

Package ‘microeco’

October 31, 2025

Type Package

Title Microbial Community Ecology Data Analysis

Version 1.16.0

Author Chi Liu [aut, cre],
Felipe R. P. Mansoldo [ctb],
Minjie Yao [ctb],
Xiangzhen Li [ctb]

Maintainer Chi Liu <liuchi0426@126.com>

Description A series of statistical and plotting approaches in microbial community ecology based on the R6 class. The classes are designed for data preprocessing, taxa abundance plotting, alpha diversity analysis, beta diversity analysis, differential abundance test, null model analysis, network analysis, machine learning, environmental data analysis and functional analysis.

URL <https://github.com/ChiLiubio/microeco>

Depends R (>= 3.5.0)

Imports R6, stats, ape, vegan, rlang, data.table, magrittr, dplyr,
tibble, scales, grid, ggplot2 (>= 3.5.0), RColorBrewer,
reshape2, igraph (>= 2.0.0), lifecycle

Suggests GUniFrac, MASS, ggpubr, randomForest, ggdendro, ggrepel,
agricolae, gridExtra, picante, pheatmap, rgexf, mice, GGally

License GPL-3

LazyData true

Encoding UTF-8

NeedsCompilation no

Repository CRAN

Date/Publication 2025-10-31 07:10:03 UTC

RoxygenNote 7.3.2

Contents

clone	2
dataset	3
dropallfactors	4
env_data_16S	5
fungi_func_FungalTraits	5
fungi_func_FUNGuild	5
microeco	6
microtable	6
otu_table_16S	19
otu_table_ITS	19
phylo_tree_16S	20
prok_func_FAPROTAX	20
prok_func_NJC19_list	20
sample_info_16S	21
sample_info_ITS	21
Tax4Fun2_KEGG	21
taxonomy_table_16S	22
taxonomy_table_ITS	22
tidy_taxonomy	22
trans_abund	24
trans_alpha	35
trans_beta	41
trans_classifier	51
trans_diff	61
trans_env	72
trans_func	88
trans_network	95
trans_norm	112
trans_nullmodel	116
trans_venn	126
Index	132

clone	<i>Copy an R6 class object</i>
-------	--------------------------------

Description

Copy an R6 class object

Usage

clone(x, deep = TRUE)

Arguments

x	R6 class object
deep	default TRUE; TRUE means deep copy, i.e. copied object is unlinked with the original one.

Value

identical but unlinked R6 object

Examples

```
data("dataset")
clone(dataset)
```

dataset	<i>The dataset structured with microtable class for the demonstration of examples</i>
---------	---

Description

The dataset arose from 16S rRNA gene amplicon sequencing of wetland soils in China <doi:10.1016/j.geoderma.2018.09.035>. In dataset\$sample_table, the 'Group' column means Chinese inland wetlands (IW), coastal wetland (CW) and Tibet plateau wetlands (TW). The column 'Type' denotes the sampling region: northeastern region (NE), northwest region (NW), North China area (NC), middle-lower reaches of the Yangtze River (YML), southern coastal area (SC), upper reaches of the Yangtze River (YU) and Qinghai-Tibet Plateau (QTP). The column 'Saline' represents the saline soils and non-saline soils.

Usage

```
data(dataset)
```

Format

R6 class object

Details

- sample_table: sample information table
- otu_table: species-sample abundance table
- tax_table: taxonomic table
- phylo_tree: phylogenetic tree
- taxa_abund: taxa abundance list with several tables for Phylum...Genus
- alpha_diversity: alpha diversity table
- beta_diversity: list with several beta diversity distance matrix

dropallfactors	<i>Remove all factors in a data frame</i>
----------------	---

Description

Remove all factors in a data frame

Usage

```
dropallfactors(x, unfac2num = FALSE, char2num = FALSE)
```

Arguments

x	data.frame object
unfac2num	default FALSE; whether try to convert all character columns to numeric directly; If TRUE, it will attempt to convert each column, including those of character and factor types. First, it tries to convert them to the character type, and then checks if they can be converted to numeric. If the conversion to numeric is possible, it outputs the numeric type; otherwise, it outputs the character type. If FALSE, only columns with the factor attribute will be attempted for conversion. Factors will first be converted to character type, and then an attempt will be made to convert them to numeric. If successful, the numeric type will be output; otherwise, the character type will be output. This process can effectively remove the factor attribute. Note that this can only transform the columns that may be transformed to numeric without using factor.
char2num	default FALSE; whether force all the character to be numeric class by using factor as an intermediate. Therefore, this parameter can enforce the conversion of all character and factor types to numeric. This operation is very useful in some cases that numerical data is required as input.

Value

data frame without factor

Examples

```
data("taxonomy_table_16S")
taxonomy_table_16S[, 1] <- as.factor(taxonomy_table_16S[, 1])
str(dropallfactors(taxonomy_table_16S))
```

env_data_16S	<i>The environmental factors for the 16S example data</i>
--------------	---

Description

The environmental factors for the 16S example data

Usage

```
data(env_data_16S)
```

Format

```
data.frame
```

fungi_func_FungalTraits	<i>The FungalTraits database for fungi trait prediction</i>
-------------------------	---

Description

The FungalTraits database for fungi trait prediction

Usage

```
data(fungi_func_FungalTraits)
```

Format

```
data.frame
```

fungi_func_FUNGuild	<i>The FUNGuild database for fungi trait prediction</i>
---------------------	---

Description

The FUNGuild database for fungi trait prediction

Usage

```
data(fungi_func_FUNGuild)
```

Format

```
data.frame
```

microeco	<i>Introduction</i>	<i>to</i>	<i>microeco</i>	<i>package</i>
	(Rhref https://github.com/ChiLiubio/microeco https://github.com/ChiLiubio/microeco)			

Description

For the detailed tutorial on microeco package, please follow the links:

Online tutorial website: https://chiliubio.github.io/microeco_tutorial/

Download tutorial: https://github.com/ChiLiubio/microeco_tutorial/releases

For each R6 class, please open the help document by searching the class name. For example, to search microtable class, please run the command `help(microtable)` or `?microtable`.

Another way to open the help document of R6 class is to click the following links collected:

[microtable](#)
[trans_abund](#)
[trans_venn](#)
[trans_alpha](#)
[trans_beta](#)
[trans_diff](#)
[trans_network](#)
[trans_nullmodel](#)
[trans_classifier](#)
[trans_env](#)
[trans_func](#)
[trans_norm](#)

To report bugs or discuss questions, please use Github Issues (<https://github.com/ChiLiubio/microeco/issues>).

Before creating a new issue, please read the guideline (https://chiliubio.github.io/microeco_tutorial/notes.html#github-issues).

To cite microeco package in publications, please run the following command to get the references:
`citation("microeco")`

Reference:

Chi Liu, Felipe R. P. Mansoldo, Hankang Li, Alane Beatriz Vermelho, Raymond Jianxiong Zeng, Xiangzhen Li & Minjie Yao. A workflow for statistical analysis and visualization of microbiome omics data using the R microeco package. Nature Protocols (2025). DOI:10.1038/s41596-025-01239-4

Chi Liu, Yaoming Cui, Xiangzhen Li & Minjie Yao. microeco: an R package for data mining in microbial community ecology. FEMS Microbiology Ecology, 2021, 97(2): faa255. DOI:10.1093/femsec/faa255

microtable	<i>Create microtable object to store and manage all the basic files.</i>
------------	--

Description

This class is a wrapper for a series of operations on the basic data manipulations, including microtable object creation, data trimming, data filtering, rarefaction based on Paul et al. (2013) <doi:10.1371/journal.pone.0061217>, taxonomic abundance calculation, alpha and beta diversity calculation based on the An et al. (2019) <doi:10.1016/j.geoderma.2018.09.035> and Lozupone et al. (2005) <doi:10.1128/AEM.71.12.8228-8235.2005> and other basic operations.

Online tutorial: https://chiliubio.github.io/microeco_tutorial/

Download tutorial: https://github.com/ChiLiubio/microeco_tutorial/releases

Format

microtable.

Methods

Public methods:

- `microtable$new()`
- `microtable$filter_pollution()`
- `microtable$filter_taxa()`
- `microtable$rarefy_samples()`
- `microtable$tidy_dataset()`
- `microtable$add_rownames2taxonomy()`
- `microtable$sample_sums()`
- `microtable$taxa_sums()`
- `microtable$sample_names()`
- `microtable$taxa_names()`
- `microtable$rename_taxa()`
- `microtable$merge_samples()`
- `microtable$merge_taxa()`
- `microtable$save_table()`
- `microtable$cal_abund()`
- `microtable$save_abund()`
- `microtable$cal_alphadiv()`
- `microtable$save_alphadiv()`
- `microtable$cal_betadiv()`
- `microtable$save_betadiv()`
- `microtable$print()`
- `microtable$clone()`

Method `new()`:

Usage:

```
microtable$new(
  otu_table,
  sample_table = NULL,
  tax_table = NULL,
  phylo_tree = NULL,
  rep_fasta = NULL,
  auto_tidy = FALSE
)
```

Arguments:

`otu_table` data.frame class; Feature abundance table; rownames are features (e.g. OTUs/ASVs/species/genes); column names are samples.

`sample_table` default NULL; data.frame; The sample information table; rownames are samples; columns are sample metadata; If not provided, the function can generate a table automatically according to the sample names in `otu_table`.

`tax_table` default NULL; data.frame class; Taxonomic information table; rownames are features; column names are taxonomic ranks. This can also be other hierarchical information tables, such as traits, genes, or metabolic pathways.

`phylo_tree` default NULL; phylo class; Phylogenetic tree; It must be read with the `read.tree` function of ape package.

`rep_fasta` default NULL; DNASTringSet, list or DNABin class; Representative sequences of OTUs/ASVs. The sequences should be read with the `readDNASTringSet` function of Biostrings package (DNASTringSet class), `read.fasta` function of seqinr package (list class), or `read.FASTA` function of ape package (DNABin class).

`auto_tidy` default FALSE; Whether tidy the data in the microtable object automatically. If TRUE, the function can invoke the `tidy_dataset` function.

Returns: an object of microtable class with the following components:

`sample_table` The sample information table.

`otu_table` The feature table.

`tax_table` The taxonomic table.

`phylo_tree` The phylogenetic tree.

`rep_fasta` The sequences.

`taxa_abund` default NULL; use `cal_abund` function to calculate.

`alpha_diversity` default NULL; use `cal_alphadiv` function to calculate.

`beta_diversity` default NULL; use `cal_betadiv` function to calculate.

Examples:

```
data(otu_table_16S)
data(taxonomy_table_16S)
data(sample_info_16S)
data(phylo_tree_16S)
m1 <- microtable$new(otu_table = otu_table_16S)
m1 <- microtable$new(sample_table = sample_info_16S, otu_table = otu_table_16S,
  tax_table = taxonomy_table_16S, phylo_tree = phylo_tree_16S)
# trim each data in the object
m1$tidy_dataset()
```

Method `filter_pollution()`: Filter the features considered pollution in `microtable$tax_table`. This operation will remove any line of the `microtable$tax_table` containing any the word in `taxa` parameter regardless of word case.

Usage:

```
microtable$filter_pollution(taxa = c("mitochondria", "chloroplast"))
```

Arguments:

`taxa` default `c("mitochondria", "chloroplast")`; filter mitochondria and chloroplast, or others as needed.

Returns: updated microtable object

Examples:

```
m1$filter_pollution(taxa = c("mitochondria", "chloroplast"))
```

Method `filter_taxa()`: Filter the features with low abundance and/or low occurrence frequency for `otu_table` or `taxa_abund` list.

Usage:

```
microtable$filter_taxa(
  rel_abund = 0,
  freq = 1,
  include_lowest = TRUE,
  for_taxa_abund = FALSE
)
```

Arguments:

`rel_abund` default 0; the relative abundance threshold, such as 0.0001.

`freq` default 1; the occurrence frequency threshold. For example, the number 2 represents filtering the feature that occurs less than 2 times. A number smaller than 1 is also allowable. For instance, the number 0.1 represents filtering the feature that occurs in less than 10% samples.

`include_lowest` default TRUE; whether include the feature with the threshold.

`for_taxa_abund` default FALSE; whether apply this function to `taxa_abund` list. FALSE means using this function for `otu_table`

Returns: updated microtable object

Examples:

```
\donttest{
d1 <- clone(m1)
d1$filter_taxa(rel_abund = 0.0001, freq = 0.2)
}
```

Method `rarefy_samples()`: Rarefy communities to make all samples have same count number.

Usage:

```
microtable$rarefy_samples(
  method = c("rarefy", "SRS")[1],
  sample.size = NULL,
  ...
)
```

Arguments:

`method` default `c("rarefy", "SRS")[1]`; "rarefy" represents the classical resampling like `rrarefy` function of `vegan` package. "SRS" is scaling with ranked subsampling method based on the SRS package provided by Lukas Beule and Petr Karlovsky (2020) <DOI:10.7717/peerj.9593>.

`sample.size` default `NULL`; library size. If not provided, use the minimum number across all samples. For "SRS" method, this parameter is passed to `Cmin` parameter of SRS function of SRS package.

... parameters pass to `norm` function of `trans_norm` class.

Returns: rarefied microtable object.

Examples:

```
\donttest{
m1$rarefy_samples(sample.size = min(m1$sample_sums()))
}
```

Method `tidy_dataset()`: Trim all the data in the microtable object to make taxa and samples consistent. The results are intersections across data.

Usage:

```
microtable$tidy_dataset(main_data = FALSE)
```

Arguments:

`main_data` default `FALSE`; if `TRUE`, only basic data in microtable object is trimmed. Otherwise, all data, including `taxa_abund`, `alpha_diversity` and `beta_diversity`, are all trimmed.

Returns: None. The data in the object are tidied up. If `tax_table` is in object, its row names are completely same with the row names of `otu_table`.

Examples:

```
m1$tidy_dataset(main_data = TRUE)
```

Method `add_rownames2taxonomy()`: Add the row names of `microtable$tax_table` as its last column. This is especially useful when the row names of `microtable$tax_table` are required as a taxonomic level for the taxonomic abundance calculation and biomarker identification.

Usage:

```
microtable$add_rownames2taxonomy(use_name = "OTU")
```

Arguments:

`use_name` default "OTU"; The name of the column added in the `tax_table`.

Returns: `tax_table` updated in the object.

Examples:

```
\donttest{
m1$add_rownames2taxonomy()
}
```

Method `sample_sums()`: Sum the abundance for each sample.

Usage:

```
microtable$sample_sums()
```

Returns: abundance in each sample.

Examples:

```
\donttest{
m1$sample_sums()
}
```

Method taxa_sums(): Sum the abundance for each taxon.

Usage:

```
microtable$taxa_sums()
```

Returns: abundance in each taxon.

Examples:

```
\donttest{
m1$taxa_sums()
}
```

Method sample_names(): Show the sample names.

Usage:

```
microtable$sample_names()
```

Returns: sample names.

Examples:

```
\donttest{
m1$sample_names()
}
```

Method taxa_names(): Show the taxa names.

Usage:

```
microtable$taxa_names()
```

Returns: taxa names.

Examples:

```
\donttest{
m1$taxa_names()
}
```

Method rename_taxa(): Rename the features, including the row names of otu_table, row names of tax_table, tip labels of phylo_tree and names in rep_fasta.

Usage:

```
microtable$rename_taxa(newname_prefix = "ASV_")
```

Arguments:

newname_prefix default "ASV_"; the prefix of new names; new names will be newname_prefix + numbers according to the order of row names in otu_table.

Returns: renamed object

Examples:

```
\donttest{
m1$rename_taxa()
}
```

Method `merge_samples()`: Merge samples according to specific groups to generate a new `microtable` object.

Usage:

```
microtable$merge_samples(group)
```

Arguments:

`group` a column name in `sample_table` of `microtable` object.

Returns: a merged `microtable` object.

Examples:

```
\donttest{
m1$merge_samples("Group")
}
```

Method `merge_taxa()`: Merge taxa according to a specific taxonomic rank to generate a new `microtable` object.

Usage:

```
microtable$merge_taxa(taxa = "Genus")
```

Arguments:

`taxa` default "Genus"; the specific rank in `tax_table`.

Returns: a merged `microtable` object.

Examples:

```
\donttest{
m1$merge_taxa(taxa = "Genus")
}
```

Method `save_table()`: Save each basic data in `microtable` object as local file.

Usage:

```
microtable$save_table(dirpath = "basic_files", sep = ",", ...)
```

Arguments:

`dirpath` default "basic_files"; directory to save the tables, phylogenetic tree and sequences in `microtable` object. It will be created if not found.

`sep` default ","; the field separator string, used to save tables. Same with `sep` parameter in [write.table](#) function. default ' ' correspond to the file name suffix 'csv'. The option '\t' correspond to the file name suffix 'tsv'. For other options, suffix are all 'txt'.

... parameters passed to [write.table](#).

Examples:

```
\dontrun{
m1$save_table()
}
```

Method `cal_abund()`: Calculate the taxonomic abundance at each taxonomic level or selected levels.

Usage:

```
microtable$cal_abund(
  select_cols = NULL,
  rel = TRUE,
  merge_by = "|",
  split_group = FALSE,
  split_by = "&",
  split_column = NULL,
  split_special_char = "&&"
)
```

Arguments:

`select_cols` default NULL; numeric vector (column sequences) or character vector (column names of `microtable$tax_table`); applied to select columns to calculate abundances according to ordered hierarchical levels. This parameter is very useful when only part of the columns are needed to calculate abundances.

`rel` default TRUE; if TRUE, relative abundance is used; if FALSE, absolute abundance (i.e. raw values) will be summed.

`merge_by` default "|"; the symbol to merge and concatenate taxonomic names of different levels.

`split_group` default FALSE; if TRUE, split the rows to multiple rows according to one or more columns in `tax_table` when there is multiple mapping information.

`split_by` default "&"; Separator delimiting collapsed values; only available when `split_group` = TRUE.

`split_column` default NULL; one column name used for the splitting in `tax_table` for each abundance calculation; only available when `split_group` = TRUE. If not provided, the function will split each column that containing the `split_by` character.

`split_special_char` default "&&"; special character that will be used forcibly to split multiple mapping information in `tax_table` by default no matter `split_group` setting. For example, the hierarchical information of MetaCyc metabolic pathways from the `file2meco` package may have multiple ontology entries linked together. In this case, the default parameters are automatically changed to `split_group` = TRUE and `split_by` = `split_special_char`, which is the default "&&". If users have other multi-label data in `tax_table`, they can adjust the `split_group`, `split_by`, and `split_column` parameters accordingly.

Returns: `taxa_abund` list in object.

Examples:

```
\donttest{
m1$cal_abund()
}
```

Method `save_abund()`: Save taxonomic abundance as local file.

Usage:

```
microtable$save_abund(
  dirpath = "taxa_abund",
  merge_all = FALSE,
```

```

rm_un = FALSE,
rm_pattern = "__$",
sep = ",",
...
)

```

Arguments:

`dirpath` default "taxa_abund"; directory to save the taxonomic abundance files. It will be created if not found.

`merge_all` default FALSE; Whether merge all tables into one. The merged file format is generally called 'mpa' style.

`rm_un` default FALSE; Whether remove unclassified taxa in which the name ends with '___' generally.

`rm_pattern` default "__\$"; The pattern searched through the merged taxonomic names. See also pattern parameter in [grepl](#) function. Only available when `rm_un` = TRUE. The default "__\$" means removing the names end with '___'.

`sep` default ","; the field separator string. Same with `sep` parameter in [write.table](#) function. default ',' correspond to the file name suffix 'csv'. The option '\t' correspond to the file name suffix 'tsv'. For other options, suffix are all 'txt'.

... parameters passed to [write.table](#).

Examples:

```

\dontrun{
m1$save_abund(dirpath = "taxa_abund")
m1$save_abund(merge_all = TRUE, rm_un = TRUE, sep = "\t")
}

```

Method `cal_alphadiv()`: Calculate alpha diversity.

Usage:

```
microtable$cal_alphadiv(measures = NULL, PD = FALSE)
```

Arguments:

`measures` default NULL; one or more indexes in `c("Observed", "Coverage", "Chao1", "ACE", "Shannon", "Simpson", "InvSimpson", "Fisher", "Pielou")`; The default NULL represents that all the measures are calculated. 'Shannon', 'Simpson' and 'InvSimpson' are calculated based on `vegan::diversity` function; 'Chao1' and 'ACE' depend on the function `vegan::estimateR`. 'Fisher' index relies on the function `vegan::fisher.alpha`. "Observed" means the observed species number in a community, i.e. richness. "Coverage" represents good's coverage. It is defined:

$$Coverage = 1 - \frac{f1}{n}$$

where n is the total abundance of a sample, and $f1$ is the number of singleton (species with abundance 1) in the sample. "Pielou" denotes the Pielou evenness index. It is defined:

$$J = \frac{H'}{\ln(S)}$$

where H' is Shannon index, and S is the species number.

PD default FALSE; whether Faith's phylogenetic diversity is calculated. The calculation depends on the function `picante::pd`. Note that the phylogenetic tree (`phylo_tree` object in the data) is required for PD.

Returns: `alpha_diversity` stored in the object. The `se.chao1` and `se.ACE` are the standard errors of Chao1 and ACE, respectively.

Examples:

```
\donttest{
m1$cal_alphadiv(measures = NULL, PD = FALSE)
class(m1$alpha_diversity)
}
```

Method `save_alphadiv()`: Save alpha diversity table to the computer.

