meaning do not remove. If it is numeric, meaning the keep p.value or the adjust p with p.adjust the factors smaller than the numeric, e.g when filter.envfit=0.05 or (filter.envfit=TRUE), meaning the factors of p <= 0.05 will be displayed.

... additional parameters, see also the stat_ellipse.

See Also

[mp_cal_pca()], [mp_cal_pcoa], [mp_cal_nmds], [mp_cal_rda], [mp_cal_cca], [mp_envfit()] and [mp_extract_internal_attr()]

Examples

## Not run:
library(vegan)
data(varespec, varechem)
mpse <- MPSE(assays=list(Abundance=t(varespec)), colData=varechem)
envformula <- paste("~", paste(colnames(varechem), collapse="+")) %>% as.formula
mpse %<>% mp_cal_cca(.abundance=Abundance, .formula=envformula, action="add") %>%
mp_envfit(.ord=CCA, .env=colnames(varechem), permutations=9999, action="add")
mpse
p1 <- mpse %>% mp_plot_ord(.ord=CCA, .group=Al, .size=Mn)
p1
p2 <- mpse %>% mp_plot_ord(.ord=CCA, .group=Al, .size=Mn, show.sample=TRUE)
p2
p3 <- mpse %>% mp_plot_ord(.ord=CCA, .group="blue", .size=Mn, .alpha=0.8, show.sample=TRUE)
p3
p4 <- mpse %>% mp_plot_ord(.ord=CCA, .group=Al, .size=Mn, show.sample=TRUE, show.envfit=TRUE)
p4
## End(Not run)
Usage

```r
mp_plot_rarecurve(
  .data,
  .rare,
  .alpha = c("Observe", "Chao1", "ACE"),
  .group = NULL,
  nrow = 1,
  plot.group = FALSE,
  ...
)
```

## S4 method for signature 'MPSE'
```r
mp_plot_rarecurve(
  .data,
  .rare,
  .alpha = c("Observe", "Chao1", "ACE"),
  .group = NULL,
  nrow = 1,
  plot.group = FALSE,
  ...
)
```

## S4 method for signature 'tbl_mpse'
```r
mp_plot_rarecurve(
  .data,
  .rare,
  .alpha = c("Observe", "Chao1", "ACE"),
  .group = NULL,
  nrow = 1,
  plot.group = FALSE,
  ...
)
```

## S4 method for signature 'grouped_df_mpse'
```r
mp_plot_rarecurve(
  .data,
  .rare,
  .alpha = c("Observe", "Chao1", "ACE"),
  .group = NULL,
  nrow = 1,
  plot.group = FALSE,
  ...
)
```

Arguments

- `.data` MPSE object or tbl_mpse after it was performed `mp_cal_rarecurve` with `action='add'`
- `.rare` the column names of
mp_plot_upset

Plotting the different number of OTU between group via UpSet plot

Description

Plotting the different number of OTU between group via UpSet plot

Usage

mp_plot_upset(.data, .group, .upset = NULL, ...)

## S4 method for signature 'MPSE'
mp_plot_upset(.data, .group, .upset = NULL, ...)

## S4 method for signature 'tbl_mpse'
mp_plot_upset(.data, .group, .upset = NULL, ...)

## S4 method for signature 'grouped_df_mpse'
mp_plot_upset(.data, .group, .upset = NULL, ...)
Arguments

.data MPSE obejct or tbl_mpse object
.group the column name of group
.upset the column name of result after run mp_cal_upset
... additional parameters, see also 'scale_x_upset' of 'ggupset'.

Author(s)

Shuangbin Xu

Examples

## Not run:
data(mouse.time.mpse)
mpse <- mouse.time.mpse %>%
  mp_rrarefy(.abundance=Abundance) %>%
  mp_cal_upset(.abundance=RareAbundance, .group=time)
mpse
p <- mpse %>% mp_plot_upset(.group=time, .upset=ggupsetOftime)
p
## End(Not run)

---

**mp_plot_venn**

Plotting the different number of OTU between groups with Venn Diagram.