Usage:

```
microtable$save_alphadiv(dirpath = "alpha_diversity")
```

Arguments:

`dirpath` default "alpha_diversity"; directory name to save the `alpha_diversity.csv` file.

Method `cal_betadiv()`: Calculate beta diversity dissimilarity matrix, such as Bray-Curtis, Jaccard, and UniFrac. See An et al. (2019) <doi:10.1016/j.geoderma.2018.09.035> and Lozupone et al. (2005) <doi:10.1128/AEM.71.12.8228-8235.2005>.

Usage:

```
microtable$cal_betadiv(
  method = NULL,
  unifrac = FALSE,
  binary = FALSE,
  force_jaccard_binary = TRUE,
  ...
)
```

Arguments:

`method` default NULL; a character vector with one or more elements; `c("bray", "jaccard")` is used when `method = NULL`; See the `method` parameter in `vegdist` function for more available options, such as 'aitchison' and 'robust.aitchison'.

`unifrac` default FALSE; whether UniFrac indexes (weighted and unweighted) are calculated. Phylogenetic tree is necessary when `unifrac = TRUE`.

`binary` default FALSE; Whether convert abundance to binary data (presence/absence).

`force_jaccard_binary` default TRUE; Whether forcibly convert abundance to binary data (presence/absence) when `method = "jaccard"`. The reason for this setting is that the Jaccard metric is commonly used for binary data. If `force_jaccard_binary = FALSE` is set, the conversion will not be enforced, but will instead be based on the setting of the `binary` parameter.

`...` parameters passed to `vegdist` function of `vegan` package.

Returns: `beta_diversity` list stored in the object.

Examples:

```
\donttest{
m1$cal_betadiv(unifrac = FALSE)
class(m1$beta_diversity)
}
```

Method `save_betadiv()`: Save beta diversity matrix to the computer.

Usage:

```
microtable$save_betadiv(dirpath = "beta_diversity")
```

Arguments:

`dirpath` default "beta_diversity"; directory name to save the beta diversity matrix files.

Method `print()`: Print the microtable object.

Usage:

```
microtable$print()
```

Method `clone()`: The objects of this class are cloneable with this method.

Usage:

```
microtable$clone(deep = FALSE)
```

Arguments:

`deep` Whether to make a deep clone.

Examples

```
## -----
## Method `microtable$new`
## -----

data(otu_table_16S)
data(taxonomy_table_16S)
data(sample_info_16S)
data(phylo_tree_16S)
m1 <- microtable$new(otu_table = otu_table_16S)
m1 <- microtable$new(sample_table = sample_info_16S, otu_table = otu_table_16S,
  tax_table = taxonomy_table_16S, phylo_tree = phylo_tree_16S)
# trim each data in the object
m1$tidy_dataset()

## -----
## Method `microtable$filter_pollution`
## -----

m1$filter_pollution(taxa = c("mitochondria", "chloroplast"))

## -----
## Method `microtable$filter_taxa`
## -----

d1 <- clone(m1)
```

```

d1$filter_taxa(rel_abund = 0.0001, freq = 0.2)

## -----
## Method `microtable$rarefy_samples`
## -----

m1$rarefy_samples(sample.size = min(m1$sample_sums()))

## -----
## Method `microtable$tidy_dataset`
## -----

m1$tidy_dataset(main_data = TRUE)

## -----
## Method `microtable$add_rownames2taxonomy`
## -----

m1$add_rownames2taxonomy()

## -----
## Method `microtable$sample_sums`
## -----

m1$sample_sums()

## -----
## Method `microtable$taxa_sums`
## -----

m1$taxa_sums()

## -----
## Method `microtable$sample_names`
## -----

m1$sample_names()

## -----
## Method `microtable$taxa_names`
## -----

```

```

m1$taxa_names()

## -----
## Method `microtable$rename_taxa`
## -----

m1$rename_taxa()

## -----
## Method `microtable$merge_samples`
## -----

m1$merge_samples("Group")

## -----
## Method `microtable$merge_taxa`
## -----

m1$merge_taxa(taxa = "Genus")

## -----
## Method `microtable$save_table`
## -----

## Not run:
m1$save_table()

## End(Not run)

## -----
## Method `microtable$cal_abund`
## -----

m1$cal_abund()

## -----
## Method `microtable$save_abund`
## -----

## Not run:
m1$save_abund(dirpath = "taxa_abund")
m1$save_abund(merge_all = TRUE, rm_un = TRUE, sep = "\t")

```

```
## End(Not run)

## -----
## Method `microtable$cal_alphadiv`
## -----

m1$cal_alphadiv(measures = NULL, PD = FALSE)
class(m1$alpha_diversity)

## -----
## Method `microtable$cal_betadiv`
## -----

m1$cal_betadiv(unifrac = FALSE)
class(m1$beta_diversity)
```

otu_table_16S	<i>The OTU table of the 16S example data</i>
---------------	--

Description

The OTU table of the 16S example data

Usage

```
data(otu_table_16S)
```

Format

```
data.frame
```

otu_table_ITS	<i>The OTU table of the ITS example data</i>
---------------	--

Description

The OTU table of the ITS example data

Usage

```
data(otu_table_ITS)
```

Format

```
data.frame
```

phylo_tree_16S	<i>The phylogenetic tree of 16S example data</i>
----------------	--

Description

The phylogenetic tree of 16S example data

Usage

```
data(phylo_tree_16S)
```

Format

data.frame

prok_func_FAPROTAX	<i>Customized FAPROTAX trait database</i>
--------------------	---

Description

Customized FAPROTAX trait database

Usage

```
data(prok_func_FAPROTAX)
```

Format

list

prok_func_NJC19_list	<i>The NJC19 database</i>
----------------------	---------------------------

Description

The NJC19 database

Usage

```
data(prok_func_NJC19_list)
```

Format

list

sample_info_16S	<i>The sample information of 16S example data</i>
-----------------	---

Description

The sample information of 16S example data

Usage

```
data(sample_info_16S)
```

Format

```
data.frame
```

sample_info_ITS	<i>The sample information of ITS example data</i>
-----------------	---

Description

The sample information of ITS example data

Usage

```
data(sample_info_ITS)
```

Format

```
data.frame
```

Tax4Fun2_KEGG	<i>The KEGG data files used in the trans_func class</i>
---------------	---

Description

The KEGG data files used in the trans_func class

Usage

```
data(Tax4Fun2_KEGG)
```

Format

```
list
```

taxonomy_table_16S	<i>The taxonomic information of 16S example data</i>
--------------------	--

Description

The taxonomic information of 16S example data

Usage

```
data(taxonomy_table_16S)
```

Format

data.frame

taxonomy_table_ITS	<i>The taxonomic information of ITS example data</i>
--------------------	--

Description

The taxonomic information of ITS example data

Usage

```
data(taxonomy_table_ITS)
```

Format

data.frame

tidy_taxonomy	<i>Clean up the taxonomic table to make taxonomic assignments consistent.</i>
---------------	---

Description

Clean up the taxonomic table to make taxonomic assignments consistent.

Usage

```
tidy_taxonomy(
  taxonomy_table,
  column = "all",
  pattern = c(".*unassigned.*", ".*uncultur.*", ".*unknown.*", ".*unidentif.*",
    ".*unclassified.*", ".*No blast hit.*", ".*Incertae.sedis.*"),
  replacement = "",
  ignore.case = TRUE,
  na_fill = ""
)
```

Arguments

taxonomy_table	a data.frame with taxonomic information (rows are features; columns are taxonomic levels); or a microtable object with tax_table in it.
column	default "all"; "all" or a number; 'all' represents cleaning up all the columns; a number represents cleaning up this specific column.
pattern	default c(".*unassigned.*", ".*uncultur.*", ".*unknown.*", ".*unidentif.*", ".*unclassified.*", ".*No blast hit.*", ".*Incertae.sedis.*"); the characters (regular expressions) to be removed or replaced; removed when parameter replacement = "", replaced when parameter replacement has something; Note that the capital and small letters are not distinguished when ignore.case = TRUE.
replacement	default ""; the characters used to replace the character in pattern parameter.
ignore.case	default TRUE; if FALSE, the pattern matching is case sensitive and if TRUE, case is ignored during matching.
na_fill	default ""; used to replace NA.

Format

`data.frame` object.

Value

`data.frame`

Examples

```
data("taxonomy_table_16S")
tidy_taxonomy(taxonomy_table_16S)
```

trans_abund

*Create trans_abund object for taxonomic abundance visualization.***Description**

This class is a wrapper for the taxonomic abundance transformations and visualization (e.g., bar plot, boxplot, heatmap, pie chart and line chart). The converted data style is the long-format for ggplot2 plot.

Methods**Public methods:**

- `trans_abund$new()`
- `trans_abund$plot_bar()`
- `trans_abund$plot_heatmap()`
- `trans_abund$plot_box()`
- `trans_abund$plot_line()`
- `trans_abund$plot_pie()`
- `trans_abund$plot_donut()`
- `trans_abund$plot_radar()`
- `trans_abund$plot_tern()`
- `trans_abund$print()`
- `trans_abund$clone()`

Method new():*Usage:*

```
trans_abund$new(
  dataset = NULL,
  taxrank = "Phylum",
  show = 0,
  ntaxa = 10,
  groupmean = NULL,
  group_morestats = FALSE,
  delete_taxonomy_lineage = TRUE,
  delete_taxonomy_prefix = TRUE,
  prefix = NULL,
  use_percentage = TRUE,
  input_taxaname = NULL,
  high_level = NULL,
  high_level_fix_nsub = NULL
)
```

Arguments:

`dataset` default NULL; the object of `microtable` class.

taxrank default "Phylum"; taxonomic level, i.e. a column name in tax_table of the input object. The function extracts the abundance from the taxa_abund list according to the names in the list. If the taxa_abund list is NULL, the function can automatically calculate the relative abundance to generate taxa_abund list.

show default 0; the mean relative abundance threshold for filtering the taxa with low abundance.

ntaxa default 10; how many taxa are selected to use. Taxa are ordered by abundance from high to low. This parameter does not conflict with the parameter show. Both can be used. ntaxa = NULL means the parameter will be invalid.

groupmean default NULL; calculate mean abundance for each group. Select a column name in microtable\$sample_table.

group_morestats default FALSE; only available when groupmean parameter is provided; Whether output more statistics for each group, including min, max, median and quantile; Thereinto, quantile25 and quantile75 denote 25% and 75% quantiles, respectively.

delete_taxonomy_lineage default TRUE; whether delete the taxonomy lineage in front of the target level.

delete_taxonomy_prefix default TRUE; whether delete the prefix of taxonomy, such as "g__".

prefix default NULL; character string; available when delete_taxonomy_prefix = T; default NULL represents using the "letter+__", e.g. "k__" for Phylum level; Please provide the customized prefix when it is not standard, otherwise the program can not correctly recognize it.

use_percentage default TRUE; whether show the abundance percentage. If TRUE, the abundance data will be multiplied by 100.

input_taxaname default NULL; character vector; input taxa names to select some taxa.

high_level default NULL; a taxonomic rank, such as "Phylum", used to add the taxonomic information of higher level. It is required for the legend with nested taxonomic levels in the bar plot or the higher taxonomic level in facets of y axis in the heatmap.

high_level_fix_nsub default NULL; an integer, used to fix the number of selected abundant taxa in each taxon from higher taxonomic level. If the total number under one taxon of higher level is less than the high_level_fix_nsub, the total number will be used. When high_level_fix_nsub is provided, the taxa number of higher level is calculated as: $\text{ceiling}(\text{ntaxa}/\text{high_level_fix_nsub})$. Note that ntaxa means either the parameter ntaxa or the taxonomic number obtained by filtering according to the show parameter.

Returns: data_abund stored in the object. The column 'all_mean_abund' represents mean relative abundance across all the samples. So the values in one taxon are all same across all the samples. If the sum of column 'Abundance' in one sample is larger than 1, the 'Abundance', 'SD' and 'SE' has been multiplied by 100.

Examples:

```
\donttest{
data(dataset)
t1 <- trans_abund$new(dataset = dataset, taxrank = "Phylum", ntaxa = 10)
}
```

Method plot_bar(): Bar plot.

Usage:

```
trans_abund$plot_bar(
  color_values = RColorBrewer::brewer.pal(8, "Dark2"),
```

```

bar_full = TRUE,
others_color = "grey90",
facet = NULL,
order_x = NULL,
x_axis_name = NULL,
barwidth = 0.9,
use_alluvium = FALSE,
clustering = FALSE,
clustering_plot = FALSE,
cluster_plot_width = 0.2,
facet_color = "grey95",
strip_text = 11,
legend_text_italic = FALSE,
xtext_angle = 0,
xtext_size = 10,
xtext_keep = TRUE,
xtitle_keep = TRUE,
ytitle_size = 17,
coord_flip = FALSE,
ggnested = FALSE,
high_level_add_other = FALSE,
sample_plot = NULL,
sample_plot_color = NULL,
sample_plot_height = NULL,
sample_plot_mainnames = FALSE,
bar_type = deprecated()
)

```

Arguments:

`color_values` default `RColorBrewer::brewer.pal(8, "Dark2")`; colors palette for the bars.

`bar_full` default `TRUE`; Whether the bar shows all the features (including 'Others'). Default `TRUE` means total abundance are summed to 1 or 100 (percentage). `FALSE` means 'Others' will not be shown.

`others_color` default "grey90"; the color for "Others" taxa.

`facet` default `NULL`; a character vector for the facet; group column name of `sample_table`, such as, "Group"; If multiple facets are needed, please provide ordered names, such as `c("Group", "Type")`. The latter should have a finer scale than the former one; Please adjust the facet orders in the plot by assigning factors in `sample_table` before creating `trans_abund` object or assigning factors in the `data_abund` table of `trans_abund` object. When multiple facets are used, please first install package `ggh4x` using the command `install.packages("ggh4x")`.

`order_x` default `NULL`; vector; used to order the sample names in x axis; must be the samples vector, such as `c("S1", "S3", "S2")`.

`x_axis_name` `NULL`; a character string; a column name of `sample_table` in dataset; used to show the sample names in x axis.

`barwidth` default 0.9; bar width, see width in `geom_bar` of `ggplot2` package.

`use_alluvium` default `FALSE`; whether add alluvium plot. If `TRUE`, please first install `ggalluvial` package.

clustering default FALSE; whether order samples by the clustering.
 clustering_plot default FALSE; whether add clustering plot. If clustering_plot = TRUE, clustering will be also TRUE in any case for the clustering.
 cluster_plot_width default 0.2, the dendrogram plot width; available when clustering_plot = TRUE.
 facet_color default "grey95"; facet background color.
 strip_text default 11; facet text size.
 legend_text_italic default FALSE; whether use italic in legend.
 xtext_angle default 0; number ranging from 0 to 90; used to adjust x axis text angle to reduce text overlap;
 xtext_size default 10; x axis text size.
 xtext_keep default TRUE; whether to keep the text on the x-axis.
 xtitle_keep default TRUE; whether to keep the title of the x-axis.
 ytitle_size default 17; y axis title size.
 coord_flip default FALSE; whether flip cartesian coordinates so that horizontal becomes vertical, and vertical becomes horizontal.
 ggnested default FALSE; whether use nested legend. Need ggnested package to be installed (<https://github.com/gmteunisse/ggnested>). To make it available, please assign high_level parameter when creating the object.
 high_level_add_other default FALSE; whether add 'Others' (all the unknown taxa) in each taxon of higher taxonomic level. Only available when ggnested = TRUE.
 sample_plot default NULL; Use the heatmap colors to represent sample information. The input should be column names from sample_table, e.g., c("Group", "pH").
 sample_plot_color default NULL; Color settings. The input must be a list that corresponds to sample_plot, e.g. list(Group = RColorBrewer::brewer.pal(6, "Set2"), pH = c("white", "red")). When the input factor is a numerical variable, it will be displayed with a color gradient; therefore, two colors should be provided for the input (as shown for "pH" above).
 sample_plot_height default NULL; Height of the sample heatmap; defaults to 1/10 of the main bar plot. The input must be a vector whose length equals that of sample_plot, e.g., c(0.1, 0.1).
 sample_plot_mainnames default FALSE; whether show the sample names in the main bar plot.
 bar_type deprecated. Please use bar_full argument instead.

Returns: ggplot2 object.

Examples:

```

\donttest{
t1$plot_bar(facet = "Group", xtext_keep = FALSE)
}

```

Method plot_heatmap(): Plot the heatmap.

Usage:

```

trans_abund$plot_heatmap(
  color_values = rev(RColorBrewer::brewer.pal(n = 11, name = "RdYlBu")),
  facet = NULL,

```

```

facet_switch = "y",
x_axis_name = NULL,
order_x = NULL,
withmargin = TRUE,
plot_numbers = FALSE,
plot_text_size = 4,
plot_breaks = NULL,
margincolor = "white",
plot_colorscale = "log10",
min_abundance = 0.01,
max_abundance = NULL,
strip_text = 11,
xtext_keep = TRUE,
xtext_angle = 0,
xtext_size = 10,
ytext_size = 11,
xtitle_keep = TRUE,
grid_clean = TRUE,
legend_title = "% Relative\nAbundance",
pheatmap = FALSE,
...
)

```

Arguments:

`color_values` default `rev(RColorBrewer::brewer.pal(n = 11, name = "RdYlBu"))`; colors palette for the plotting.

`facet` default `NULL`; a character vector for the facet; a group column name of `sample_table`, such as, "Group"; If multiple facets are needed, please provide ordered names, such as `c("Group", "Type")`. The latter should have a finer scale than the former one; Please adjust the facet orders in the plot by assigning factors in `sample_table` before creating `trans_abund` object or assigning factors in the `data_abund` table of `trans_abund` object. When multiple facets are used, please first install package `ggh4x` using the command `install.packages("ggh4x")`.

`facet_switch` default "y"; By default, the labels in facets are displayed on the top and right of the plot. If "x", the top labels will be displayed to the bottom. If "y", the right-hand side labels will be displayed to the left. Can also be set to "both". When the `high_level` is found in the object, the function will generate facets for the higher taxonomy in y axis. So the default "y" of the parameter is to make the visualization better when `high_level` is found. This parameter will be passed to the switch parameter in `ggplot2::facet_grid` or `ggh4x::facet_nested` function.

`x_axis_name` `NULL`; a character string; a column name of `sample_table` used to show the sample names in x axis.

`order_x` default `NULL`; vector; used to order the sample names in x axis; must be the samples vector, such as, `c("S1", "S3", "S2")`.

`withmargin` default `TRUE`; whether retain the tile margin.

`plot_numbers` default `FALSE`; whether plot the number in heatmap.

`plot_text_size` default 4; If `plot_numbers TRUE`, text size in plot.

`plot_breaks` default `NULL`; The legend breaks.

margincolor default "white"; If withmargin TRUE, use this as the margin color.

plot_colorscale default "log10"; color scale.

min_abundance default .01; the minimum abundance percentage in plot.

max_abundance default NULL; the maximum abundance percentage in plot, NULL represent the max percentage.

strip_text default 11; facet text size.

xtext_keep default TRUE; whether to keep the text on the x-axis.

xtext_angle default 0; number ranging from 0 to 90; used to adjust x axis text angle to reduce text overlap;

xtext_size default 10; x axis text size.

ytext_size default 11; y axis text size.

xtitle_keep default TRUE; whether to keep the title of the x-axis.

grid_clean default TRUE; whether remove grid lines.

legend_title default "% Relative\nAbundance"; legend title text.

pheatmap default FALSE; whether use pheatmap package to plot the heatmap.

... parameters pass to pheatmap when pheatmap = TRUE.

Returns: ggplot2 object or grid object based on pheatmap.

Examples:

```
\donttest{
t1 <- trans_abund$new(dataset = dataset, taxrank = "Genus", ntaxa = 40)
t1$plot_heatmap(facet = "Group", xtext_keep = FALSE, withmargin = FALSE)
}
```

Method plot_box(): Box plot.

Usage:

```
trans_abund$plot_box(
  color_values = RColorBrewer::brewer.pal(8, "Dark2"),
  group = NULL,
  show_point = FALSE,
  point_color = "black",
  point_size = 3,
  point_alpha = 0.3,
  plot_flip = FALSE,
  boxfill = TRUE,
  middlecolor = "grey95",
  middlesize = 1,
  xtext_angle = 0,
  xtext_size = 10,
  ytitle_size = 17,
  ...
)
```

Arguments:

color_values default RColorBrewer::brewer.pal(8, "Dark2"); colors palette for the box.

group default NULL; a column name of sample table to show abundance across groups.

show_point default FALSE; whether show points in plot.
 point_color default "black"; If show_point TRUE; use the color
 point_size default 3; If show_point TRUE; use the size
 point_alpha default .3; If show_point TRUE; use the transparency.
 plot_flip default FALSE; Whether rotate plot.
 boxfill default TRUE; Whether fill the box with colors.
 middlecolor default "grey95"; The middle line color.
 middlesize default 1; The middle line size.
 xtext_angle default 0; number ranging from 0 to 90; used to adjust x axis text angle to reduce text overlap;
 xtext_size default 10; x axis text size.
 ytitle_size default 17; y axis title size.
 ... parameters pass to geom_boxplot function.

Returns: ggplot2 object.

Examples:

```
\donttest{
t1$plot_box(group = "Group")
}
```

Method plot_line(): Plot the line chart.

Usage:

```
trans_abund$plot_line(
  color_values = RColorBrewer::brewer.pal(8, "Dark2"),
  plot_SE = TRUE,
  position = position_dodge(0.1),
  errorbar_size = 1,
  errorbar_width = 0.1,
  point_size = 3,
  point_alpha = 0.8,
  line_size = 0.8,
  line_alpha = 0.8,
  line_type = 1,
  xtext_angle = 0,
  xtext_size = 10,
  ytitle_size = 17
)
```

Arguments:

color_values default RColorBrewer::brewer.pal(8, "Dark2"); colors palette for the points and lines.
 plot_SE default TRUE; TRUE: the errorbar is *meanse*; FALSE: the errorbar is *meansd*.
 position default position_dodge(0.1); Position adjustment, either as a string (such as "identity"), or the result of a call to a position adjustment function.
 errorbar_size default 1; errorbar line size.
 errorbar_width default 0.1; errorbar width.

point_size default 3; point size for taxa.
 point_alpha default 0.8; point transparency.
 line_size default 0.8; line size.
 line_alpha default 0.8; line transparency.
 line_type default 1; an integer; line type.
 xtext_angle default 0; number ranging from 0 to 90; used to adjust x axis text angle to reduce text overlap;
 xtext_size default 10; x axis text size.
 ytitle_size default 17; y axis title size.

Returns: ggplot2 object.

Examples:

```
\donttest{
t1 <- trans_abund$new(dataset = dataset, taxrank = "Genus", ntaxa = 5)
t1$plot_line(point_size = 3)
t1 <- trans_abund$new(dataset = dataset, taxrank = "Genus", ntaxa = 5, groupmean = "Group")
t1$plot_line(point_size = 5, errorbar_size = 1, xtext_angle = 30)
}
```

Method plot_pie(): Pie chart.

Usage:

```
trans_abund$plot_pie(
  color_values = RColorBrewer::brewer.pal(8, "Dark2"),
  facet_nrow = 1,
  strip_text = 11,
  add_label = FALSE,
  legend_text_italic = FALSE
)
```

Arguments:

color_values default RColorBrewer::brewer.pal(8, "Dark2"); colors palette for each section.
 facet_nrow default 1; how many rows in the plot.
 strip_text default 11; sample title size.
 add_label default FALSE; Whether add the percentage label in each section of pie chart.
 legend_text_italic default FALSE; whether use italic in legend.

Returns: ggplot2 object.

Examples:

```
\donttest{
t1 <- trans_abund$new(dataset = dataset, taxrank = "Phylum", ntaxa = 6, groupmean = "Group")
t1$plot_pie(facet_nrow = 1)
}
```

Method plot_donut(): Donut chart based on the ggpubr::ggdonutchart function.

Usage:

```
trans_abund$plot_donut(
  color_values = RColorBrewer::brewer.pal(8, "Dark2"),
  label = TRUE,
  facet_nrow = 1,
  legend_text_italic = FALSE,
  ...
)
```

Arguments:

color_values default RColorBrewer::brewer.pal(8, "Dark2"); colors palette for the donut.

label default TRUE; whether show the percentage label.

facet_nrow default 1; how many rows in the plot.

legend_text_italic default FALSE; whether use italic in legend.

... parameters passed to ggpubr::ggdonutchart.

Returns: combined ggplot2 objects list, generated by ggpubr::ggarrange function.

Examples:

```
\dontrun{
t1 <- trans_abund$new(dataset = dataset, taxrank = "Phylum", ntaxa = 6, groupmean = "Group")
t1$plot_donut(label = TRUE)
}
```

Method plot_radar(): Radar chart based on the ggradar package (<https://github.com/ricardobion/ggradar>).

Usage:

```
trans_abund$plot_radar(
  color_values = RColorBrewer::brewer.pal(8, "Dark2"),
  ...
)
```

Arguments:

color_values default RColorBrewer::brewer.pal(8, "Dark2"); colors palette for samples.

... parameters passed to ggradar::ggradar function except group.colours parameter.

Returns: ggplot2 object.

Examples:

```
\dontrun{
t1 <- trans_abund$new(dataset = dataset, taxrank = "Phylum", ntaxa = 6, groupmean = "Group")
t1$plot_radar()
}
```

Method plot_tern(): Ternary diagrams based on the ggtern package.

Usage:

```
trans_abund$plot_tern(
  color_values = RColorBrewer::brewer.pal(8, "Dark2"),
  color_legend_guide_size = 4
)
```

Arguments:

color_values default RColorBrewer::brewer.pal(8, "Dark2"); colors palette for the samples.

color_guide_size default 4; The size of legend guide for color.

Returns: ggplot2 object.

Examples:

```
\dontrun{
t1 <- trans_abund$new(dataset = dataset, taxrank = "Phylum", ntaxa = 6, groupmean = "Group")
t1$plot_tern()
}
```

Method print(): Print the trans_abund object.

Usage:

```
trans_abund$print()
```

Method clone(): The objects of this class are cloneable with this method.

Usage:

```
trans_abund$clone(deep = FALSE)
```

Arguments:

deep Whether to make a deep clone.

Examples

```
## -----
## Method `trans_abund$new`
## -----

data(dataset)
t1 <- trans_abund$new(dataset = dataset, taxrank = "Phylum", ntaxa = 10)

## -----
## Method `trans_abund$plot_bar`
## -----

t1$plot_bar(facet = "Group", xtext_keep = FALSE)

## -----
## Method `trans_abund$plot_heatmap`
## -----

t1 <- trans_abund$new(dataset = dataset, taxrank = "Genus", ntaxa = 40)
t1$plot_heatmap(facet = "Group", xtext_keep = FALSE, withmargin = FALSE)

## -----
```

```

## Method `trans_abund$plot_box`
## -----

t1$plot_box(group = "Group")

## -----
## Method `trans_abund$plot_line`
## -----

t1 <- trans_abund$new(dataset = dataset, taxrank = "Genus", ntaxa = 5)
t1$plot_line(point_size = 3)
t1 <- trans_abund$new(dataset = dataset, taxrank = "Genus", ntaxa = 5, groupmean = "Group")
t1$plot_line(point_size = 5, errorbar_size = 1, xtext_angle = 30)

## -----
## Method `trans_abund$plot_pie`
## -----

t1 <- trans_abund$new(dataset = dataset, taxrank = "Phylum", ntaxa = 6, groupmean = "Group")
t1$plot_pie(facet_nrow = 1)

## -----
## Method `trans_abund$plot_donut`
## -----

## Not run:
t1 <- trans_abund$new(dataset = dataset, taxrank = "Phylum", ntaxa = 6, groupmean = "Group")
t1$plot_donut(label = TRUE)

## End(Not run)

## -----
## Method `trans_abund$plot_radar`
## -----

## Not run:
t1 <- trans_abund$new(dataset = dataset, taxrank = "Phylum", ntaxa = 6, groupmean = "Group")
t1$plot_radar()

## End(Not run)

## -----
## Method `trans_abund$plot_tern`
## -----

## Not run:
t1 <- trans_abund$new(dataset = dataset, taxrank = "Phylum", ntaxa = 6, groupmean = "Group")

```

```
t1$plot_tern()

## End(Not run)
```

trans_alpha	Create trans_alpha object for alpha diversity statistics and visualization.
-------------	---

Description

This class is a wrapper for a series of alpha diversity analysis, including the statistics and visualization.

Methods

Public methods:

- `trans_alpha$new()`
- `trans_alpha$cal_diff()`
- `trans_alpha$plot_alpha()`
- `trans_alpha$print()`
- `trans_alpha$clone()`

Method `new()`:

Usage:

```
trans_alpha$new(
  dataset = NULL,
  group = NULL,
  by_group = NULL,
  by_ID = NULL,
  order_x = NULL
)
```

Arguments:

dataset `microtable` object.

group default NULL; a column name of sample_table in the input microtable object used for the statistics across groups.

by_group default NULL; a column name of sample_table used to perform the differential test among groups (from group parameter) for each group (from by_group parameter) separately.

by_ID default NULL; a column name of sample_table used to perform paired T test or paired Wilcoxon test for the paired data, such as continuous sampling of individual animals or plant compartments for different plant species (ID). So by_ID in sample_table should be the smallest unit of sample collection without any repetition in it. When the by_ID parameter is provided, the function can automatically perform paired test, and no more parameters is required.

order_x default NULL; a column name of sample_table or a vector with sample names. If provided, sort samples using factor.

Returns: data_alpha and data_stat stored in the object.

Examples:

```
\donttest{
data(dataset)
t1 <- trans_alpha$new(dataset = dataset, group = "Group")
}
```

Method cal_diff(): Differential test on alpha diversity.

Usage:

```
trans_alpha$cal_diff(
  measure = NULL,
  method = c("KW", "KW_dunn", "wilcox", "t.test", "anova", "scheirerRayHare", "lm",
    "lme", "betareg", "glmm", "glmm_beta")[1],
  formula = NULL,
  p_adjust_method = "fdr",
  KW_dunn_letter = TRUE,
  alpha = 0.05,
  anova_post_test = "duncan.test",
  anova_varequal_test = FALSE,
  return_model = FALSE,
  ...
)
```

Arguments:

measure default NULL; character vector; If NULL, all indexes will be used; see names of microtable\$alpha_diversity, e.g. c("Observed", "Chao1", "Shannon").

method default "KW"; see the following available options:

'KW' Kruskal-Wallis Rank Sum Test for all groups (≥ 2)

'KW_dunn' Dunn's Kruskal-Wallis Multiple Comparisons <10.1080/00401706.1964.10490181> based on dunnTest function in FSA package

'wilcox' Wilcoxon Rank Sum Test for all paired groups When by_ID parameter is provided in creating the object of the class, paired Wilcoxon test will be performed.

't.test' Student's t-Test for all paired groups. When by_ID parameter is provided in creating the object of the class, paired t-test will be performed.

'anova' Variance analysis. For one-way anova, the default post hoc test is Duncan's new multiple range test. Please use anova_post_test parameter to change the post hoc method. For multi-way anova, Please use formula parameter to specify the model and see [aov](#) for more details

'scheirerRayHare' Scheirer-Ray-Hare test (nonparametric test) for a two-way factorial experiment; see scheirerRayHare function of rcompanion package

'lm' Linear Model based on the lm function

'lme' Linear Mixed Effect Model based on the lmerTest package

'betareg' Beta Regression for Rates and Proportions based on the betareg package

'glmm' Generalized linear mixed model (GLMM) based on the glmmTMB package. A family function can be provided using parameter passing, such as: `family = glmmTMB::lognormal(link = "log")`

'glmm_beta' Generalized linear mixed model (GLMM) with a family function of beta distribution. This is an extension of the GLMM model in 'glmm' option. The only difference is in glmm_beta the family function is fixed with the beta distribution function, facilitating the fitting for proportional data (ranging from 0 to 1). The link function is fixed with "logit".

formula default NULL; applied to two-way or multi-factor analysis when method is "anova", "scheirerRayHare", "lm", "lme", "betareg" or "glmm"; specified set for independent variables, i.e. the latter part of a general formula, such as 'block + N*P*K'.

p_adjust_method default "fdr" (for "KW", "wilcox", "t.test" methods) or "holm" (for "KW_dunn"); P value adjustment method; For method = 'KW', 'wilcox' or 't.test', please see method parameter of p.adjust function for available options; For method = 'KW_dunn', please see dunn.test::p.adjustment.methods for available options.

KW_dunn_letter default TRUE; For method = 'KW_dunn', TRUE denotes significances are presented by letters; FALSE means significances are shown by asterisk for paired comparison.

alpha default 0.05; Significant level; used for generating significance letters when method is 'anova' or 'KW_dunn'.

anova_post_test default "duncan.test". The post hoc test method for one-way anova. The default option represents the Duncan's new multiple range test. Other available options include "LSD.test" (LSD post hoc test) and "HSD.test" (HSD post hoc test). All those are the function names from agricolae package.

anova_varequal_test default FALSE; whether conduct Levene's Test for equality of variances. Only available for one-way anova. Significant P value means the variance among groups is not equal.

return_model default FALSE; whether return the original "lm", "lmer" or "glmm" model list in the object.

... parameters passed to `kruskal.test` (when method = "KW") or `wilcox.test` function (when method = "wilcox") or `dunnTest` function of FSA package (when method = "KW_dunn") or `agricolae::duncan.test/agricolae::LSD.test/agricolae::HSD.test` (when method = "anova", one-way anova) or `rcompanion::scheirerRayHare` (when method = "scheirerRayHare") or `stats::lm` (when method = "lm") or `lmerTest::lmer` (when method = "lme") or `betareg::betareg` (when method = "betareg") or `glmmTMB::glmmTMB` (when method = "glmm").

Returns: `res_diff`, stored in object with the format `data.frame`.

When method is "betareg", "lm", "lme" or "glmm", "Estimate" and "Std.Error" columns represent the fitted coefficient and its standard error, respectively.

Examples:

```
\donttest{
t1$cal_diff(method = "KW")
t1$cal_diff(method = "anova")
t1 <- trans_alpha$new(dataset = dataset, group = "Type", by_group = "Group")
t1$cal_diff(method = "anova")
}
```

Method plot_alpha(): Plot the alpha diversity. Box plot (and others for visualizing data in groups of single factor) is used for the visualization of alpha diversity when the group is found in

the object. When the formula is found in the `res_diff` table in the object, heatmap is employed automatically to show the significances of differential test for multiple indexes, and errorbar (coefficient and standard errors) can be used for single index.

Usage:

```
trans_alpha$plot_alpha(
  plot_type = "ggboxplot",
  color_values = RColorBrewer::brewer.pal(8, "Dark2"),
  measure = "Shannon",
  group = NULL,
  add = NULL,
  add_sig = TRUE,
  add_sig_label = "Significance",
  add_sig_text_size = 3.88,
  add_sig_label_num_dec = 4,
  order_x_mean = FALSE,
  y_start = 0.1,
  y_increase = 0.05,
  xtext_angle = 30,
  xtext_size = 13,
  ytitle_size = 17,
  bar_width = 0.9,
  bar_alpha = 0.8,
  dodge_width = 0.9,
  plot_SE = TRUE,
  errorbar_size = 1,
  errorbar_width = 0.2,
  errorbar_addpoint = TRUE,
  errorbar_color_black = FALSE,
  point_size = 3,
  point_alpha = 0.8,
  add_line = FALSE,
  line_size = 0.8,
  line_type = 2,
  line_color = "grey50",
  line_alpha = 0.5,
  heatmap_cell = "P.unadj",
  heatmap_sig = "Significance",
  heatmap_x = "Factors",
  heatmap_y = "Measure",
  heatmap_lab_fill = "P value",
  coefplot_sig_pos = 2,
  ...
)
```

Arguments:

`plot_type` default "ggboxplot"; plot type; available options include "ggboxplot", "ggdotplot", "ggviolin", "ggstripchart", "ggerrorplot", "errorbar" and "barerrorbar". The options starting with "gg" are function names coming from ggpubr package. All those methods with ggpubr package use the `data_alpha` table in the object. "errorbar" represents Mean±SD or

Mean±SE plot based on ggplot2 package by invoking the data_stat table in the object.

"bareerrorbar" denotes "bar plot + error bar". It is similar with "errorbar" and has a bar plot.

color_values default RColorBrewer::brewer.pal(8, "Dark2"); color pallete for groups.

measure default "Shannon"; one alpha diversity index in the object.

group default NULL; group name used for the plot.

add default NULL; add another plot element; passed to the add parameter of the function (e.g., ggboxplot) from ggpubr package when plot_type starts with "gg" (functions coming from ggpubr package).

add_sig default TRUE; whether add significance label using the result of cal_diff function, i.e. object\$res_diff; This is mainly designed to add post hoc test of anova or other significances to make the label mapping easy.

add_sig_label default "Significance"; select a colname of object\$res_diff for the label text when 'Letter' is not in the table, such as 'P.adj' or 'Significance'.

add_sig_text_size default 3.88; the size of text in added label.

add_sig_label_num_dec default 4; reserved decimal places when the parameter add_sig_label use numeric column, like 'P.adj'.

order_x_mean default FALSE; whether order x axis by the means of groups from large to small.

y_start default 0.1; the y axis value from which to add the significance asterisk label; the default 0.1 means $\max(\text{values}) + 0.1 * (\max(\text{values}) - \min(\text{values}))$.

y_increase default 0.05; the increasing y axis space to add the label (asterisk or letter); the default 0.05 means $0.05 * (\max(\text{values}) - \min(\text{values}))$; this parameter is also used to label the letters of anova result with the fixed space.

xtext_angle default 30; number (e.g. 30). Angle of text in x axis.

xtext_size default 13; x axis text size. NULL means the default size in ggplot2.

ytitle_size default 17; y axis title size.

bar_width default 0.9; the bar width when plot_type = "bareerrorbar".

bar_alpha default 0.8; the alpha of bar color when plot_type = "bareerrorbar".

dodge_width default 0.9; the dodge width used in position_dodge function of ggplot2 package when plot_type is "errorbar" or "bareerrorbar".

plot_SE default TRUE; TRUE: the errorbar is *meanse*; FALSE: the errorbar is *meansd*. Available when plot_type is "errorbar" or "bareerrorbar".

errorbar_size default 1; errorbar size. Available when plot_type is "errorbar" or "bareerrorbar".

errorbar_width default 0.2; errorbar width. Available when plot_type is "errorbar" or "bareerrorbar" and by_group is NULL.

errorbar_addpoint default TRUE; whether add point for mean. Available when plot_type is "errorbar" or "bareerrorbar" and by_group is NULL.

errorbar_color_black default FALSE; whether use black for the color of errorbar when plot_type is "errorbar" or "bareerrorbar".

point_size default 3; point size for taxa. Available when plot_type is "errorbar" or "bareerrorbar".

point_alpha default 0.8; point transparency. Available when plot_type is "errorbar" or "bareerrorbar".

`add_line` default FALSE; whether add line. Available when `plot_type` is "errorbar" or "bar-errorbar".
`line_size` default 0.8; line size when `add_line` = TRUE. Available when `plot_type` is "errorbar" or "bareerrorbar".
`line_type` default 2; an integer; line type when `add_line` = TRUE. The available case is same with `line_size`.
`line_color` default "grey50"; line color when `add_line` = TRUE. Available when `by_group` is NULL. Other available case is same with `line_size`.
`line_alpha` default 0.5; line transparency when `add_line` = TRUE. The available case is same with `line_size`.
`heatmap_cell` default "P.unadj"; the column of `res_diff` table for the cell of heatmap when formula with multiple factors is found in the method.
`heatmap_sig` default "Significance"; the column of `res_diff` for the significance label of heatmap.
`heatmap_x` default "Factors"; the column of `res_diff` for the x axis of heatmap.
`heatmap_y` default "Taxa"; the column of `res_diff` for the y axis of heatmap.
`heatmap_lab_fill` default "P value"; legend title of heatmap.
`coefplot_sig_pos` default 2; Significance label position in the coefficient point and errorbar plot. The formula is $\text{Estimate} + \text{coefplot_sig_pos} * \text{Std.Error}$. This plot is used when there is only one measure found in the table, and 'Estimate' and 'Std.Error' are both in the column names (such as for `lm` and `lme` methods). The x axis is 'Estimate', and y axis denotes 'Factors'. When `coefplot_sig_pos` is a negative value, the label is in the left of the errorbar. Errorbar size and width in the coefficient point plot can be adjusted with the parameters `errorbar_size` and `errorbar_width`. Point size and alpha can be adjusted with parameters `point_size` and `point_alpha`. The significance label size can be adjusted with parameter `add_sig_text_size`. Furthermore, the vertical line around 0 can be adjusted with parameters `line_size`, `line_type`, `line_color` and `line_alpha`.
... parameters passing to `ggpubr::ggboxplot` function (or other functions shown by `plot_type` parameter when it starts with "gg") or `plot_cor` function in [trans_env](#) class for the heatmap of multiple factors when formula is found in the `res_diff` of the object.

Returns: ggplot.

Examples:

```

\donttest{
t1 <- trans_alpha$new(dataset = dataset, group = "Group")
t1$cal_diff(method = "wilcox")
t1$plot_alpha(measure = "Shannon", add_sig = TRUE)
t1 <- trans_alpha$new(dataset = dataset, group = "Type", by_group = "Group")
t1$cal_diff(method = "wilcox")
t1$plot_alpha(measure = "Shannon", add_sig = TRUE)
}

```

Method `print()`: Print the `trans_alpha` object.

Usage:

```
trans_alpha$print()
```

Method `clone()`: The objects of this class are cloneable with this method.

Usage:

```
trans_alpha$clone(deep = FALSE)
```

Arguments:

deep Whether to make a deep clone.

Examples

```
## -----
## Method `trans_alpha$new`
## -----

data(dataset)
t1 <- trans_alpha$new(dataset = dataset, group = "Group")

## -----
## Method `trans_alpha$cal_diff`
## -----

t1$cal_diff(method = "KW")
t1$cal_diff(method = "anova")
t1 <- trans_alpha$new(dataset = dataset, group = "Type", by_group = "Group")
t1$cal_diff(method = "anova")

## -----
## Method `trans_alpha$plot_alpha`
## -----

t1 <- trans_alpha$new(dataset = dataset, group = "Group")
t1$cal_diff(method = "wilcox")
t1$plot_alpha(measure = "Shannon", add_sig = TRUE)
t1 <- trans_alpha$new(dataset = dataset, group = "Type", by_group = "Group")
t1$cal_diff(method = "wilcox")
t1$plot_alpha(measure = "Shannon", add_sig = TRUE)
```

trans_beta

*Create trans_beta object for beta-diversity analysis***Description**

This class is a wrapper for a series of beta-diversity related analysis, including ordination analysis based on An et al. (2019) <doi:10.1016/j.geoderma.2018.09.035>, group distance comparison, clustering, perMANOVA based on Anderson al. (2008) <doi:10.1111/j.1442-9993.2001.01070.pp.x>, ANOSIM and PERMDISP. Note that the beta diversity analysis methods related with environmental variables are encapsulated within the trans_env class.