Description

Plotting the different number of OTU between groups with Venn Diagram.

Usage

`mp_plot_venn(.data, .group, .venn = NULL, ...)`

## S4 method for signature 'MPSE'
`mp_plot_venn(.data, .group, .venn = NULL, ...)`

## S4 method for signature 'tbl_mpse'
`mp_plot_venn(.data, .group, .venn = NULL, ...)`

## S4 method for signature 'grouped_df_mpse'
`mp_plot_venn(.data, .group, .venn = NULL, ...)`
Arguments

- `.data` MPSE object or tbl_mpse object
- `.group` the column names of group to be visualized
- `.venn` the column names of result after run `mp_cal_venn`
- `...` additional parameters, such as `size`, `label_size`, `edge_size` etc, see also `ggVennDiagram`.

Author(s)

Shuangbin Xu

Examples

```r
## Not run:
data(mouse.time.mpse)
mpse <- mouse.time.mpse %>%
  mp_rrarefy() %>%
  mp_cal_venn(.abundance=RareAbundance, .group=time, action="add")
mpse
p <- mpse %>% mp_plot_venn(.group=time, .venn=vennOftime)
p
## End(Not run)
```

---

### Description

**mp_rrarefy** method

### Usage

```r
mp_rrarefy(
  .data,
  .abundance = NULL,
  raresize,
  trimOTU = FALSE,
  trimSample = FALSE,
  seed = 123,
  ...)
```

---

```r
## S4 method for signature 'MPSE'
mp_rrarefy(
  .data,
  .abundance = NULL,
  ...)
```
raresize,
trimOTU = FALSE,
trimSample = FALSE,
seed = 123,
...
)

## S4 method for signature 'tbl_mpse'
mp_rrarefy(
  .data,
  .abundance = NULL,
  raresize,
  trimOTU = FALSE,
  trimSample = FALSE,
  seed = 123,
  ...
)

## S4 method for signature 'grouped_df_mpse'
mp_rrarefy(
  .data,
  .abundance = NULL,
  raresize,
  trimOTU = FALSE,
  trimSample = FALSE,
  seed = 123,
  ...
)

Arguments

- `.data` MPSE or tbl_mpse object
- `.abundance` the name of OTU(feature) abundance column, default is Abundance.
- `raresize` integer Subsample size for rarefying community.
- `trimOTU` logical Whether to remove the otus that are no longer present in any sample after rarefaction
- `trimSample` logical whether to remove the samples that do not have enough abundance (raresize), default is FALSE.
- `seed` a random seed to make the rarefy reproducible, default is 123.
- `...` additional parameters, meaningless now.

Value

update object

Author(s)

Shuangbin Xu
mp_select_as_tip

See Also

[mp_extract_assays()] and [mp_decostand()]

Examples

data(mouse.time.mpse)
mouse.time.mpse %>% mp_rrarefy()

mp_select_as_tip

select specific taxa level as rownames of MPSE

Description

select specific taxa level as rownames of MPSE

Usage

mp_select_as_tip(x, tip.level = "OTU")

## S4 method for signature 'MPSE'
mp_select_as_tip(x, tip.level = "OTU")

## S4 method for signature 'tbl_mpse'
mp_select_as_tip(x, tip.level = "OTU")

## S4 method for signature 'grouped_df_mpse'
mp_select_as_tip(x, tip.level = "OTU")

Arguments

x MPSE object
tip.level the taxonomy level, default is 'OTU'.