Methods

Public methods:

- `trans_beta$new()`
- `trans_beta$cal_ordination()`
- `trans_beta$plot_ordination()`
- `trans_beta$cal_manova()`
- `trans_beta$cal_anosim()`
- `trans_beta$cal_betadisper()`
- `trans_beta$cal_group_distance()`
- `trans_beta$cal_group_distance_diff()`
- `trans_beta$plot_group_distance()`
- `trans_beta$plot_clustering()`
- `trans_beta$clone()`

Method `new()`:

Usage:

```
trans_beta$new(dataset = NULL, measure = NULL, group = NULL)
```

Arguments:

`dataset` an object of `microtable` class.

`measure` default NULL; a matrix name stored in `microtable$beta_diversity` list, such as "bray" or "jaccard", or a customized matrix; used for ordination, manova, group distance comparison, etc.; Please see `cal_betadiv` function of `microtable` class for more details.

`group` default NULL; a column name of `sample_table` in the input dataset; group information will be used for manova, betadisper or distance comparison.

Returns: measure, group and dataset stored in the object.

Examples:

```
data(dataset)
t1 <- trans_beta$new(dataset = dataset, measure = "bray", group = "Group")
```

Method `cal_ordination()`: Unconstrained ordination.

Usage:

```
trans_beta$cal_ordination(
  method = "PCoA",
  ncomp = 2,
  taxa_level = NULL,
  NMDS_matrix = TRUE,
  trans = FALSE,
  scale_species = FALSE,
  scale_species_ratio = 0.8,
  orthoI = NA,
  ordination = deprecated(),
  ...
)
```

Arguments:

method default "PCoA"; "PCoA", "NMDS", "PCA", "DCA", "PLS-DA" or "OPLS-DA". PCoA: principal coordinates analysis; NMDS: non-metric multidimensional scaling, PCA: principal component analysis; DCA: detrended correspondence analysis; PLS-DA: partial least squares discriminant analysis; OPLS-DA: orthogonal partial least squares discriminant analysis. For the methods details, please refer to the papers <doi:10.1111/j.1574-6941.2007.00375.x> (for PCoA, NMDS, PCA and DCA) and <doi:10.1186/s12859-019-3310-7> (for PLS-DA or OPLS-DA).

ncomp default 2; dimensions in the result. For the method option "PCA", "PCoA" or "DCA", the corresponding dimension information will be selected from the original model based on this parameter. For all the dimension information, please refer to `model` in the results. For the method option "NMDS", this argument will be passed to the `k` parameter in the `vegan::metaMDS` function.

taxa_level default NULL; available for PCA, DCA or NMDS (`NMDS_matrix = TRUE`). Default NULL means using the `otu_table` in the `microtable` object. For other options, please provide the taxonomic rank names in `tax_table`, such as "Phylum" or "Genus". In such cases, the data will be merged according to the provided taxonomic levels to generate a new abundance table.

NMDS_matrix default TRUE; For the NMDS method, whether use a distance matrix as input like PCoA. If it is FALSE, the input will be the abundance table like PCA.

trans default FALSE; whether species abundance will be square root transformed; only available when method is "PCA" or "DCA". For method "NMDS" and `NMDS_matrix = FALSE`, please set the `autotransform` parameter, which will be passed to `vegan::metaMDS` function directly.

scale_species default FALSE; whether species loading in PCA, DCA or NMDS (`NMDS_matrix = FALSE`) is scaled.

scale_species_ratio default 0.8; the ratio to scale up the loading; multiply by the maximum distance between samples and origin. Only available when `scale_species = TRUE`.

orthoI default NA; number of orthogonal components (for OPLS-DA only). Default NA means the number of orthogonal components is automatically computed. Please also see `orthoI` parameter in `opls` function of `ropls` package.

ordination deprecated. Please use `method` argument instead.

... parameters passed to `vegan::rda` function when `method = "PCA"`, or `vegan::decorana` function when `method = "DCA"`, or `ape::pcoa` function when `method = "PCoA"`, or `vegan::metaMDS` function when `method = "NMDS"`, or `ropls::opls` function when `method = "PLS-DA"` or `method = "OPLS-DA"`.

Returns: `res_ordination` list stored in the object. In the list, `model` is the original analysis results; `scores` is the sample scores table; `loading` is the feature loading table.

Examples:

```
t1$cal_ordination(method = "PCoA")
```

Method `plot_ordination()`: Plot the ordination result.

Usage:

```
trans_beta$plot_ordination(
  plot_type = "point",
  choices = c(1, 2),
```

```

color_values = RColorBrewer::brewer.pal(8, "Dark2"),
shape_values = c(16, 17, 7, 8, 15, 18, 11, 10, 12, 13, 9, 3, 4, 0, 1, 2, 14),
plot_color = NULL,
plot_shape = NULL,
plot_group_order = NULL,
add_sample_label = NULL,
point_size = 3,
point_alpha = 0.8,
centroid_segment_alpha = 0.6,
centroid_segment_size = 1,
centroid_segment_linetype = 3,
ellipse_chull_fill = TRUE,
ellipse_chull_alpha = 0.1,
ellipse_level = 0.9,
ellipse_type = "t",
NMDS_stress_pos = c(1, 1),
NMDS_stress_text_prefix = "",
loading_arrow = FALSE,
loading_taxa_num = 10,
loading_text_taxlevel = NULL,
loading_text_color = "black",
loading_arrow_color = "grey30",
loading_text_size = 3,
loading_text_prefix = FALSE,
loading_text_italic = FALSE
)

```

Arguments:

`plot_type` default "point"; one or more elements of "point", "ellipse", "chull" and "centroid".

'point' add sample points

'ellipse' add confidence ellipse for points of each group

'chull' add convex hull for points of each group

'centroid' add centroid line for points in each group

`choices` default c(1, 2); selected axis for the visualization; must be numeric vector. The maximum value must not exceed the parameter `ncomp` in the `cal_ordination` function.

`color_values` default `RColorBrewer::brewer.pal(8, "Dark2")`; colors palette for different groups.

`shape_values` default `c(16, 17, 7, 8, 15, 18, 11, 10, 12, 13, 9, 3, 4, 0, 1, 2, 14)`; a vector for point shape types of groups, see `ggplot2` tutorial.

`plot_color` default `NULL`; a colname of `sample_table` to assign colors to different groups in plot.

`plot_shape` default `NULL`; a colname of `sample_table` to assign shapes to different groups in plot.

`plot_group_order` default `NULL`; a vector used to order the groups in the legend of plot.

`add_sample_label` default `NULL`; a column name in `sample_table`; If provided, show the point name in plot.

`point_size` default 3; point size when "point" is in `plot_type` parameter. `point_size` can also be a variable name in `sample_table`, such as "pH".

point_alpha default .8; point transparency in plot when "point" is in plot_type parameter.

centroid_segment_alpha default 0.6; segment transparency in plot when "centroid" is in plot_type parameter.

centroid_segment_size default 1; segment size in plot when "centroid" is in plot_type parameter.

centroid_segment_linetype default 3; the line type related with centroid in plot when "centroid" is in plot_type parameter.

ellipse_chull_fill default TRUE; whether fill colors to the area of ellipse or chull.

ellipse_chull_alpha default 0.1; color transparency in the ellipse or convex hull depending on whether "ellipse" or "centroid" is in plot_type parameter.

ellipse_level default .9; confidence level of ellipse when "ellipse" is in plot_type parameter.

ellipse_type default "t"; ellipse type when "ellipse" is in plot_type parameter; see type in stat_ellipse.

NMDS_stress_pos default c(1, 1); a numerical vector with two values used to represent the insertion position of the stress text. The first one denotes the x-axis, while the second one corresponds to the y-axis. The assigned position is determined by multiplying the respective value with the maximum point on the corresponding coordinate axis. Thus, the x-axis position is equal to $\max(\text{points of x axis}) * \text{NMDS_stress_pos}[1]$, and the y-axis position is equal to $\max(\text{points of y axis}) * \text{NMDS_stress_pos}[2]$. Negative values can also be utilized for the negative part of the axis. NMDS_stress_pos = NULL denotes no stress text to show.

NMDS_stress_text_prefix default ""; If NMDS_stress_pos is not NULL, this parameter can be used to add text in front of the stress value.

loading_arrow default FALSE; whether show the loading using arrow.

loading_taxa_num default 10; the number of taxa used for the loading. Only available when loading_arrow = TRUE.

loading_text_taxlevel default NULL; which level of taxonomic table will be used. Default NULL means using the taxa_level parameter in the previous cal_ordination function.

loading_text_color default "black"; the color of taxa text. Only available when loading_arrow = TRUE.

loading_arrow_color default "grey30"; the color of taxa arrow. Only available when loading_arrow = TRUE.

loading_text_size default 3; the size of taxa text. Only available when loading_arrow = TRUE.

loading_text_prefix default FALSE; whether show the prefix (e.g., g__) in the taxa text. Only available when loading_arrow = TRUE.

loading_text_italic default FALSE; whether using italic for the taxa text. Only available when loading_arrow = TRUE.

Returns: ggplot.

Examples:

```
t1$plot_ordination(plot_type = "point")
t1$plot_ordination(plot_color = "Group", plot_shape = "Group", plot_type = "point")
t1$plot_ordination(plot_color = "Group", plot_type = c("point", "ellipse"))
t1$plot_ordination(plot_color = "Group", plot_type = c("point", "centroid"),
  centroid_segment_linetype = 1)
```

Method `cal_manova()`: Calculate perMANOVA (Permutational Multivariate Analysis of Variance) based on the `adonis2` function of `vegan` package <doi:10.1111/j.1442-9993.2001.01070.pp.x>.

Usage:

```
trans_beta$cal_manova(
  manova_all = TRUE,
  manova_set = NULL,
  group = NULL,
  by_group = NULL,
  p_adjust_method = "fdr",
  by = "terms",
  by_auto_set = TRUE,
  permutations = 999,
  ...
)
```

Arguments:

`manova_all` default TRUE; TRUE represents test for all the groups, i.e. the overall test; FALSE represents test for all the paired groups.

`manova_set` default NULL; other specified group set for manova, such as "Group + Type" and "Group*Type". Please also see the `formula` parameter (only right-hand side) in `adonis2` function of `vegan` package. The parameter `manova_set` has higher priority than `manova_all` parameter. If `manova_set` is provided; `manova_all` is disabled.

`group` default NULL; a column name of `sample_table` used for manova. If NULL, search group variable stored in the object. Available when `manova_set` is not provided.

`by_group` default NULL; one column name in `sample_table`; used to perform paired comparisons within each group. Only available when `manova_all` = FALSE and `manova_set` is not provided.

`p_adjust_method` default "fdr"; p.adjust method; available when `manova_all` = FALSE; see `method` parameter of `p.adjust` function for available options.

`by` default "terms"; same with the `by` parameter in `adonis2` function of `vegan` package.

`by_auto_set` default TRUE; Whether automatically set the options for `by` parameter ("marginal" or "terms") when `manova_set` is provided. The primary reason for setting this parameter is that using marginal effects (also known as "Type III" effects) is more robust for unbalanced experimental designs. Since the option `by = "margin"` in the `adonis2` function ignores main effects when interaction effects are present, we automatically set `by = "margin"` when there are no interaction effects, and set `by = "terms"` when interaction effects exist. If the user wants to use parameter `by`, please set `by_auto_set` = FALSE. Note that this parameter is only available when `manova_set` is provided.

`permutations` default 999; same with the `permutations` parameter in `adonis2` function of `vegan` package.

... parameters passed to `adonis2` function of `vegan` package.

Returns: `res_manova` stored in object with `data.frame` class.

Examples:

```
t1$cal_manova(manova_all = TRUE)
```

Method `cal_anosim()`: Analysis of similarities (ANOSIM) based on the `anosim` function of `vegan` package.

Usage:

```
trans_beta$cal_anosim(
  paired = FALSE,
  group = NULL,
  by_group = NULL,
  p_adjust_method = "fdr",
  permutations = 999,
  ...
)
```

Arguments:

`paired` default FALSE; whether perform paired test between any two combined groups from all the input groups.

`group` default NULL; a column name of `sample_table`. If NULL, search group variable stored in the object.

`by_group` default NULL; one column name in `sample_table`; used to perform paired comparisons within each group. Only available when `paired = TRUE`.

`p_adjust_method` default "fdr"; p.adjust method; available when `paired = TRUE`; see method parameter of p.adjust function for available options.

`permutations` default 999; same with the permutations parameter in anosim function of vegan package.

... parameters passed to anosim function of vegan package.

Returns: `res_anosim` stored in object with `data.frame` class.

Examples:

```
t1$cal_anosim()
```

Method `cal_betadisper()`: Multivariate homogeneity test of groups dispersions (PERMDISP) based on `betadisper` function in `vegan` package.

Usage:

```
trans_beta$cal_betadisper(...)
```

Arguments:

... parameters passed to `betadisper` function.

Returns: `res_betadisper` stored in object.

Examples:

```
t1$cal_betadisper()
```

Method `cal_group_distance()`: Convert symmetric distance matrix to distance table of paired samples that are within groups or between groups.

Usage:

```
trans_beta$cal_group_distance(
  within_group = TRUE,
  by_group = NULL,
  ordered_group = NULL,
  sep = " vs "
)
```

Arguments:

`within_group` default TRUE; whether obtain distance table of paired samples within groups; if FALSE, obtain distances of paired samples between any two groups.

`by_group` default NULL; one colname name of `sample_table` in `microtable` object. If provided, transform distances by the provided `by_group` parameter. This is especially useful for ordering and filtering values further. When `within_group = TRUE`, the result of `by_group` parameter is the format of paired groups. When `within_group = FALSE`, the result of `by_group` parameter is the format same with the group information in `sample_table`.

`ordered_group` default NULL; a vector representing the ordered elements of group parameter; only useful when `within_group = FALSE`.

`sep` default TRUE; a character string to separate the group names after merging them into a new name.

Returns: `res_group_distance` stored in object.

Examples:

```
\donttest{
t1$cal_group_distance(within_group = TRUE)
}
```

Method `cal_group_distance_diff()`: Differential test of converted distances across groups.

Usage:

```
trans_beta$cal_group_distance_diff(
  group = NULL,
  by_group = NULL,
  by_ID = NULL,
  ...
)
```

Arguments:

`group` default NULL; a column name of `object$res_group_distance` used for the statistics; If NULL, use the group inside the object.

`by_group` default NULL; a column of `object$res_group_distance` used to perform the differential test among elements in `group` parameter for each element in `by_group` parameter. So `by_group` has a larger scale than `group` parameter. This `by_group` is very different from the `by_group` parameter in the `cal_group_distance` function.

`by_ID` default NULL; a column of `object$res_group_distance` used to perform paired t test or paired wilcox test for the paired data, such as the data of plant compartments for different plant species (ID). So `by_ID` should be the smallest unit of sample collection without any repetition in it.

... parameters passed to `cal_diff` function of `trans_alpha` class.

Returns: `res_group_distance_diff` stored in object.

Examples:

```
\donttest{
t1$cal_group_distance_diff()
}
```

Method `plot_group_distance()`: Plot the distances of paired groups within or between groups.

Usage:

```
trans_beta$plot_group_distance(plot_group_order = NULL, ...)
```

Arguments:

`plot_group_order` default NULL; a vector used to order the groups in the plot.

`...` parameters (except measure) passed to `plot_alpha` function of `trans_alpha` class.

Returns: ggplot.

Examples:

```
\donttest{
t1$plot_group_distance()
}
```

Method `plot_clustering()`: Plot clustering result based on the `ggdendro` package.

Usage:

```
trans_beta$plot_clustering(
  color_values = RColorBrewer::brewer.pal(8, "Dark2"),
  measure = NULL,
  group = NULL,
  replace_name = NULL
)
```

Arguments:

`color_values` default `RColorBrewer::brewer.pal(8, "Dark2")`; color palette for the text.

`measure` default NULL; beta diversity index; If NULL, using the measure when creating object

`group` default NULL; if provided, use this group to assign color.

`replace_name` default NULL; if provided, use this as label.

Returns: ggplot.

Examples:

```
t1$plot_clustering(group = "Group", replace_name = c("Saline", "Type"))
```

Method `clone()`: The objects of this class are cloneable with this method.

Usage:

```
trans_beta$clone(deep = FALSE)
```

Arguments:

`deep` Whether to make a deep clone.

Examples

```
## -----
## Method `trans_beta$new`
## -----

data(dataset)
t1 <- trans_beta$new(dataset = dataset, measure = "bray", group = "Group")

## -----
```

```

## Method `trans_beta$cal_ordination`
## -----

t1$cal_ordination(method = "PCoA")

## -----
## Method `trans_beta$plot_ordination`
## -----

t1$plot_ordination(plot_type = "point")
t1$plot_ordination(plot_color = "Group", plot_shape = "Group", plot_type = "point")
t1$plot_ordination(plot_color = "Group", plot_type = c("point", "ellipse"))
t1$plot_ordination(plot_color = "Group", plot_type = c("point", "centroid"),
  centroid_segment_linetype = 1)

## -----
## Method `trans_beta$cal_manova`
## -----

t1$cal_manova(manova_all = TRUE)

## -----
## Method `trans_beta$cal_anosim`
## -----

t1$cal_anosim()

## -----
## Method `trans_beta$cal_betadisper`
## -----

t1$cal_betadisper()

## -----
## Method `trans_beta$cal_group_distance`
## -----

t1$cal_group_distance(within_group = TRUE)

## -----
## Method `trans_beta$cal_group_distance_diff`
## -----

t1$cal_group_distance_diff()

## -----
## Method `trans_beta$plot_group_distance`
## -----

```

```
t1$plot_group_distance()

## -----
## Method `trans_beta$plot_clustering`
## -----

t1$plot_clustering(group = "Group", replace_name = c("Saline", "Type"))
```

trans_classifier	Create trans_classifier object for machine-learning-based model prediction.
------------------	---

Description

This class is a wrapper for methods of machine-learning-based classification or regression models, including data pre-processing, feature selection, data split, model training, prediction, confusion-Matrix and ROC (Receiver Operator Characteristic) or PR (Precision-Recall) curve.

Author(s): Felipe Mansoldo and Chi Liu

Methods

Public methods:

- `trans_classifier$new()`
- `trans_classifier$cal_split()`
- `trans_classifier$cal_preProcess()`
- `trans_classifier$cal_feature_sel()`
- `trans_classifier$set_trainControl()`
- `trans_classifier$cal_train()`
- `trans_classifier$cal_feature_imp()`
- `trans_classifier$plot_feature_imp()`
- `trans_classifier$cal_predict()`
- `trans_classifier$plot_confusionMatrix()`
- `trans_classifier$cal_ROC()`
- `trans_classifier$plot_ROC()`
- `trans_classifier$cal_caretList()`
- `trans_classifier$cal_caretList_resamples()`
- `trans_classifier$plot_caretList_resamples()`
- `trans_classifier$clone()`

Method `new()`: Create a trans_classifier object.

Usage:

```
trans_classifier$new(
  dataset,
  x.predictors = "Genus",
  y.response = NULL,
  n.cores = 1
)
```

Arguments:

dataset an object of `microtable` class.

x.predictors default "Genus"; character string or data.frame; a character string represents selecting the corresponding data from `microtable$taxa_abund`; data.frame denotes other customized input. See the following available options:

'Genus' use Genus level table in `microtable$taxa_abund`, or other specific taxonomic rank, e.g., 'Phylum'. If an input level (e.g., ASV) is not found in the names of `taxa_abund` list, the function will use `otu_table` to calculate relative abundance of features.

'all' use all the levels stored in `microtable$taxa_abund`.

other input must be a data.frame object. It should have the same format with the tables in `microtable$taxa_abund`, i.e. rows are features; columns are samples with same names in `sample_table`.

y.response default NULL; the response variable in `sample_table` of input `microtable` object.

n.cores default 1; the CPU thread used.

Returns: `data_feature` and `data_response` stored in the object.

Examples:

```
\donttest{
data(dataset)
t1 <- trans_classifier$new(
  dataset = dataset,
  x.predictors = "Genus",
  y.response = "Group")
}
```

Method `cal_split()`: Split data for training and testing.

Usage:

```
trans_classifier$cal_split(prop.train = 3/4)
```

Arguments:

prop.train default 3/4; the ratio of the data used for the training.

Returns: `data_train` and `data_test` in the object.

Examples:

```
\dontrun{
t1$cal_split(prop.train = 3/4)
}
```

Method `cal_preProcess()`: Pre-process (centering, scaling etc.) of features based on the `caret::preProcess` function. See <https://topepo.github.io/caret/pre-processing.html> for more details.

Usage:

```
trans_classifier$cal_preProcess(...)
```

Arguments:

... parameters pass to preProcess function of caret package.

Returns: data_preProcess, data_train and data_test in the object. data_preProcess is the return data generated by the preProcess function of caret package based on the training data. data_train and data_test are preprocessed training and testing data based on the data_preProcess.

Examples:

```
\dontrun{
# "nzv" removes near zero variance predictors
t1$cal_preProcess(method = c("center", "scale", "nzv"))
}
```

Method cal_feature_sel(): Perform feature selection. See <https://topepo.github.io/caret/feature-selection-overview.html> for more details.

Usage:

```
trans_classifier$cal_feature_sel(
  boruta.maxRuns = 300,
  boruta.pValue = 0.01,
  boruta.repetitions = 4,
  ...
)
```

Arguments:

boruta.maxRuns default 300; maximal number of importance source runs; passed to the maxRuns parameter in Boruta function of Boruta package.

boruta.pValue default 0.01; p value passed to the pValue parameter in Boruta function of Boruta package.

boruta.repetitions default 4; repetition runs for the feature selection.

... parameters pass to Boruta function of Boruta package.

Returns: optimized data_train and data_test in the object.

Examples:

```
\dontrun{
t1$cal_feature_sel(boruta.maxRuns = 300, boruta.pValue = 0.01)
}
```

Method set_trainControl(): Control parameters for the following training. Please see trainControl function of caret package for details.

Usage:

```
trans_classifier$set_trainControl(
  method = "repeatedcv",
  classProbs = TRUE,
  savePredictions = TRUE,
  ...
)
```

Arguments:

method default 'repeatedcv'; 'repeatedcv': Repeated k-Fold cross validation; see method parameter in trainControl function of caret package for available options.
 classProbs default TRUE; should class probabilities be computed for classification models?; see classProbs parameter in caret::trainControl function.
 savePredictions default TRUE; see savePredictions parameter in caret::trainControl function.
 ... parameters pass to trainControl function of caret package.

Returns: trainControl in the object.

Examples:

```
\dontrun{
t1$set_trainControl(method = 'repeatedcv')
}
```

Method cal_train(): Run the model training. Please see <https://topepo.github.io/caret/available-models.html> for available models.

Usage:

```
trans_classifier$cal_train(method = "rf", max.mtry = 2, ntree = 500, ...)
```

Arguments:

method default "rf"; "rf": random forest; see method in train function of caret package for other options. For method = "rf", the tuneGrid is set: expand.grid(mtry = seq(from = 1, to = max.mtry))
 max.mtry default 2; for method = "rf"; maximum mtry used in the tuneGrid to do hyperparameter tuning to optimize the model.
 ntree default 500; for method = "rf"; Number of trees to grow. The default 500 is same with the ntree parameter in randomForest function in randomForest package. When it is a vector with more than one element, the function will try to optimize the model to select a best one, such as c(100, 500, 1000).
 ... parameters pass to caret::train function.

Returns: res_train in the object.

Examples:

```
\dontrun{
# random forest
t1$cal_train(method = "rf")
# Support Vector Machines with Radial Basis Function Kernel
t1$cal_train(method = "svmRadial", tuneLength = 15)
}
```

Method cal_feature_imp(): Get feature importance from the training model.

Usage:

```
trans_classifier$cal_feature_imp(rf_feature_sig = FALSE, ...)
```

Arguments:

rf_feature_sig default FALSE; whether calculate feature significance in 'rf' model using rfPermute package; only available for method = "rf" in cal_train function.

... parameters pass to varImp function of caret package. If rf_feature_sig is TRUE and train_method is "rf", the parameters will be passed to rfPermute function of rfPermute package.

Returns: res_feature_imp in the object. One row for each predictor variable. The column(s) are different importance measures. For the method 'rf', it is MeanDecreaseGini (classification) or IncNodePurity (regression) when rf_feature_sig = FALSE.

Examples:

```
\dontrun{
t1$cal_feature_imp()
}
```

Method plot_feature_imp(): Bar plot for feature importance.

Usage:

```
trans_classifier$plot_feature_imp(
  rf_sig_show = NULL,
  show_sig_group = FALSE,
  ...
)
```

Arguments:

rf_sig_show default NULL; "MeanDecreaseAccuracy" (Default) or "MeanDecreaseGini" for random forest classification; "%IncMSE" (Default) or "IncNodePurity" for random forest regression; Only available when rf_feature_sig = TRUE in function cal_feature_imp, which generate "MeanDecreaseGini" (and "MeanDecreaseAccuracy") or "%IncMSE" (and "IncNodePurity") in the column names of res_feature_imp; Function can also generate "Significance" according to the p value.

show_sig_group default FALSE; whether show the features with different significant groups; Only available when "Significance" is found in the data.

... parameters pass to plot_diff_bar function of trans_diff package.

Returns: ggplot2 object.

Examples:

```
\dontrun{
t1$plot_feature_imp(use_number = 1:20, coord_flip = FALSE)
}
```

Method cal_predict(): Run the prediction.

Usage:

```
trans_classifier$cal_predict(positive_class = NULL)
```

Arguments:

positive_class default NULL; see positive parameter in confusionMatrix function of caret package; If positive_class is NULL, use the first group in data as the positive class automatically.

Returns: res_predict, res_confusion_fit and res_confusion_stats stored in the object. The res_predict is the predicted result for data_test. Several evaluation metrics in res_confusion_fit are defined as follows:

$$Accuracy = \frac{TP + TN}{TP + TN + FP + FN}$$

$$Sensitivity = Recall = TPR = \frac{TP}{TP + FN}$$

$$Specificity = TNR = 1 - FPR = \frac{TN}{TN + FP}$$

$$Precision = \frac{TP}{TP + FP}$$

$$Prevalence = \frac{TP + FN}{TP + TN + FP + FN}$$

$$F1 - Score = \frac{2 * Precision * Recall}{Precision + Recall}$$

$$Kappa = \frac{Accuracy - Pe}{1 - Pe}$$

where TP is true positive; TN is true negative; FP is false positive; and FN is false negative; FPR is False Positive Rate; TPR is True Positive Rate; TNR is True Negative Rate; Pe is the hypothetical probability of chance agreement on the classes for reference and prediction in the confusion matrix. Accuracy represents the ratio of correct predictions. Precision identifies how the model accurately predicted the positive classes. Recall (sensitivity) measures the ratio of actual positives that are correctly identified by the model. F1-score is the weighted average score of recall and precision. The value at 1 is the best performance and at 0 is the worst. Prevalence represents how often positive events occurred. Kappa identifies how well the model is predicting.

Examples:

```
\dontrun{
t1$cal_predict()
}
```

Method `plot_confusionMatrix()`: Plot the cross-tabulation of observed and predicted classes with associated statistics based on the results of function `cal_predict`.

Usage:

```
trans_classifier$plot_confusionMatrix(
  plot_confusion = TRUE,
  plot_statistics = TRUE
)
```

Arguments:

`plot_confusion` default TRUE; whether plot the confusion matrix.
`plot_statistics` default TRUE; whether plot the statistics.

Returns: ggplot object.

Examples:

```
\dontrun{
t1$plot_confusionMatrix()
}
```

Method `cal_ROC()`: Get ROC (Receiver Operator Characteristic) curve data and the performance data.

Usage:

```
trans_classifier$cal_ROC(input = "pred")
```

Arguments:

input default "pred"; 'pred' or 'train'; 'pred' represents using prediction results; 'train' represents using training results.

Returns: a list res_ROC stored in the object. It has two tables: res_roc and res_pr. AUC: Area Under the ROC Curve. For the definition of metrics, please refer to the return part of function cal_predict.

Examples:

```
\dontrun{
t1$cal_ROC()
}
```

Method plot_ROC(): Plot ROC curve.

Usage:

```
trans_classifier$plot_ROC(
  plot_type = c("ROC", "PR")[1],
  plot_group = "all",
  color_values = RColorBrewer::brewer.pal(8, "Dark2"),
  add_AUC = TRUE,
  plot_method = FALSE,
  ...
)
```

Arguments:

plot_type default c("ROC", "PR")[1]; 'ROC' represents ROC (Receiver Operator Characteristic) curve; 'PR' represents PR (Precision-Recall) curve.

plot_group default "all"; 'all' represents all the classes in the model; 'add' represents all adding micro-average and macro-average results, see https://scikit-learn.org/stable/auto_examples/model_selection/ other options should be one or more class names, same with the names in Group column of res_ROC\$res_roc from cal_ROC function.

color_values default RColorBrewer::brewer.pal(8, "Dark2"); colors used in the plot.

add_AUC default TRUE; whether add AUC in the legend.

plot_method default FALSE; If TRUE, show the method in the legend though only one method is found.

... parameters pass to geom_path function of ggplot2 package.

Returns: ggplot2 object.

Examples:

```
\dontrun{
t1$plot_ROC(size = 1, alpha = 0.7)
}
```

Method cal_caretList(): Use caretList function of caretEnsemble package to run multiple models. For the available models, please run names(getModelInfo()).

Usage:

```
trans_classifier$cal_caretList(...)
```

Arguments:

... parameters pass to caretList function of caretEnsemble package.

Returns: res_caretList_models in the object.

Examples:

```
\dontrun{
t1$cal_caretList(methodList = c('rf', 'svmRadial'))
}
```

Method cal_caretList_resamples(): Use resamples function of caret package to collect the metric values based on the res_caretList_models data.

Usage:

```
trans_classifier$cal_caretList_resamples(...)
```

Arguments:

... parameters pass to resamples function of caret package.

Returns: res_caretList_resamples list and res_caretList_resamples_resaped table in the object.

Examples:

```
\dontrun{
t1$cal_caretList_resamples()
}
```

Method plot_caretList_resamples(): Visualize the metric values based on the res_caretList_resamples_resaped data.

Usage:

```
trans_classifier$plot_caretList_resamples(
  color_values = RColorBrewer::brewer.pal(8, "Dark2"),
  ...
)
```

Arguments:

color_values default RColorBrewer::brewer.pal(8, "Dark2"); colors palette for the box.

... parameters pass to geom_boxplot function of ggplot2 package.

Returns: ggplot object.

Examples:

```
\dontrun{
t1$plot_caretList_resamples()
}
```

Method clone(): The objects of this class are cloneable with this method.

Usage:

```
trans_classifier$clone(deep = FALSE)
```

Arguments:

deep Whether to make a deep clone.

Examples

```
## -----
## Method `trans_classifier$new`
## -----

data(dataset)
t1 <- trans_classifier$new(
  dataset = dataset,
  x.predictors = "Genus",
  y.response = "Group")

## -----
## Method `trans_classifier$cal_split`
## -----

## Not run:
t1$cal_split(prop.train = 3/4)

## End(Not run)

## -----
## Method `trans_classifier$cal_preProcess`
## -----

## Not run:
# "nzv" removes near zero variance predictors
t1$cal_preProcess(method = c("center", "scale", "nzv"))

## End(Not run)

## -----
## Method `trans_classifier$cal_feature_sel`
## -----

## Not run:
t1$cal_feature_sel(boruta.maxRuns = 300, boruta.pValue = 0.01)

## End(Not run)

## -----
## Method `trans_classifier$set_trainControl`
## -----

## Not run:
t1$set_trainControl(method = 'repeatedcv')

## End(Not run)

## -----
## Method `trans_classifier$cal_train`
```

```

## -----

## Not run:
# random forest
t1$cal_train(method = "rf")
# Support Vector Machines with Radial Basis Function Kernel
t1$cal_train(method = "svmRadial", tuneLength = 15)

## End(Not run)

## -----
## Method `trans_classifier$cal_feature_imp`
## -----

## Not run:
t1$cal_feature_imp()

## End(Not run)

## -----
## Method `trans_classifier$plot_feature_imp`
## -----

## Not run:
t1$plot_feature_imp(use_number = 1:20, coord_flip = FALSE)

## End(Not run)

## -----
## Method `trans_classifier$cal_predict`
## -----

## Not run:
t1$cal_predict()

## End(Not run)

## -----
## Method `trans_classifier$plot_confusionMatrix`
## -----

## Not run:
t1$plot_confusionMatrix()

## End(Not run)

## -----
## Method `trans_classifier$cal_ROC`
## -----

## Not run:
t1$cal_ROC()

```

```
## End(Not run)

## -----
## Method `trans_classifier$plot_ROC`
## -----

## Not run:
t1$plot_ROC(size = 1, alpha = 0.7)

## End(Not run)

## -----
## Method `trans_classifier$cal_caretList`
## -----

## Not run:
t1$cal_caretList(methodList = c('rf', 'svmRadial'))

## End(Not run)

## -----
## Method `trans_classifier$cal_caretList_resamples`
## -----

## Not run:
t1$cal_caretList_resamples()

## End(Not run)

## -----
## Method `trans_classifier$plot_caretList_resamples`
## -----

## Not run:
t1$plot_caretList_resamples()

## End(Not run)
```

trans_diff

Create trans_diff object for the differential analysis on the taxonomic abundance

Description

This class is a wrapper for a series of differential abundance test and indicator analysis methods, including LEfSe based on the Segata et al. (2011) <doi:10.1186/gb-2011-12-6-r60>, random forest <doi:10.1016/j.geoderma.2018.09.035>, metastat based on White et al. (2009) <doi:10.1371/journal.pcbi.1000352>, non-parametric Kruskal-Wallis Rank Sum Test, Dunn's Kruskal-Wallis Multiple Comparisons based on the FSA package, Wilcoxon Rank Sum and Signed Rank Tests, t-test, anova, Scheirer Ray Hare test, R package metagenomeSeq Paulson et al. (2013) <doi:10.1038/nmeth.2658>, R package

ANCOMBC <doi:10.1038/s41467-020-17041-7>, R package ALDEx2 <doi:10.1371/journal.pone.0067019; 10.1186/2049-2618-2-15>, R package MicrobiomeStat <doi:10.1186/s13059-022-02655-5>, beta regression <doi:10.18637/jss.v034.i02>, R package maaslin2, linear mixed-effects model and generalized linear mixed model.

Methods

Public methods:

- `trans_diff$new()`
- `trans_diff$plot_diff_abund()`
- `trans_diff$plot_diff_bar()`
- `trans_diff$plot_diff_cladogram()`
- `trans_diff$clone()`

Method `new()`:

Usage:

```
trans_diff$new(
  dataset = NULL,
  method = c("lefse", "rf", "metastat", "metagenomeSeq", "KW", "KW_dunn", "wilcox",
    "t.test", "anova", "scheirerRayHare", "lm", "ancombc2", "ALDEx2_t", "ALDEx2_kw",
    "DESeq2", "edgeR", "linda", "maaslin2", "betareg", "lme", "glmm", "glmm_beta")[1],
  group = NULL,
  taxa_level = "all",
  filter_thres = 0,
  alpha = 0.05,
  p_adjust_method = "fdr",
  transformation = NULL,
  remove_unknown = TRUE,
  lefse_subgroup = NULL,
  lefse_min_subsam = 10,
  lefse_sub_strict = FALSE,
  lefse_sub_alpha = NULL,
  lefse_norm = 1e+06,
  nresam = 0.6667,
  boots = 30,
  rf_imp_type = 2,
  group_choose_paired = NULL,
  metagenomeSeq_count = 1,
  ALDEx2_sig = c("wi.eBH", "kw.eBH"),
  by_group = NULL,
  by_ID = NULL,
  beta_pseudo = .Machine$double.eps,
  ...
)
```

Arguments:

`dataset` default NULL; `microtable` object.

method default "lfe". Some methods (e.g., "lfe", "KW", "wilcox", "anova", "lm", "betareg", "glmm" and "glmm_beta") invoke the taxa_abund list (generally relative abundance data) of input microtable object for the analysis. Some (e.g., "metastat", "metagenomeSeq", "ALDEx2_t", "DESeq2", "edgeR", "ancombc2" and "linda") use the otu_table of input microtable object for the analysis. Available options include:

'lfe' LEfSe method based on Segata et al. (2011) <doi:10.1186/gb-2011-12-6-r60>

'rf' random forest and non-parametric test method based on An et al. (2019) <doi:10.1016/j.geoderma.2018.09.035>

'metastat' Metastat method for all paired groups based on White et al. (2009) <doi:10.1371/journal.pcbi.1000352>

'metagenomeSeq' zero-inflated log-normal model-based differential test method from metagenomeSeq package <doi:10.1038/nmeth.2658>

'KW' KW: Kruskal-Wallis Rank Sum Test for all groups (≥ 2)

'KW_dunn' Dunn's Kruskal-Wallis Multiple Comparisons when group number > 2 ; see dunnTest function in FSA package

'wilcox' Wilcoxon Rank Sum and Signed Rank Tests for all paired groups

't.test' Student's t-Test for all paired groups

'anova' ANOVA for one-way or multi-factor analysis; see cal_diff function of trans_alpha class

'scheirerRayHare' Scheirer Ray Hare test for nonparametric test used for a two-way factorial experiment; see scheirerRayHare function of rcompanion package

'lm' Linear Model based on the lm function

'ALDEx2_t' runs Welch's t and Wilcoxon tests with ALDEx2 package; see also the test parameter in ALDEx2::aldex function; ALDEx2 uses the centred log-ratio (clr) transformation and estimates per-feature technical variation within each sample using Monte-Carlo instances drawn from the Dirichlet distribution; Reference: <doi:10.1371/journal.pone.0067019> and <doi:10.1186/2049-2618-2-15>; require ALDEx2 package to be installed (<https://bioconductor.org/packages/release/bioc/html/ALDEx2.html>)

'ALDEx2_kw' runs Kruskal-Wallis and generalized linear model (glm) test with ALDEx2 package; see also the test parameter in ALDEx2::aldex function.

'DESeq2' Differential expression analysis based on the Negative Binomial (a.k.a. Gamma-Poisson) distribution based on the DESeq2 package.

'edgeR' The exactTest method of edgeR package is implemented.

'ancombc2' Analysis of Compositions of Microbiomes with Bias Correction (ANCOM-BC) based on the ancombc2 function from ANCOMBC package <doi:10.1038/s41467-020-17041-7><doi:10.1038/s41592-023-02092-7>. If the fix_formula parameter is not provided, the function can automatically assign it by using group parameter. For this method, the group parameter is directly passed to the group parameter of ancombc2 function. Require ANCOMBC package to be installed (<https://bioconductor.org/packages/release/bioc/html/ANCOMBC.html>)

'linda' Linear Model for Differential Abundance Analysis of High-dimensional Compositional Data based on the linda function of MicrobiomeStat package <doi:10.1186/s13059-022-02655-5>. For linda method, please provide either the group parameter or the formula parameter. When the formula parameter is provided, it should start with '~' as it is directly used by the linda function. If the group parameter is used, the prefix '~' is not necessary as the function can automatically add it. The parameter feature.dat.type = 'count' has been fixed. Other parameters can be passed to the linda function.

'maaslin2' finding associations between metadata and potentially high-dimensional microbial multi-omics data based on the Maaslin2 package <doi:10.1371/journal.pcbi.1009442>. Using this option can invoke the trans_env\$cal_cor function with method = "maaslin2".

'betareg' Beta Regression based on the betareg package <doi:10.18637/jss.v034.i02>. Please see the beta_pseudo parameter for the use of pseudo value when there is 0 or 1 in the data

'lme' Linear Mixed Effect Model based on the lmerTest package. In the return table, the significance of fixed factors are tested by function anova. The significance of 'Estimate' in each term of fixed factors comes from the model.

'glmm' Generalized linear mixed model (GLMM) based on the glmmTMB package <doi:10.32614/RJ-2017-066>. The formula and family parameters are needed. Please refer to glmmTMB package to select the family function, e.g. family = glmmTMB::lognormal(link = "log"). The usage of formula is similar with that in 'lme' method. For more available parameters, please see glmmTMB::glmmTMB function and use parameter passing. In the result, Conditional R2 and Marginal R2 represent the variance explained by both fixed and random effects and the variance explained by fixed effects, respectively. For more details on R2 calculation, please refer to the article <doi: 10.1098/rsif.2017.0213>. The significance of fixed factors are tested by Chi-square test from function car::Anova. The significance of 'Estimate' in each term of fixed factors comes from the model.

'glmm_beta' Generalized linear mixed model with a family function of beta distribution, developed for the relative abundance (ranging from 0 to 1) of taxa specifically. This is an extension of the GLMM model in 'glmm' option. The only difference is in glmm_beta the family function is fixed with the beta distribution function, i.e. family = glmmTMB::beta_family(link = "logit"). Please see the beta_pseudo parameter for the use of pseudo value when there is 0 or 1 in the data

group default NULL; sample group used for the comparison; a colname of input microtable\$sample_table; It is necessary when method is not "anova" or method is "anova" but formula is not provided. Once group is provided, the return res_abund will have mean and sd values for group.

taxa_level default "all"; 'all' represents using abundance data of all taxonomic ranks; For testing at a specific rank, provide taxonomic rank name, such as "Genus". If the provided taxonomic name is neither 'all' nor a colname in tax_table of input dataset (e.g., "ASV"), the function will use the features in input microtable\$otu_table automatically. Note that a specific level (e.g., "ASV") should be provided for method: "metastat", "metagenomeSeq", "ALDEx2_t", "DESeq2", "edgeR", "ancombc2", "linda", "maaslin2".

filter_thres default 0; the abundance threshold, such as 0.0005 when the input is relative abundance; only available when method != "metastat". The features with abundances lower than filter_thres will be filtered.

alpha default 0.05; significance threshold to select taxa when method is "lfe" or "rf"; or used to generate significance letters when method is 'anova' or 'KW_dunn' like the alpha parameter in cal_diff of trans_alpha class.

p_adjust_method default "fdr"; p.adjust method; see method parameter of p.adjust function for other available options; "none" means disable p value adjustment; So when p_adjust_method = "none", P.adj is same with P.unadj.

transformation default NULL; feature abundance transformation method in the class trans_norm, such as 'AST' for the arc sine square root transformation. Only available when method is one of "KW", "KW_dunn", "wilcox", "t.test", "anova", "scheirerRayHare", "betareg" and "lme".

remove_unknown default TRUE; whether remove unknown features that donot have clear classification information.

lefse_subgroup default NULL; sample sub group used for sub-comparison in lefse; Segata et al. (2011) <doi:10.1186/gb-2011-12-6-r60>.

lefse_min_subsam default 10; sample numbers required in the subgroup test.

lefse_sub_strict default FALSE; whether remove the features strictly in the sub-checking. FALSE means only removing the features that have different orders of medians across sub-groups with those across groups and the statistics are also significant. TRUE means removing the features that are not significant in one (or more) sub-test or have different orders of medians across sub-groups with those across groups.

lefse_sub_alpha default NULL; The significance threshold in the test for lefse sub-groups. NULL means it is same with alpha.

lefse_norm default 1000000; normalization value used in lefse to scale abundances for each level. A lefse_norm value < 0 (e.g., -1) means no normalization same with the LEfSe python version.

nresam default 0.6667; sample number ratio used in each bootstrap for method = "lefse" or "rf".

boots default 30; bootstrap test number for method = "lefse" or "rf".

rf_imp_type default 2; the type of feature importance in random forest when method = "rf". Same with type parameter in importance function of randomForest package. 1=mean decrease in accuracy (MeanDecreaseAccuracy), 2=mean decrease in node impurity (MeanDecreaseGini).

group_choose_paired default NULL; a vector used for selecting the required groups for paired testing instead of all paired combinations across groups; Available when method is "metastat", "metagenomeSeq", "ALDEx2_t" or "edgeR".

metagenomeSeq_count default 1; Filter features to have at least 'counts' counts.; see the count parameter in MRcoefs function of metagenomeSeq package.