Examples

## Not run:
data(mouse.time.mpse)
newmpse <- mouse.time.mpse %>%
    mp_select_as_tip(tip.level = Species)
newmpse

## End(Not run)
mp_stat_taxa  
Count the number and total number taxa for each sample at different taxonomy levels

Description
Count the number and total number taxa for each sample at different taxonomy levels

Usage

```r
mp_stat_taxa(.data, .abundance, action = "add", ...)
```

## S4 method for signature 'MPSE'

```r
mp_stat_taxa(.data, .abundance, action = "add", ...)
```

## S4 method for signature 'tbl_mpse'

```r
mp_stat_taxa(.data, .abundance, action = "add", ...)
```

## S4 method for signature 'grouped_df_mpse'

```r
mp_stat_taxa(.data, .abundance, action = "add", ...)
```

Arguments
- `.data` MPSE or tbl_mpse object
- `.abundance` the column name of abundance to be calculated
- `action` a character "get" returns a table only contained the number and total number for each sample at different taxonomy levels, "only" returns a non-redundant tibble contained a nest column (StatTaxaInfo) and other sample information, "add" returns a update object (.data) contained a nest column (StatTaxaInfo).
- `...` additional parameter

Value
update object or tbl_df according action argument

Author(s)
Shuangbin Xu

Examples
```r
data(mouse.time.mpse)
mouse.time.mpse %>%
  mp_stat_taxa(.abundance=Abundance, action="only")
```
multi_compare

Description

A container for performing two or more sample test.

Usage

multi_compare(
    fun = wilcox.test,
    data,
    feature,
    factorNames,
    subgroup = NULL,
    ...
)

Arguments

fun character, the method for test, optional ""

data data.frame, nrow sample * ncol feature+factorNames.

feature vector, the features wanted to test.

factorNames character, the name of a factor giving the corresponding groups.

subgroup vector, the names of groups, default is NULL.

..., additional arguments for fun.

Value

The result of fun, if fun is wilcox.test, it will return the list with class "htest".

Author(s)

Shuangbin Xu

Examples

datest <- data.frame(A=rnorm(1:10,mean=5),
    B=rnorm(2:11, mean=6),
    group=c(rep("case",5),rep("control",5)))
head(datest)
multi_compare(fun=wilcox.test,data=datest,
    feature=c("A", "B"),factorNames="group")
da2 <- data.frame(A=rnorm(1:15,mean=5),
    B=rnorm(2:16,mean=6),
    group=c(rep("case1",5),rep("case2",5),rep("control",5)))
multi_compare(fun=wilcox.test,data=d2,
feature=c("A", "B"), factorNames="group", subgroup=c("case1", "case2"))

ordplotClass-class  ordplotClass class

Description

ordplotClass class

Slots

coord  matrix object contained the coordinate for ordination plot.

xlab  character object contained the text of xlab for ordination plot.

ylab  character object contained the text of ylab for ordination plot.

title  character object contained the text of title for ordination plot.

pcasample-class  pcasample class

Description

pcasample class

Slots

pca  prcomp or pcoa object

sampleda  associated sample information

pcoa-class  pcoa class

Description

pcoa class

See Also

pcoa
prcomp-class

Description

prcomp class

See Also

prcomp

print

print some objects

Description

print some objects

Usage

## S3 method for class 'MPSE'
print(
x,
..., 
n = NULL,
width = NULL,
max_extra_cols = NULL,
max_footer_lines = NULL
)

## S3 method for class 'tbl_mpse'
print(x, ..., n = NULL, width = NULL, max_extra_cols = NULL)

## S3 method for class 'grouped_df_mpse'
print(x, ..., n = NULL, width = NULL, max_extra_cols = NULL)

## S3 method for class 'rarecurve'
print(x, ..., n = NULL, width = NULL, max_extra_cols = NULL)

Arguments

x  Object to format or print.

...  Other arguments passed on to individual methods.

n  Number of rows to show. If ‘NULL’, the default, will print all rows if less than option ‘tibble.print_max’. Otherwise, will print ‘tibble.print_min’ rows.
read_qza

width Width of text output to generate. This defaults to `NULL`, which means use `getOption("tibble.width")` or (if also `NULL`) `getOption("width")`; the latter displays only the columns that fit on one screen. You can also set `options(tibble.width = Inf)` to override this default and always print all columns.

max_extra_cols Number of extra columns to print abbreviated information for, if the width is too small for the entire tibble. If `NULL`, the default, will print information about at most `tibble.max_extra_cols` extra columns.

max_footer_lines integer maximum number of lines for the footer.