ALDEx2_sig default c("wi.eBH", "kw.eBH"); which column of the final result is used as the significance asterisk assignment; applied to method = "ALDEx2_t" or "ALDEx2_kw"; the first element is provided to "ALDEx2_t"; the second is provided to "ALDEx2_kw"; for "ALDEx2_t", the available choice is "wi.eBH" (Expected Benjamini-Hochberg corrected P value of Wilcoxon test) and "we.eBH" (Expected BH corrected P value of Welch's t test); for "ALDEx2_kw"; for "ALDEx2_t", the available choice is "kw.eBH" (Expected BH corrected P value of Kruskal-Wallis test) and "glm.eBH" (Expected BH corrected P value of glm test).

by_group default NULL; a column of sample_table used to perform the differential test among groups (group parameter) for each group (by_group parameter). So by_group has a higher level than group parameter. Same with the by_group parameter in trans_alpha class. Only available when method is one of c("KW", "KW_dunn", "wilcox", "t.test", "anova", "scheirerRayHare").

by_ID default NULL; a column of sample_table used to perform paired t test or paired wilcox test for the paired data, such as the data of plant compartments for different plant species (ID). So by_ID in sample_table should be the smallest unit of sample collection without any repetition in it. Same with the by_ID parameter in trans_alpha class.

beta_pseudo default .Machine\$double.eps; the pseudo value used when the parameter method is 'betareg' or 'glm_beta'. As the beta distribution function limits $0 < \text{response value} < 1$, a pseudo value will be added for the data that equal to 0. The data that equal to 1 will be replaced by $1/(1 + \text{beta_pseudo})$.

... parameters passed to `cal_diff` function of `trans_alpha` class when method is one of "KW", "KW_dunn", "wilcox", "t.test", "anova", "betareg", "lme", "glmm" or "glmm_beta"; passed to `randomForest::randomForest` function when method = "rf"; passed to `ANCOMBC::ancombc2` function when method is "ancombc2" (except `tax_level`, `global` and `fix_formula` parameters); passed to `ALDEx2::aldex` function when method = "ALDEx2_t" or "ALDEx2_kw"; passed to `DESeq2::DESeq` function when method = "DESeq2"; passed to `MicrobiomeStat::linda` function when method = "linda"; passed to `trans_env$cal_cor` function when method = "maaslin2".

Returns: `res_diff` and `res_abund`.

res_abund includes mean abundance of each taxa (Mean), standard deviation (SD), standard error (SE) and sample number (N) in the group (Group).

res_diff is the detailed differential test result depending on the method choice, may containing: **"Comparison"**: The groups for the comparison, maybe all groups or paired groups. If this column is not found, all groups are used;

"Group": Which group has the maximum median or mean value across the test groups; For non-parametric methods, median value; For t.test, mean value;

"Taxa": which taxa is used in this comparison;

"Method": Test method used in the analysis depending on the method input;

"LDA" or others: LDA: linear discriminant score in LEfSe; MeanDecreaseAccuracy and MeanDecreaseGini: mean decreasing in accuracy or in node impurity (gini index) in random forest;

"P.unadj": original p value;

"P.adj": adjusted p value;

"Estimate" and "Std.Error": When method is "betareg", "lm", "lme" or "glmm", "Estimate" and "Std.Error" represent fitted coefficient and its standard error, respectively;

Others: qvalue: qvalue in metastat analysis.

Examples:

```
\donttest{
data(dataset)
t1 <- trans_diff$new(dataset = dataset, method = "lefse", group = "Group")
t1 <- trans_diff$new(dataset = dataset, method = "rf", group = "Group")
t1 <- trans_diff$new(dataset = dataset, method = "metastat", group = "Group", taxa_level = "Genus")
t1 <- trans_diff$new(dataset = dataset, method = "wilcox", group = "Group")
}
```

Method `plot_diff_abund()`: Plot the abundance of taxa.

The significance can be optionally added in the plot. The taxa displayed are based on the taxa in the 'res_diff' table, selected using parameters. If the user filters out the non-significant taxa from the 'res_diff' table, these taxa will also be filtered from the plot.

Usage:

```
trans_diff$plot_diff_abund(
  use_number = 1:10,
  color_values = RColorBrewer::brewer.pal(8, "Dark2"),
  select_taxa = NULL,
  simplify_names = TRUE,
  keep_prefix = TRUE,
  group_order = NULL,
  order_x_mean = FALSE,
```

```

    coord_flip = TRUE,
    add_sig = TRUE,
    xtext_angle = 45,
    xtext_size = 13,
    ytitle_size = 17,
    ...
  )

```

Arguments:

`use_number` default 1:10; numeric vector; the sequences of taxa (1:n) selected in the plot; If n is larger than the number of total significant taxa, automatically use the total number as n.

`color_values` default `RColorBrewer::brewer.pal(8, "Dark2")`; color pallete for groups.

`select_taxa` default NULL; character vector to provide taxa names. The taxa names should be same with the names shown in the plot, not the 'Taxa' column names in `object$res_diff$Taxa`.

`simplify_names` default TRUE; whether use the simplified taxonomic name.

`keep_prefix` default TRUE; whether retain the taxonomic prefix.

`group_order` default NULL; a vector to order groups, i.e. reorder the legend and colors in plot; If NULL, the function can first check whether the group column of `sample_table` is factor. If yes, use the levels in it. If provided, overlook the levels in the group of `sample_table`.

`order_x_mean` default FALSE; whether order the taxa in x axis by the means of abundances from large to small. If TRUE, all other factors in the data will become invalid.

`coord_flip` default TRUE; whether flip cartesian coordinates so that horizontal becomes vertical, and vertical becomes horizontal.

`add_sig` default TRUE; whether add the significance label to the plot.

`xtext_angle` default 45; number (e.g. 45). Angle of text in x axis.

`xtext_size` default 13; x axis text size. NULL means the default size in `ggplot2`. If `coord_flip` = TRUE, it represents the text size of the y axis.

`ytitle_size` default 17; y axis title size. If `coord_flip` = TRUE, it represents the title size of the x axis (i.e. "Relative abundance").

... parameters passed to `plot_alpha` function of `trans_alpha` class.

Returns: `ggplot`.

Examples:

```

\donttest{
t1 <- trans_diff$new(dataset = dataset, method = "anova", group = "Group", taxa_level = "Genus")
t1$plot_diff_abund(use_number = 1:10)
t1$plot_diff_abund(use_number = 1:10, add_sig = TRUE)
t1 <- trans_diff$new(dataset = dataset, method = "wilcox", group = "Group")
t1$plot_diff_abund(use_number = 1:20)
t1$plot_diff_abund(use_number = 1:20, add_sig = TRUE)
t1 <- trans_diff$new(dataset = dataset, method = "lefse", group = "Group")
t1$plot_diff_abund(use_number = 1:20)
t1$plot_diff_abund(use_number = 1:20, add_sig = TRUE)
}

```

Method `plot_diff_bar()`: Bar plot for indicator index, such as LDA score and P value.

Usage:

```

trans_diff$plot_diff_bar(
  color_values = RColorBrewer::brewer.pal(8, "Dark2"),
  color_group_map = FALSE,
  use_number = 1:10,
  threshold = NULL,
  select_group = NULL,
  keep_full_name = FALSE,
  keep_prefix = TRUE,
  group_order = NULL,
  group_aggre = TRUE,
  group_two_sep = TRUE,
  coord_flip = TRUE,
  add_sig = FALSE,
  add_sig_increase = 0.1,
  add_sig_text_size = 5,
  xtext_angle = 45,
  xtext_size = 10,
  ytext_size = NULL,
  axis_text_y = deprecated(),
  heatmap_cell = "P.unadj",
  heatmap_sig = "Significance",
  heatmap_x = "Factors",
  heatmap_y = "Taxa",
  heatmap_lab_fill = "P value",
  ...
)

```

Arguments:

`color_values` default `RColorBrewer::brewer.pal(8, "Dark2")`; colors palette for different groups.

`color_group_map` default `FALSE`; whether match the colors to groups in order to fix the color in each group when part of groups are not shown in the plot. When `color_group_map = TRUE`, the `group_order` inside the object will be used as full groups set to guide the color extraction.

`use_number` default `1:10`; numeric vector; the taxa numbers used in the plot, i.e. `1:n`.

`threshold` default `NULL`; threshold value of indicators for selecting taxa, such as 3 for LDA score of LEfSe.

`select_group` default `NULL`; this is used to select the paired group when multiple comparisons are generated; The input `select_group` must be one of `object$res_diff$Comparison`.

`keep_full_name` default `FALSE`; whether keep the taxonomic full lineage names.

`keep_prefix` default `TRUE`; whether retain the taxonomic prefix, such as "g__".

`group_order` default `NULL`; a vector to order the legend and colors in plot; If `NULL`, the function can first determine whether the group column of `microtable$sample_table` is factor. If yes, use the levels in it. If provided, this parameter can overwrite the levels in the group of `microtable$sample_table`.

`group_aggre` default `TRUE`; whether aggregate the features for each group.

`group_two_sep` default `TRUE`; whether display the features of two groups on opposite sides of the coordinate axes when there are only two groups in total.

coord_flip default TRUE; whether flip cartesian coordinates so that horizontal becomes vertical, and vertical becomes horizontal.
 add_sig default FALSE; whether add significance label (asterisk) above the bar.
 add_sig_increase default 0.1; the axis position (Value + add_sig_increase * max(Value)) from which to add the significance label; only available when add_sig = TRUE.
 add_sig_text_size default 5; the size of added significance label; only available when add_sig = TRUE.
 xtext_angle default 45; number ranging from 0 to 90; used to make x axis text generate angle to reduce text overlap; only available when coord_flip = FALSE.
 xtext_size default 10; text size of x axis.
 ytext_size default NULL; text size of y axis. NULL means default ggplot2 value.
 axis_text_y deprecated. Please use ytext_size argument instead.
 heatmap_cell default "P.unadj"; the column of data for the cell of heatmap when formula with multiple factors is found in the method.
 heatmap_sig default "Significance"; the column of data for the significance label of heatmap.
 heatmap_x default "Factors"; the column of data for the x axis of heatmap.
 heatmap_y default "Taxa"; the column of data for the y axis of heatmap.
 heatmap_lab_fill default "P value"; legend title of heatmap.
 ... parameters passing to geom_bar for the bar plot or plot_cor function in [trans_env](#) class for the heatmap of multiple factors when formula is found in the method.

Returns: ggplot.

Examples:

```

\donttest{
t1$plot_diff_bar(use_number = 1:20)
}

```

Method plot_diff_cladogram(): Plot the cladogram using taxa with significant difference.

Usage:

```

trans_diff$plot_diff_cladogram(
  color = RColorBrewer::brewer.pal(8, "Dark2"),
  group_order = NULL,
  use_taxa_num = 200,
  filter_taxa = NULL,
  use_feature_num = NULL,
  clade_label_level = 4,
  select_show_labels = NULL,
  only_select_show = FALSE,
  sep = "|",
  branch_size = 0.2,
  alpha = 0.2,
  clade_label_size = 2,
  clade_label_size_add = 5,
  clade_label_size_log = exp(1),
  node_size_scale = 1,
  node_size_offset = 1,

```

```

    annotation_shape = 22,
    annotation_shape_size = 5
)

```

Arguments:

`color` default `RColorBrewer::brewer.pal(8, "Dark2")`; color palette used in the plot.

`group_order` default `NULL`; a vector to order the legend in plot; If `NULL`, the function can first check whether the `group` column of `sample_table` is factor. If yes, use the levels in it. If provided, this parameter can overwrite the levels in the group of `sample_table`. If the number of provided `group_order` is less than the number of groups in `res_diff$Group`, the function will select the groups of `group_order` automatically.

`use_taxa_num` default 200; integer; The taxa number used in the background tree plot; select the taxa according to the mean abundance .

`filter_taxa` default `NULL`; The mean relative abundance used to filter the taxa with low abundance.

`use_feature_num` default `NULL`; integer; The feature number used in the plot; select the features according to the metric (method = "lefse" or "rf") from high to low.

`clade_label_level` default 4; the taxonomic level for marking the label with letters, root is the largest.

`select_show_labels` default `NULL`; character vector; The features to show in the plot with full label names, not the letters.

`only_select_show` default `FALSE`; whether only use the the select features in the parameter `select_show_labels`.

`sep` default "|"; the separate character in the taxonomic information.

`branch_size` default 0.2; numeric, size of branch.

`alpha` default 0.2; shading of the color.

`clade_label_size` default 2; basic size for the clade label; please also see `clade_label_size_add` and `clade_label_size_log`.

`clade_label_size_add` default 5; added basic size for the clade label; see the formula in `clade_label_size_log` parameter.

`clade_label_size_log` default `exp(1)`; the base of log function for added size of the clade label; the size formula: `clade_label_size + log(clade_label_level + clade_label_size_add, base = clade_label_size_log)`; so use `clade_label_size_log`, `clade_label_size_add` and `clade_label_size` can totally control the label size for different taxonomic levels.

`node_size_scale` default 1; scale for the node size.

`node_size_offset` default 1; offset for the node size.

`annotation_shape` default 22; shape used in the annotation legend.

`annotation_shape_size` default 5; size used in the annotation legend.

Returns: ggplot.

Examples:

```

\dontrun{
t1$plot_diff_cladogram(use_taxa_num = 100, use_feature_num = 30, select_show_labels = NULL)
}

```

Method `clone()`: The objects of this class are cloneable with this method.

Usage:

```
trans_diff$clone(deep = FALSE)
```

Arguments:

deep Whether to make a deep clone.

Examples

```
## -----
## Method `trans_diff$new`
## -----

data(dataset)
t1 <- trans_diff$new(dataset = dataset, method = "lefse", group = "Group")
t1 <- trans_diff$new(dataset = dataset, method = "rf", group = "Group")
t1 <- trans_diff$new(dataset = dataset, method = "metastat", group = "Group", taxa_level = "Genus")
t1 <- trans_diff$new(dataset = dataset, method = "wilcox", group = "Group")

## -----
## Method `trans_diff$plot_diff_abund`
## -----

t1 <- trans_diff$new(dataset = dataset, method = "anova", group = "Group", taxa_level = "Genus")
t1$plot_diff_abund(use_number = 1:10)
t1$plot_diff_abund(use_number = 1:10, add_sig = TRUE)
t1 <- trans_diff$new(dataset = dataset, method = "wilcox", group = "Group")
t1$plot_diff_abund(use_number = 1:20)
t1$plot_diff_abund(use_number = 1:20, add_sig = TRUE)
t1 <- trans_diff$new(dataset = dataset, method = "lefse", group = "Group")
t1$plot_diff_abund(use_number = 1:20)
t1$plot_diff_abund(use_number = 1:20, add_sig = TRUE)

## -----
## Method `trans_diff$plot_diff_bar`
## -----

t1$plot_diff_bar(use_number = 1:20)

## -----
## Method `trans_diff$plot_diff_cladogram`
## -----

## Not run:
t1$plot_diff_cladogram(use_taxa_num = 100, use_feature_num = 30, select_show_labels = NULL)

## End(Not run)
```

trans_env

Create trans_env object to analyze the association between environmental factor and microbial community.

Description

This class is a wrapper for a series of operations associated with environmental measurements, including redundancy analysis, mantel test, correlation analysis and linear fitting.

Methods

Public methods:

- `trans_env$new()`
- `trans_env$scal_diff()`
- `trans_env$plot_diff()`
- `trans_env$scal_autocor()`
- `trans_env$scal_ordination()`
- `trans_env$scal_ordination_anova()`
- `trans_env$scal_ordination_envfit()`
- `trans_env$trans_ordination()`
- `trans_env$plot_ordination()`
- `trans_env$scal_mantel()`
- `trans_env$scal_cor()`
- `trans_env$plot_cor()`
- `trans_env$plot_scatterfit()`
- `trans_env$print()`
- `trans_env$clone()`

Method `new()`:

Usage:

```
trans_env$new(
  dataset = NULL,
  env_cols = NULL,
  add_data = NULL,
  character2numeric = FALSE,
  standardize = FALSE,
  complete_na = FALSE
)
```

Arguments:

`dataset` the object of `microtable` Class.

`env_cols` default `NULL`; either numeric vector or character vector to select columns in `microtable$sample_table`, i.e. `dataset$sample_table`. This parameter should be used in the case that all the required environmental data is in `sample_table` of your `microtable` object. Otherwise, please use `add_data` parameter.

add_data default NULL; data.frame format; provide the environmental data in the format data.frame; rownames should be sample names. This parameter should be used when the microtable\$sample_table object does not have environmental data. Under this circumstance, the env_cols parameter can not be used because no data can be selected.

character2numeric default FALSE; whether convert all the character or factor columns to numeric type using the dropallfactors function. If TRUE, character columns will first be attempted to convert to numeric. If that fails, they will be converted to the factor type and then to numeric.

standardize default FALSE; whether scale environmental variables to zero mean and unit variance.

complete_na default FALSE; Whether fill the NA (missing value) in the environmental data; If TRUE, the function can run the interpolation with the mice package.

Returns: data_env stored in the object.

Examples:

```
data(dataset)
data(env_data_16S)
t1 <- trans_env$new(dataset = dataset, add_data = env_data_16S[, 4:11])
```

Method cal_diff(): Differential test of environmental variables across groups.

Usage:

```
trans_env$cal_diff(
  group = NULL,
  by_group = NULL,
  method = c("KW", "KW_dunn", "wilcox", "t.test", "anova", "scheirerRayHare", "lm",
    "lme", "glmm")[1],
  ...
)
```

Arguments:

group default NULL; a colname of sample_table used to compare values across groups.

by_group default NULL; perform differential test among groups (group parameter) within each group (by_group parameter).

method default "KW"; see the following available options:

'KW' KW: Kruskal-Wallis Rank Sum Test for all groups (≥ 2)

'KW_dunn' Dunn's Kruskal-Wallis Multiple Comparisons, see dunnTest function in FSA package

'wilcox' Wilcoxon Rank Sum and Signed Rank Tests for all paired groups

't.test' Student's t-Test for all paired groups

'anova' Duncan's new multiple range test for one-way anova; see duncan.test function of agricolae package. For multi-factor anova, see aov

'scheirerRayHare' Scheirer Ray Hare test for nonparametric test used for a two-way factorial experiment; see scheirerRayHare function of rcompanion package

'lm' Linear model based on the lm function

'lme' lme: Linear Mixed Effect Model based on the lmerTest package. The formula parameter should be provided.

'glmm' Generalized linear mixed model (GLMM) based on the glmmTMB package. The formula and family parameters are needed. Please refer to glmmTMB package to select the family function, e.g. `family = glmmTMB::lognormal(link = "log")`. The usage of formula is similar with that in 'lme' method. For the details of return table, please refer to the help document of trans_diff class.

... parameters passed to cal_diff function of [trans_alpha](#) class.

Returns: res_diff stored in the object. In the data frame, 'Group' column means that the group has the maximum median or mean value across the test groups; For non-parametric methods, median value; For t.test, mean value.

Examples:

```
\donttest{
t1$cal_diff(group = "Group", method = "KW")
t1$cal_diff(group = "Group", method = "anova")
}
```

Method plot_diff(): Plot environmental variables across groups and add the significance label.

Usage:

```
trans_env$plot_diff(...)
```

Arguments:

... parameters passed to plot_alpha in [trans_alpha](#) class. Please see plot_alpha function of [trans_alpha](#) for all the available parameters.

Method cal_autocor(): Calculate the autocorrelations among environmental variables.

Usage:

```
trans_env$cal_autocor(
  group = NULL,
  ggpairs = TRUE,
  color_values = RColorBrewer::brewer.pal(8, "Dark2"),
  alpha = 0.8,
  ...
)
```

Arguments:

group default NULL; a colname of sample_table; used to perform calculations for different groups.

ggpairs default TRUE; whether use GGally::ggpairs function to plot the correlation results.

If ggpairs = FALSE, the function will output a table with all the values instead of a graph.

In this case, the function will call cal_cor to calculate autocorrelation instead of using the ggpairs function in GGally, so please use parameter passing to control more options.

color_values default RColorBrewer::brewer.pal(8, "Dark2"); colors palette.

alpha default 0.8; the alpha value to add transparency in colors; useful when group is not NULL.

... parameters passed to GGally::ggpairs when ggpairs = TRUE or passed to cal_cor of trans_env class when ggpairs = FALSE.

Returns: ggmatrix when ggpairs = TRUE or data.frame object when ggpairs = FALSE.

Examples:

```
\dontrun{
# Spearman correlation
t1$cal_autocor(upper = list(continuous = GGally::wrap("cor", method= "spearman")))
}
```

Method `cal_ordination()`: Constrained ordination analysis.

Usage:

```
trans_env$cal_ordination(
  method = c("RDA", "dbRDA", "CCA")[1],
  feature_sel = FALSE,
  taxa_level = NULL,
  taxa_filter_thres = NULL,
  use_measure = NULL,
  add_matrix = NULL,
  ...
)
```

Arguments:

`method` default `c("RDA", "dbRDA", "CCA")[1]`; the ordination method; "RDA": redundancy analysis, "dbRDA": distance-based RDA, "CCA": correspondence analysis.

`feature_sel` default `FALSE`; whether perform the feature selection based on forward selection method.

`taxa_level` default `NULL`; the taxonomic level used in RDA or CCA. Default `NULL` means using the merged data at "Genus" level. "ASV" or "OTU" can also be provided for the use of `otu_table` in `microtable` object.

`taxa_filter_thres` default `NULL`; relative abundance threshold used to filter taxa when method is "RDA" or "CCA".

`use_measure` default `NULL`; a name of beta diversity matrix; only available when parameter `method = "dbRDA"`; If not provided, use the first beta diversity matrix in the `microtable$beta_diversity` automatically.

`add_matrix` default `NULL`; additional distance matrix provided, when the user does not want to use the beta diversity matrix within the dataset; only available when `method = "dbRDA"`.

... paremeters passed to `rda`, `dbRda` or `cca` function of `vegan` package according to the method parameter.

Returns: `res_ordination` and `res_ordination_R2` stored in the object.

Examples:

```
\donttest{
t1$cal_ordination(method = "dbRDA", use_measure = "bray")
t1$cal_ordination(method = "RDA", taxa_level = "Genus")
t1$cal_ordination(method = "CCA", taxa_level = "Genus")
}
```

Method `cal_ordination_anova()`: Use anova to test the significance of the terms and axis in ordination.

Usage:

```
trans_env$cal_ordination_anova(...)
```

Arguments:

... parameters passed to anova function.

Returns: res_ordination_terms and res_ordination_axis stored in the object.

Examples:

```
\donttest{
t1$cal_ordination_anova()
}
```

Method cal_ordination_envfit(): Fit each environmental vector onto the ordination to obtain the contribution of each variable.

Usage:

```
trans_env$cal_ordination_envfit(...)
```

Arguments:

... the parameters passed to vegan::envfit function.

Returns: res_ordination_envfit stored in the object.

Examples:

```
\donttest{
t1$cal_ordination_envfit()
}
```

Method trans_ordination(): Transform ordination results for the following plot.

Usage:

```
trans_env$trans_ordination(
  show_taxa = 10,
  adjust_arrow_length = FALSE,
  min_perc_env = 0.1,
  max_perc_env = 0.8,
  min_perc_tax = 0.1,
  max_perc_tax = 0.8
)
```

Arguments:

show_taxa default 10; taxa number shown in the plot.

adjust_arrow_length default FALSE; whether adjust the arrow length to be clearer.

min_perc_env default 0.1; used for scaling up the minimum of env arrow; multiply by the maximum distance between samples and origin.

max_perc_env default 0.8; used for scaling up the maximum of env arrow; multiply by the maximum distance between samples and origin.

min_perc_tax default 0.1; used for scaling up the minimum of tax arrow; multiply by the maximum distance between samples and origin.

max_perc_tax default 0.8; used for scaling up the maximum of tax arrow; multiply by the maximum distance between samples and origin.

Returns: res_ordination_trans stored in the object.

Examples:

```
\donttest{
t1$trans_ordination(adjust_arrow_length = TRUE, min_perc_env = 0.1, max_perc_env = 1)
}
```

Method plot_ordination(): plot ordination result.

Usage:

```
trans_env$plot_ordination(
  plot_color = NULL,
  plot_shape = NULL,
  color_values = RColorBrewer::brewer.pal(8, "Dark2"),
  shape_values = c(16, 17, 7, 8, 15, 18, 11, 10, 12, 13, 9, 3, 4, 0, 1, 2, 14),
  env_text_color = "black",
  env_arrow_color = "grey30",
  taxa_text_color = "firebrick1",
  taxa_arrow_color = "firebrick1",
  env_text_size = 3.7,
  taxa_text_size = 3,
  taxa_text_prefix = FALSE,
  taxa_text_italic = TRUE,
  plot_type = "point",
  point_size = 3,
  point_alpha = 0.8,
  centroid_segment_alpha = 0.6,
  centroid_segment_size = 1,
  centroid_segment_linetype = 3,
  ellipse_chull_fill = TRUE,
  ellipse_chull_alpha = 0.1,
  ellipse_level = 0.9,
  ellipse_type = "t",
  add_sample_label = NULL,
  env_nudge_x = NULL,
  env_nudge_y = NULL,
  taxa_nudge_x = NULL,
  taxa_nudge_y = NULL,
  ...
)
```

Arguments:

plot_color default NULL; a colname of sample_table to assign colors to different groups.

plot_shape default NULL; a colname of sample_table to assign shapes to different groups.

color_values default RColorBrewer::brewer.pal(8, "Dark2"); color pallete for different groups.

shape_values default c(16, 17, 7, 8, 15, 18, 11, 10, 12, 13, 9, 3, 4, 0, 1, 2, 14); a vector for point shape types of groups, see ggplot2 tutorial.

env_text_color default "black"; environmental variable text color.

env_arrow_color default "grey30"; environmental variable arrow color.

taxa_text_color default "firebrick1"; taxa text color.

taxa_arrow_color default "firebrick1"; taxa arrow color.
 env_text_size default 3.7; environmental variable text size.
 taxa_text_size default 3; taxa text size.
 taxa_text_prefix default FALSE; whether show the prefix (e.g., g__) of taxonomic information in the text.
 taxa_text_italic default TRUE; "italic"; whether use "italic" style for the taxa text.
 plot_type default "point"; plotting type of samples; one or more elements of "point", "ellipse", "chull", "centroid" and "none"; "none" denotes nothing.
 'point' add point
 'ellipse' add confidence ellipse for points of each group
 'chull' add convex hull for points of each group
 'centroid' add centroid line of each group
 point_size default 3; point size in plot when "point" is in plot_type. point_size can also be a variable name in sample_table, such as "pH".
 point_alpha default .8; point transparency in plot when "point" is in plot_type.
 centroid_segment_alpha default 0.6; segment transparency in plot when "centroid" is in plot_type.
 centroid_segment_size default 1; segment size in plot when "centroid" is in plot_type.
 centroid_segment_linetype default 3; an integer; the line type related with centroid in plot when "centroid" is in plot_type.
 ellipse_chull_fill default TRUE; whether fill colors to the area of ellipse or chull.
 ellipse_chull_alpha default 0.1; color transparency in the ellipse or convex hull depending on whether "ellipse" or "centroid" is in plot_type.
 ellipse_level default .9; confidence level of ellipse when "ellipse" is in plot_type.
 ellipse_type default "t"; ellipse type when "ellipse" is in plot_type; see type parameter in stat_ellipse function of ggplot2 package.
 add_sample_label default NULL; the column name in sample table, if provided, show the point name in plot.
 env_nudge_x default NULL; numeric vector to adjust the env text x axis position; passed to nudge_x parameter of ggrepel::geom_text_repel function; default NULL represents automatic adjustment; the length must be same with the row number of object\$res_ordination_trans\$df_arrows. For example, if there are 5 env variables, env_nudge_x should be something like c(0.1, 0, -0.2, 0, 0). Note that this parameter and env_nudge_y is generally used when the automatic text adjustment is not very well.
 env_nudge_y default NULL; numeric vector to adjust the env text y axis position; passed to nudge_y parameter of ggrepel::geom_text_repel function; default NULL represents automatic adjustment; the length must be same with the row number of object\$res_ordination_trans\$df_arrows. For example, if there are 5 env variables, env_nudge_y should be something like c(0.1, 0, -0.2, 0, 0).
 taxa_nudge_x default NULL; numeric vector to adjust the taxa text x axis position; passed to nudge_x parameter of ggrepel::geom_text_repel function; default NULL represents automatic adjustment; the length must be same with the row number of object\$res_ordination_trans\$df_arrows. For example, if 3 taxa are shown, taxa_nudge_x should be something like c(0.3, -0.2, 0).

taxa_nudge_y default NULL; numeric vector to adjust the taxa text y axis position; passed to nudge_y parameter of ggrepel::geom_text_repel function; default NULL represents automatic adjustment; the length must be same with the row number of object\$res_ordination_trans\$df_arrows_sp
For example, if 3 taxa are shown, taxa_nudge_y should be something like c(-0.2, 0, 0.4).

... parameters passed to geom_point for controlling sample points.

Returns: ggplot object.

Examples:

```
\donttest{
t1$cal_ordination(method = "RDA")
t1$trans_ordination(adjust_arrow_length = TRUE, max_perc_env = 1.5)
t1$plot_ordination(plot_color = "Group")
t1$plot_ordination(plot_color = "Group", plot_shape = "Group", plot_type = c("point", "ellipse"))
t1$plot_ordination(plot_color = "Group", plot_type = c("point", "chull"))
t1$plot_ordination(plot_color = "Group", plot_type = c("point", "centroid"),
  centroid_segment_linetype = 1)
t1$plot_ordination(plot_color = "Group", env_nudge_x = c(0.4, 0, 0, 0, 0, -0.2, 0, 0),
  env_nudge_y = c(0.6, 0, 0.2, 0.5, 0, 0.1, 0, 0.2))
}
```

Method cal_mantel(): Mantel test between beta diversity matrix and environmental data.

Usage:

```
trans_env$cal_mantel(
  partial_mantel = FALSE,
  add_matrix = NULL,
  use_measure = NULL,
  method = "pearson",
  p_adjust_method = "fdr",
  by_group = NULL,
  ...
)
```

Arguments:

partial_mantel default FALSE; whether use partial mantel test; If TRUE, use other all measurements as the zdis in each calculation.

add_matrix default NULL; additional distance matrix provided when the beta diversity matrix in the dataset is not used.

use_measure default NULL; a name of beta diversity matrix. If necessary and not provided, use the first beta diversity matrix.

method default "pearson"; one of "pearson", "spearman" and "kendall"; correlation method; see method parameter in vegan::mantel function.

p_adjust_method default "fdr"; p.adjust method; see method parameter of p.adjust function for available options.

by_group default NULL; one column name or number in sample_table; used to perform mantel test for different groups separately.

... parameters passed to mantel of vegan package.

Returns: res_mantel in object.

Examples:

```
\donttest{
t1$cal_mantel(use_measure = "bray")
t1$cal_mantel(partial_mantel = TRUE, use_measure = "bray")
}
```

Method `cal_cor()`: Calculate the correlations between taxonomic abundance and environmental variables. Actually, it can also be applied to other correlation between any two variables from two tables.

Usage:

```
trans_env$cal_cor(
  use_data = c("Genus", "all", "other")[1],
  method = c("pearson", "spearman", "kendall", "maaslin2")[1],
  partial = FALSE,
  partial_fix = NULL,
  add_abund_table = NULL,
  filter_thres = 0,
  use_taxa_num = NULL,
  other_taxa = NULL,
  p_adjust_method = "fdr",
  p_adjust_type = c("All", "Taxa", "Env")[1],
  by_group = NULL,
  group_use = NULL,
  group_select = NULL,
  taxa_name_full = TRUE,
  tmp_input_maaslin2 = "tmp_input",
  tmp_output_maaslin2 = "tmp_output",
  cor_method = deprecated(),
  ...
)
```

Arguments:

`use_data` default "Genus"; "Genus", "all" or "other"; "Genus" or other taxonomic names (e.g., "Phylum", "ASV"): invoke taxonomic abundance table in `taxa_abund` list of the `microtable` object; "all": merge all the taxonomic abundance tables in `taxa_abund` list into one; "other": provide additional taxa names by assigning `other_taxa` parameter.

`method` default "pearson"; "pearson", "spearman", "kendall" or "maaslin2"; correlation method. "pearson", "spearman" or "kendall" all refer to the correlation analysis based on the `cor.test` function in R. "maaslin2" is the method in `Maaslin2` package for finding associations between metadata and potentially high-dimensional microbial multi-omics data.

`partial` default FALSE; whether perform partial correlation based on the `ppcor` package. Available when method is "pearson", "spearman" or "kendall".

`partial_fix` default NULL; selected environmental variable names used as third group of variables in all the partial correlations. If NULL; all the variables (except the one for correlation) in the environmental data will be used as the third group of variables. Otherwise, the function will control for the provided variables (one or more) in all the partial correlations, and the variables in `partial_fix` will not be employed anymore in the correlation analysis.

`add_abund_table` default NULL; additional data table to be used. Row names must be sample names.

`filter_thres` default 0; the abundance threshold, such as 0.0005 when the input is relative abundance. The features with abundances lower than `filter_thres` will be filtered. This parameter cannot be applied when `add_abund_table` parameter is provided.

`use_taxa_num` default NULL; integer; a number used to select high abundant taxa; only useful when `use_data` parameter is a taxonomic level, e.g., "Genus".

`other_taxa` default NULL; character vector containing a series of feature names; available when `use_data` = "other"; provided names should be standard full names used to select taxa from all the tables in `taxa_abund` list of the `microtable` object; please refer to the example.

`p_adjust_method` default "fdr"; p.adjust method; see method parameter of `p.adjust` function for available options. `p_adjust_method` = "none" can disable the p value adjustment.

`p_adjust_type` default "All"; "All", "Taxa" or "Env"; P value adjustment type. "Env": adjustment for each environmental variable separately; "Taxa": adjustment for each taxon separately; "All": adjustment for all the data together no matter whether `by_group` is provided.

`by_group` default NULL; one column name or number in `sample_table`; calculate correlations for different groups separately.

`group_use` default NULL; numeric or character vector to select one column in `sample_table` for selecting samples; together with `group_select`.

`group_select` default NULL; the group name used; remain samples within the group.

`taxa_name_full` default TRUE; Whether use the complete taxonomic name of taxa.

`tmp_input_maaslin2` default "tmp_input"; the temporary folder used to save the input files for Maaslin2.

`tmp_output_maaslin2` default "tmp_output"; the temporary folder used to save the output files of Maaslin2.

`cor_method` deprecated. Please use method argument instead.

... parameters passed to Maaslin2 function of Maaslin2 package.

Returns: `res_cor` stored in the object.

Examples:

```
\donttest{
t2 <- trans_diff$new(dataset = dataset, method = "rf", group = "Group", rf_taxa_level = "Genus")
t1 <- trans_env$new(dataset = dataset, add_data = env_data_16S[, 4:11])
t1$cal_cor(use_data = "other", p_adjust_method = "fdr", other_taxa = t2$res_diff$Taxa[1:40])
}
```

Method `plot_cor()`: Plot correlation heatmap.

Usage:

```
trans_env$plot_cor(
  color_vector = c("#053061", "white", "#A50026"),
  color_palette = NULL,
  filter_feature = NULL,
  filter_env = NULL,
  keep_full_name = FALSE,
  keep_prefix = TRUE,
  text_y_order = NULL,
```

```

text_x_order = NULL,
xtext_angle = 30,
xtext_size = 10,
xtext_color = "black",
ytext_italic = FALSE,
ytext_size = NULL,
ytext_color = "black",
ytext_position = "right",
sig_label_size = 4,
font_family = NULL,
legend_title = NULL,
cluster_ggplot = "none",
cluster_height_rows = 0.2,
cluster_height_cols = 0.2,
na.value = "grey50",
trans = "identity",
ylab_type_italic = deprecated(),
text_y_position = deprecated(),
...
)

```

Arguments:

color_vector default `c("#053061", "white", "#A50026")`; colors with only three values representing low, middle and high values.

color_palette default `NULL`; a customized palette with more color values to be used instead of the parameter **color_vector**.

filter_feature default `NULL`; character vector; used to filter features that only have labels in the **filter_feature** vector. For example, **filter_feature** = `""` can be used to remove features that only have `""`, no any `"*"`.

filter_env default `NULL`; character vector; used to filter environmental variables that only have labels in the **filter_env** vector. For example, **filter_env** = `""` can be used to remove features that only have `""`, no any `"*"`.

keep_full_name default `FALSE`; whether use the complete taxonomic name.

keep_prefix default `TRUE`; whether retain the taxonomic prefix.

text_y_order default `NULL`; character vector; customized text for y axis; shown in the plot from the top down. The input should be consistent with the feature names in the **res_cor** table.

text_x_order default `NULL`; character vector; customized text for x axis.

xtext_angle default 30; number ranging from 0 to 90; used to adjust x axis text angle.

xtext_size default 10; x axis text size.

xtext_color default "black"; x axis text color.

ytext_italic default `FALSE`; whether use italic for y axis text.

ytext_size default `NULL`; y axis text size. `NULL` means default ggplot2 value.

ytext_color default "black"; y axis text color.

ytext_position default "right"; "left" or "right"; the y axis text position.

sig_label_size default 4; the size of significance label shown in the cell.

font_family default `NULL`; font family used.

`legend_title` default NULL; legend title; default NULL means 'Pearson' for pearson method and 'Spearman' for spearman method.

`cluster_ggplot` default "none"; add clustering dendrogram for ggplot2 based heatmap. Available options: "none", "row", "col" or "both". "none": no any clustering used; "row": add clustering for rows; "col": add clustering for columns; "both": add clustering for both rows and columns.

`cluster_height_rows` default 0.2, the dendrogram plot height for rows; available when `cluster_ggplot` is not "none".

`cluster_height_cols` default 0.2, the dendrogram plot height for columns; available when `cluster_ggplot` is not "none".

`na.value` default "grey50"; the color for the missing values.

`trans` default "identity"; the transformation for continuous scales in the legend; see the `trans` item in `ggplot2::scale_colour_gradientn`.

`ylab_type_italic` deprecated. Please use `ytext_italic` argument instead.

`text_y_position` deprecated. Please use `ytext_position` argument instead.

... parameters passed to `ggplot2::geom_tile`.

Returns: ggplot2 object.

Examples:

```
\donttest{
t1$plot_cor()
}
```

Method `plot_scatterfit()`: Scatter plot with fitted line based on the correlation or regression. The most important thing is to make sure that the input `x` and `y` have corresponding sample orders. If one of `x` and `y` is a matrix, the other will be also transformed to matrix with Euclidean distance. Then, both of them are transformed to be vectors. If `x` or `y` is a vector with a single value, `x` or `y` will be assigned according to the column selection of the `data_env` in the object.

Usage:

```
trans_env$plot_scatterfit(
  x = NULL,
  y = NULL,
  group = NULL,
  group_order = NULL,
  color_values = RColorBrewer::brewer.pal(8, "Dark2"),
  shape_values = NULL,
  type = c("cor", "lm")[1],
  cor_method = "pearson",
  label_sep = ";",
  label.x.npc = "left",
  label.y.npc = "top",
  label.x = NULL,
  label.y = NULL,
  x_axis_title = "",
  y_axis_title = "",
  point_size = 5,
  point_alpha = 0.6,
```

```

    line_size = 0.8,
    line_color = "black",
    line_se = TRUE,
    line_se_color = "grey70",
    line_alpha = 0.5,
    pvalue_trim = 4,
    cor_coef_trim = 3,
    lm_equation = TRUE,
    lm_fir_trim = 2,
    lm_sec_trim = 2,
    lm_squ_trim = 2,
    ...
  )

```

Arguments:

- x default NULL; a single numeric or character value, a vector, or a distance matrix used for the x axis. If x is a single value, it will be used to select the column of data_env in the object. If x is a distance matrix, it will be transformed to be a vector.
- y default NULL; a single numeric or character value, a vector, or a distance matrix used for the y axis. If y is a single value, it will be used to select the column of data_env in the object. If y is a distance matrix, it will be transformed to be a vector.
- group default NULL; a character vector; if length is 1, must be a colname of sample_table in the input dataset; Otherwise, group should be a vector having same length with x/y (for vector) or column number of x/y (for matrix).
- group_order default NULL; a vector used to order groups, i.e. reorder the legend and colors in plot when group is not NULL; If group_order is NULL and group is provided, the function can first check whether the group column of sample_table is factor. If group_order is provided, disable the group orders or factor levels in the group column of sample_table.
- color_values default RColorBrewer::brewer.pal(8, "Dark2"); color pallete for different groups.
- shape_values default NULL; a numeric vector for point shape types of groups when group is not NULL, see ggplot2 tutorial.
- type default c("cor", "lm")[1]; "cor": correlation; "lm" for regression.
- cor_method default "pearson"; one of "pearson", "kendall" and "spearman"; correlation method.
- label_sep default ";"; the separator string between different label parts.
- label.x.npc default "left"; can be numeric or character vector of the same length as the number of groups and/or panels. If too short, they will be recycled.
 - numeric** value should be between 0 and 1. Coordinates to be used for positioning the label, expressed in "normalized parent coordinates"
 - character** allowed values include: i) one of c('right', 'left', 'center', 'centre', 'middle') for x-axis; ii) and one of c('bottom', 'top', 'center', 'centre', 'middle') for y-axis.
- label.y.npc default "top"; same usage with label.x.npc; also see label.y.npc parameter of ggpubr::stat_cor function.
- label.x default NULL; x axis absolute position for adding the statistic label.
- label.y default NULL; x axis absolute position for adding the statistic label.
- x_axis_title default ""; the title of x axis.
- y_axis_title default ""; the title of y axis.

point_size default 5; point size value.
 point_alpha default 0.6; alpha value for the point color transparency.
 line_size default 0.8; line size value.
 line_color default "black"; fitted line color; only available when group = NULL.
 line_se default TRUE; Whether show the confidence interval for the fitting.
 line_se_color default "grey70"; the color to fill the confidence interval when line_se = TRUE.
 line_alpha default 0.5; alpha value for the color transparency of line confidence interval.
 pvalue_trim default 4; trim the decimal places of p value.
 cor_coef_trim default 3; trim the decimal places of correlation coefficient.
 lm_equation default TRUE; whether include the equation in the label when type = "lm".
 lm_fir_trim default 2; trim the decimal places of first coefficient in regression.
 lm_sec_trim default 2; trim the decimal places of second coefficient in regression.
 lm_squ_trim default 2; trim the decimal places of R square in regression.
 ... other arguments passed to geom_text or geom_label.