Value

print information

Description

the function was designed to read the output of qiime2.

Usage

read_qza(qzafile, parallel = FALSE)

Arguments

qzafile character, the format of file should be one of `BIOMV210DirFmt`, `TSVTaxonomyDirectoryFormat`, `NewickDirectoryFormat` and `DNASequencesDirectoryFormat`.

parallel logical, whether parsing the taxonomy by multi-parallel, default is FALSE.

Value

list contained one or multiple object of feature table, taxonomy table, tree and represent sequences.

Examples

## Not run:
otuqzafile <- system.file("extdata", "table.qza",
  package="MicrobiotaProcess")

otuqza <- read_qza(otuqzafile)
str(otuqza)

## End(Not run)
Objects exported from other packages

Description

These objects are imported from other packages. Follow the links below to see their documentation.

dplyr arrange, distinct, filter, group_by, left_join, mutate, pull, rename, select, slice, ungroup
ggplot2 fortify, remove_missing
ggtree td_filter, td_unnest
magrittr %<>%, %>%, extract

SummarizedExperiment colData, colData<-, rowData
tibble as_tibble
tidytr nest, unnest
tidytree as.treedata

scale_fill_diff_cladogram

Create the scale of mp_plot_diff_cladogram.

Description

Create the scale of mp_plot_diff_cladogram.

Usage

scale_fill_diff_cladogram(values, breaks = waiver(), na.value = "grey50", ...)

Arguments

values a set of aesthetic values (different group (default)) to map data values to.
breaks One of ‘NULL’ for no breaks, ‘waiver()’ for the default breaks, A character vector of breaks.
na.value The aesthetic value to use for missing (‘NA’) values.
... see also ’discrete_scale’ of ‘ggplot2’.
set_diff_boxplot_color

*set the color scale of plot generated by mp_plot_diff_boxplot*

**Description**

set the color scale of plot generated by mp_plot_diff_boxplot

**Usage**

set_diff_boxplot_color(.data, values, ...)

**Arguments**

- `.data` the aplot object generated by mp_plot_diff_boxplot.
- `values` the color vector, required.
- `...` additional parameters, see also the 'scale_fill_manual' of 'ggplot2'

---

set_scale_theme

*adjust the color of heatmap of mp_plot_dist*

**Description**

adjust the color of heatmap of mp_plot_dist

**Usage**

set_scale_theme(.data, x, aes_var)

**Arguments**

- `.data` the plot of heatmap of mp_plot_dist
- `x` the scale or theme
- `aes_var` character the variable (column) name of color or size.
Description

method extensions to show for diffAnalysisClass or alphasample objects.

Usage

## S4 method for signature 'diffAnalysisClass'
show(object)

## S4 method for signature 'alphasample'
show(object)

## S4 method for signature 'MPSE'
show(object)

Arguments

object object, diffAnalysisClass or alphasample class

Value

print info

Author(s)

Shuangbin Xu

Examples

## Not run:
data(kostic2012crc)
kostic2012crc %<>% as.phyloseq()
head(phyloseq::sample_data(kostic2012crc),3)
kostic2012crc <- phyloseq::rarefy_even_depth(kostic2012crc,rngseed=1024)
table(phyloseq::sample_data(kostic2012crc)$DIAGNOSIS)
set.seed(1024)
diffres <- diff_analysis(kostic2012crc, classgroup="DIAGNOSIS",
                        mlfun="lda", filtermod="Fdr",
                        firstcomfun = "kruskal.test",
                        firstalpha=0.05, strictmod=TRUE,
                        secondcomfun = "wilcox.test",
                        subclmin=3, subclwilc=TRUE,
                        secondalpha=0.01, lda=3)

show(diffres)
## split_data

### Split Large Vector or DataFrame

**Description**

Split large vector or dataframe to list class, which contain subset vectors or dataframe of origin vector or dataframe.

**Usage**

```r
split_data(x, nums, chunks = NULL, random = FALSE)
```

**Arguments**

- `x`: vector class or data.frame class.
- `nums`: integer.
- `chunks`: integer. use chunks if nums is missing. Note nums and chunks shouldn’t concurrently be NULL, default is NULL.
- `random`: bool, whether split randomly, default is `FALSE`, if you want to split data randomly, you can set `TRUE`, and if you want the results are reproducible, you should add seed before.

**Value**

the subset of x, vector or data.frame class.

**Author(s)**

Shuangbin Xu

**Examples**

```r
data(iris)
irislist <- split_data(iris, 40)
dalist <- c(1:100)
dalist <- split_data(dalist, 30)
```
split_str_to_list  

split a dataframe contained one column

Description
split a dataframe contained one column with a specify field separator character.

Usage

split_str_to_list(
  strdataframe,
  prefix = "tax",
  sep = "; ",
  extra = "drop",
  fill = "right",
  ...
)

Arguments
strdataframe  dataframe; a dataframe contained one column to split.
prefix  character; the result dataframe columns names prefix, default is "tax".
sep  character; the field separator character, default is " ; ".
extra  character; See separate details.
fill  character; See separate details.
...

Value
data.frame of strdataframe by sep.

Author(s)
Shuangbin Xu

Examples

## Not run:
otudafile <- system.file("extdata", "otu_tax_table.txt",
  package="MicrobiotaProcess")
samplefile <- system.file("extdata",
  "sample_info.txt", package="MicrobiotaProcess")
otuda <- read.table(otudafile, sep="\t", header=TRUE,
  row.names=1, check.names=FALSE,
  skip=1, comment.char="")
sampled <- read.table(samplefile,
  sep="\t", header=TRUE, row.names=1)
```r
taxdf <- otuda[!sapply(otuda, is.numeric)]
taxdf <- split_str_to_list(taxdf)
head(taxdf)

## End(Not run)
```

---

**taxonomy**

*extract the taxonomy annotation in MPSE object*

---

**Description**

extract the taxonomy annotation in MPSE object

**Usage**

```r
taxonomy(x, ...)

## S4 method for signature 'MPSE'
taxonomy(x, ...)

## S4 method for signature 'tbl_mpse'
taxonomy(x, ...)

## S4 method for signature 'grouped_df_mpse'
taxonomy(x, ...)

mp_extract_taxonomy(x, ...)

## S4 method for signature 'MPSE'
mp_extract_taxonomy(x, ...)

## S4 method for signature 'tbl_mpse'
mp_extract_taxonomy(x, ...)

## S4 method for signature 'grouped_df_mpse'
mp_extract_taxonomy(x, ...)
```

**Arguments**

- `x` MPSE object
- `...` additional arguments

**Value**

data.frame contained taxonomy information
data.frame contained taxonomy annotation.
Description

theme_taxbar

Usage

```r
theme_taxbar(
  axis.text.x = element_text(angle = -45, hjust = 0, size = 8),
  legend.position = "bottom",
  legend.box = "horizontal",
  legend.text = element_text(size = 8),
  legend.title = element_blank(),
  strip.text.x = element_text(size = 12, face = "bold"),
  strip.background = element_rect(colour = "white", fill = "grey"),
  ...)
```

Arguments

- `axis.text.x` `element_text`, x axis tick labels.
- `legend.position` character, default is "bottom".
- `legend.box` character, arrangement of legends, default is "horizontal".
- `legend.text` `element_text`, legend labels text.
- `legend.title` `element_text`, legend title text
- `strip.text.x` `element_text`, strip text of x
- `strip.background` `element_rect`, the background of x
- `...` additional parameters

Value

updated ggplot object with new theme

See Also

- `theme`
Examples

```r
## Not run:
library(ggplot2)
data(test_otu_data)
test_otu_data %<>% as.phyloseq()
otubar <- ggbartax(test_otu_data, settheme=FALSE) +
         xlab(NULL) + ylab("relative abundance(%)") +
         theme_taxbar()

## End(Not run)
```
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