Returns: ggplot.

Examples:

```

\donttest{
t1$plot_scatterfit(x = 1, y = 2, type = "cor")
t1$plot_scatterfit(x = 1, y = 2, type = "lm", point_alpha = .3)
t1$plot_scatterfit(x = "pH", y = "TOC", type = "lm", group = "Group", line_se = FALSE)
t1$plot_scatterfit(x =
  dataset$beta_diversity$bray[rownames(t1$data_env), rownames(t1$data_env)], y = "pH")
}

```

Method print(): Print the trans_env object.

Usage:

```
trans_env$print()
```

Method clone(): The objects of this class are cloneable with this method.

Usage:

```
trans_env$clone(deep = FALSE)
```

Arguments:

deep Whether to make a deep clone.

Examples

```

## -----
## Method `trans_env$new`
## -----

data(dataset)
data(env_data_16S)
t1 <- trans_env$new(dataset = dataset, add_data = env_data_16S[, 4:11])

```

```

## -----
## Method `trans_env$cal_diff`
## -----

t1$cal_diff(group = "Group", method = "KW")
t1$cal_diff(group = "Group", method = "anova")

## -----
## Method `trans_env$cal_autocor`
## -----

## Not run:
# Spearman correlation
t1$cal_autocor(upper = list(continuous = GGally::wrap("cor", method= "spearman")))

## End(Not run)

## -----
## Method `trans_env$cal_ordination`
## -----

t1$cal_ordination(method = "dbRDA", use_measure = "bray")
t1$cal_ordination(method = "RDA", taxa_level = "Genus")
t1$cal_ordination(method = "CCA", taxa_level = "Genus")

## -----
## Method `trans_env$cal_ordination_anova`
## -----

t1$cal_ordination_anova()

## -----
## Method `trans_env$cal_ordination_envfit`
## -----

t1$cal_ordination_envfit()

## -----
## Method `trans_env$trans_ordination`
## -----

t1$trans_ordination(adjust_arrow_length = TRUE, min_perc_env = 0.1, max_perc_env = 1)

```

```

## -----
## Method `trans_env$plot_ordination`
## -----

t1$cal_ordination(method = "RDA")
t1$trans_ordination(adjust_arrow_length = TRUE, max_perc_env = 1.5)
t1$plot_ordination(plot_color = "Group")
t1$plot_ordination(plot_color = "Group", plot_shape = "Group", plot_type = c("point", "ellipse"))
t1$plot_ordination(plot_color = "Group", plot_type = c("point", "chull"))
t1$plot_ordination(plot_color = "Group", plot_type = c("point", "centroid"),
  centroid_segment_linetype = 1)
t1$plot_ordination(plot_color = "Group", env_nudge_x = c(0.4, 0, 0, 0, 0, -0.2, 0, 0),
  env_nudge_y = c(0.6, 0, 0.2, 0.5, 0, 0.1, 0, 0.2))

## -----
## Method `trans_env$cal_mantel`
## -----

t1$cal_mantel(use_measure = "bray")
t1$cal_mantel(partial_mantel = TRUE, use_measure = "bray")

## -----
## Method `trans_env$cal_cor`
## -----

t2 <- trans_diff$new(dataset = dataset, method = "rf", group = "Group", rf_taxa_level = "Genus")
t1 <- trans_env$new(dataset = dataset, add_data = env_data_16S[, 4:11])
t1$cal_cor(use_data = "other", p_adjust_method = "fdr", other_taxa = t2$res_diff$Taxa[1:40])

## -----
## Method `trans_env$plot_cor`
## -----

t1$plot_cor()

## -----
## Method `trans_env$plot_scatterfit`
## -----

t1$plot_scatterfit(x = 1, y = 2, type = "cor")
t1$plot_scatterfit(x = 1, y = 2, type = "lm", point_alpha = .3)
t1$plot_scatterfit(x = "pH", y = "TOC", type = "lm", group = "Group", line_se = FALSE)
t1$plot_scatterfit(x =
  dataset$beta_diversity$bray[rownames(t1$data_env), rownames(t1$data_env)], y = "pH")

```

trans_func

*Create trans_func object for functional prediction.***Description**

This class is a wrapper for a series of functional prediction analysis on ASVs/OTUs/species and communities, including the prokaryotic function/trait prediction based on Louca et al. (2016) <doi:10.1126/science.aaf4507> and Lim et al. (2020) <10.1038/s41597-020-0516-5>, or fungal function/trait prediction based on Nguyen et al. (2016) <10.1016/j.funeco.2015.06.006> and Polme et al. (2020) <doi:10.1007/s13225-020-00466-2>; functional redundancy calculation and metabolic pathway abundance prediction Abhauer et al. (2015) <10.1093/bioinformatics/btv287>.

Active bindings

func_group_list store and show the function group list

Methods**Public methods:**

- trans_func\$new()
- trans_func\$cal_func()
- trans_func\$cal_func_FR()
- trans_func\$trans_func_FR()
- trans_func\$plot_func_FR()
- trans_func\$cal_func_FR_comm()
- trans_func\$show_prok_func()
- trans_func\$cal_tax4fun2()
- trans_func\$cal_tax4fun2_FRI()
- trans_func\$cal_spe_func()
- trans_func\$cal_spe_func_perc()
- trans_func\$trans_spe_func_perc()
- trans_func\$plot_spe_func_perc()
- trans_func\$clone()

Method new(): Create the trans_func object. This function can identify the data type for Prokaryotes or Fungi automatically.

Usage:

```
trans_func$new(dataset = NULL)
```

Arguments:

dataset the object of `microtable` Class.

Returns: for_what: "prok" or "fungi" or NA, "prok" represent prokaryotes. "fungi" represent fungi. NA stand for unknown according to the Kingdom information. In this case, if the user still want to use the function to identify species traits, please provide "prok" or "fungi" manually, e.g. `t1$for_what <- "prok"`.

Examples:

```
data(dataset)
t1 <- trans_func$new(dataset = dataset)
```

Method `cal_func()`: Predict the functions or traits for each ASV/OTU/species by matching taxonomic assignments to functional database.

Usage:

```
trans_func$cal_func(
  prok_database = c("FAPROTAX", "NJC19")[1],
  fungi_database = c("FUNGuild", "FungalTraits")[1],
  FUNGuild_confidence = c("Highly Probable", "Probable", "Possible")
)
```

Arguments:

`prok_database` default "FAPROTAX"; "FAPROTAX" or "NJC19"; select a prokaryotic database:

'FAPROTAX' FAPROTAX; Reference: Louca et al. (2016). Decoupling function and taxonomy in the global ocean microbiome. *Science*, 353(6305), 1272. <doi:10.1126/science.aaf4507>

'NJC19' NJC19: Lim et al. (2020). Large-scale metabolic interaction network of the mouse and human gut microbiota. *Scientific Data*, 7(1). <10.1038/s41597-020-0516-5>. Note that the matching in this database is performed at the species level, hence utilizing it demands a higher level of precision in regards to the assignments of species in the taxonomic information table.

`fungi_database` default "FUNGuild"; "FUNGuild" or "FungalTraits"; select a fungal database:

'FUNGuild' Nguyen et al. (2016) FUNGuild: An open annotation tool for parsing fungal community datasets by ecological guild. *Fungal Ecology*, 20(1), 241-248, <doi:10.1016/j.funeco.2015.06.006>

'FungalTraits' version: FungalTraits_1.2_ver_16Dec_2020V.1.2; Polme et al. Fungal-Traits: a user-friendly traits database of fungi and fungus-like stramenopiles. *Fungal Diversity* 105, 1-16 (2020). <doi:10.1007/s13225-020-00466-2>

`FUNGuild_confidence` default c("Highly Probable", "Probable", "Possible"). Selected 'confidenceRanking' when `fungi_database = "FUNGuild"`.

Returns: `res_func` stored in the object.

Examples:

```
\donttest{
t1$cal_func(prok_database = "FAPROTAX")
}
```

Method `cal_func_FR()`: Calculating the functional redundancy (FR) of communities for each function/trait. For each sample and each function/trait, there will be a FR value in the result table. The FR is defined:

$$FR_{kj}^{unweighted} = \frac{N_j}{N_k} \cdot AF$$

$$FR_{kj}^{weighted} = \frac{\sum_{i=1}^{N_j} A_i}{\sum_{i=1}^{N_k} A_i} \cdot AF$$

where FR_{kj} denotes the FR for sample k and function j . N_k is the species number in sample k . N_j is the number of species with function j in sample k . A_i is the abundance (counts) of species i in sample k . AF is adjustment factor based on taxonomic information, representing the taxonomic dispersion of ASVs/OTUs/Species. It is 1 when `adj_tax = FALSE`. Please see the parameter `adj_tax` for detailed explanation.

Usage:

```
trans_func$cal_func_FR(
  abundance_weighted = FALSE,
  adj_tax = FALSE,
  adj_tax_by = "Genus",
  perc = FALSE,
  dec = 6
)
```

Arguments:

`abundance_weighted` default FALSE; whether use abundance of ASVs/OTUs/species. FALSE corresponds to $FR_{kj}^{unweighted}$ in the formula. TRUE corresponds to $FR_{kj}^{weighted}$ in the formula.

`adj_tax` default FALSE; Whether the adjustment factor (AF) is used. The default FALSE represents the AF is 1, meaning no adjustment is made based on the taxonomic distribution. The principle behind the calculation of the adjustment factor is that species with a certain function that are more dispersed taxonomically usually correspond to higher redundancy. It is defined:

$$AF = \frac{NU_{jk}}{NU_k}$$

where NU_{jk} denotes the number of unique taxon (at `adj_tax_by` level) for those ASVs/OTUs/species with function j in sample k . NU_k denotes the number of total unique taxon (at `adj_tax_by` level) in sample k . Please use the parameter `adj_tax_by` to select other taxonomic rank (default Genus level). Here is an example: Suppose a sample k contains a total of 10 genera (including unclassified ones in different lineages), and 3 ASVs with function j are distributed among 2 genera. In this case, the AF would be $\frac{2}{10}$, which is 0.2.

`adj_tax_by` default "Genus"; When `adj_tax = TRUE`, at which taxonomic level is the adjustment factor (AF) calculated?

`perc` default FALSE; whether to use percentages in the result. The default value of FALSE means that the result is in the range of 0 to 1. If it is TRUE, the result will be multiplied by 100, meaning the range will be from 0 to 100.

`dec` default 6; remained decimal places in the result table.

Returns: `res_func_FR` stored in the object.

Examples:

```
\donttest{
t1$cal_func_FR(abundance_weighted = TRUE)
}
```

Method `trans_func_FR()`: Transform the `res_func_FR` table to the long table format for the following visualization. Also add the group information if the database has hierarchical groups.

Usage:

```
trans_func$trans_func_FR()
```

Returns: res_func_FR_trans stored in the object.

Examples:

```
\donttest{
t1$trans_func_FR()
}
```

Method plot_func_FR(): Plot the functional redundancy (FR) results generated by cal_func_FR and trans_func_FR.

Usage:

```
trans_func$plot_func_FR(
  add_facet = TRUE,
  order_x = NULL,
  color_gradient_low = "#00008B",
  color_gradient_high = "#9E0142"
)
```

Arguments:

add_facet default TRUE; whether use group names as the facets in the plot, see trans_func\$func_group_list object.

order_x default NULL; character vector; to sort the x axis text; can be also used to select partial samples to show.

color_gradient_low default "#00008B"; the color used as the low end in the color gradient.

color_gradient_high default "#9E0142"; the color used as the high end in the color gradient.

Returns: ggplot2.

Examples:

```
\donttest{
t1$plot_func_FR()
}
```

Method cal_func_FR_comm(): Calculate the functional redundancy (FR) of communities based on the the result of cal_func_FR function. It is the geometric mean of FR for each function/trait in a community. It is defined:

$$FR_k = \sqrt[n]{FR_{k1} \times FR_{k2} \times \cdots \times FR_{kn}}$$

where FR_k denotes the FR at community level for sample k. FR_{kn} represents the FR of function n for sample k.

Usage:

```
trans_func$cal_func_FR_comm()
```

Returns: vector.

Examples:

```
\donttest{
t1$cal_func_FR_comm()
}
```

Method show_prok_func(): Show the annotation information for a function of prokaryotes from FAPROTAX database.

Usage:

```
trans_func$show_prok_func(use_func = NULL)
```

Arguments:

use_func default NULL; the function name in FAPROTAX database.

Returns: none.

Examples:

```
\donttest{
t1$show_prok_func(use_func = "methanotrophy")
}
```

Method cal_tax4fun2(): Predict functional potential of communities with Tax4Fun2 method. The function was adapted from the raw Tax4Fun2 package to make it compatible with the microtable object. Please cite: Tax4Fun2: prediction of habitat-specific functional profiles and functional redundancy based on 16S rRNA gene sequences. Environmental Microbiome 15, 11 (2020). <doi:10.1186/s40793-020-00358-7>

Usage:

```
trans_func$cal_tax4fun2(
  blast_tool_path = NULL,
  path_to_reference_data = "Tax4Fun2_ReferenceData_v2",
  path_to_temp_folder = NULL,
  database_mode = "Ref99NR",
  normalize_by_copy_number = T,
  min_identity_to_reference = 97,
  use_uproc = T,
  num_threads = 1,
  normalize_pathways = F
)
```

Arguments:

blast_tool_path default NULL; the folder path, e.g., ncbi-blast-2.5.0+/bin ; blast tools folder downloaded from "ftp://ftp.ncbi.nlm.nih.gov/blast/executables/blast+" ; e.g., ncbi-blast-2.5.0+-x64-win64.tar.gz for windows system; if blast_tool_path is NULL, search the tools in the environmental path variable.

path_to_reference_data default "Tax4Fun2_ReferenceData_v2"; the path that points to files used in the prediction; The directory must contain the Ref99NR or Ref100NR folder; download Ref99NR.zip from "https://cloudstor.aarnet.edu.au/plus/s/DkoZIyZpMNbrzSw/download" or Ref100NR.zip from "https://cloudstor.aarnet.edu.au/plus/s/jIByczak9ZAFUB4/download".

path_to_temp_folder default NULL; The temporary folder to store the logfile, intermediate file and result files; if NULL, use the default temporary in the computer system.

database_mode default 'Ref99NR'; "Ref99NR" or "Ref100NR"; Ref99NR: 99% clustering reference database; Ref100NR: no clustering.

normalize_by_copy_number default TRUE; whether normalize the result by the 16S rRNA copy number in the genomes.

min_identity_to_reference default 97; the sequences identity threshold used for finding the nearest species.

use_uproc default TRUE; whether use UProC to functionally annotate the genomes in the reference data.

num_threads default 1; the threads used in the blastn.
 normalize_pathways default FALSE; Different to Tax4Fun, when converting from KEGG functions to KEGG pathways, Tax4Fun2 does not equally split KO gene abundances between pathways a functions is affiliated to. The full predicted abundance is affiliated to each pathway. Use TRUE to split the abundances (default is FALSE).

Returns: res_tax4fun2_KO and res_tax4fun2_pathway in object.

Examples:

```
\dontrun{
t1$cal_tax4fun2(blast_tool_path = "ncbi-blast-2.5.0+/bin",
  path_to_reference_data = "Tax4Fun2_ReferenceData_v2")
}
```

Method cal_tax4fun2_FRI(): Calculate (multi-) functional redundancy index (FRI) of prokaryotic community with Tax4Fun2 method. This function is used to calculating aFRI and rFRI use the intermediate files generated by the function cal_tax4fun2(). please also cite: Tax4Fun2: prediction of habitat-specific functional profiles and functional redundancy based on 16S rRNA gene sequences. Environmental Microbiome 15, 11 (2020). <doi:10.1186/s40793-020-00358-7>

Usage:

```
trans_func$cal_tax4fun2_FRI()
```

Returns: res_tax4fun2_aFRI and res_tax4fun2_rFRI in object.

Examples:

```
\dontrun{
t1$cal_tax4fun2_FRI()
}
```

Method cal_spe_func(): This is a deprecated function. Please use cal_func function instead.

Usage:

```
trans_func$cal_spe_func(...)
```

Arguments:

... paremeters pass to cal_func.

Method cal_spe_func_perc(): This is a deprecated function. Please use cal_func_FR function instead.

Usage:

```
trans_func$cal_spe_func_perc(...)
```

Arguments:

... paremeters pass to cal_func_FR.

Method trans_spe_func_perc(): This is a deprecated function. Please use trans_func_FR function instead.

Usage:

```
trans_func$trans_spe_func_perc(...)
```

Arguments:

... paremeters pass to trans_func_FR.

Method plot_spe_func_perc(): This is a deprecated function. Please use plot_func_FR function instead.

Usage:

```
trans_func$plot_spe_func_perc(...)
```

Arguments:

... paremeters pass to plot_func_FR.

Method clone(): The objects of this class are cloneable with this method.

Usage:

```
trans_func$clone(deep = FALSE)
```

Arguments:

deep Whether to make a deep clone.

Examples

```
## -----
## Method `trans_func$new`
## -----

data(dataset)
t1 <- trans_func$new(dataset = dataset)

## -----
## Method `trans_func$cal_func`
## -----

t1$cal_func(prok_database = "FAPROTAX")

## -----
## Method `trans_func$cal_func_FR`
## -----

t1$cal_func_FR(abundance_weighted = TRUE)

## -----
## Method `trans_func$trans_func_FR`
## -----

t1$trans_func_FR()

## -----
## Method `trans_func$plot_func_FR`
```

```

## -----

t1$plot_func_FR()

## -----
## Method `trans_func$cal_func_FR_comm`
## -----

t1$cal_func_FR_comm()

## -----
## Method `trans_func$show_prok_func`
## -----

t1$show_prok_func(use_func = "methanotrophy")

## -----
## Method `trans_func$cal_tax4fun2`
## -----

## Not run:
t1$cal_tax4fun2(blast_tool_path = "ncbi-blast-2.5.0+/bin",
  path_to_reference_data = "Tax4Fun2_ReferenceData_v2")

## End(Not run)

## -----
## Method `trans_func$cal_tax4fun2_FRI`
## -----

## Not run:
t1$cal_tax4fun2_FRI()

## End(Not run)

```

trans_network

Create trans_network object for network analysis.

Description

This class is a wrapper for a series of network analysis methods, including the network construction, topological attributes analysis, eigengene analysis, network subsetting, node and edge properties, network visualization and other operations.

Methods

Public methods:

- `trans_network$new()`
- `trans_network$cal_network()`
- `trans_network$cal_module()`
- `trans_network$save_network()`
- `trans_network$cal_network_attr()`
- `trans_network$get_node_table()`
- `trans_network$get_edge_table()`
- `trans_network$get_adjacency_matrix()`
- `trans_network$plot_network()`
- `trans_network$cal_eigen()`
- `trans_network$plot_taxa_roles()`
- `trans_network$subset_network()`
- `trans_network$cal_powerlaw()`
- `trans_network$cal_sum_links()`
- `trans_network$plot_sum_links()`
- `trans_network$random_network()`
- `trans_network$trans_comm()`
- `trans_network$print()`
- `trans_network$clone()`

Method `new()`: Create the `trans_network` object, store the important intermediate data and calculate correlations if `cor_method` parameter is not NULL.

Usage:

```
trans_network$new(
  dataset = NULL,
  cor_method = NULL,
  use_WGCNA_pearson_spearman = FALSE,
  use_NetCoMi_pearson_spearman = FALSE,
  use_sparcc_method = c("NetCoMi", "SpiecEasi")[1],
  taxa_level = "OTU",
  filter_thres = 0,
  nThreads = 1,
  SparCC_simu_num = 100,
  env_cols = NULL,
  add_data = NULL,
  ...
)
```

Arguments:

`dataset` default NULL; the object of `microtable` class. Default NULL means customized analysis.

cor_method default NULL; NULL or one of "bray", "pearson", "spearman", "sparcc", "bicor", "cclasso" and "ccrepe"; All the methods referred to NetCoMi package are performed based on netConstruct function of NetCoMi package and require NetCoMi to be installed from Github (<https://github.com/stefpeschel/NetCoMi>); For the algorithm details, please see Peschel et al. 2020 Brief. Bioinform <doi: 10.1093/bib/bbaa290>;

NULL NULL denotes non-correlation network, i.e. do not use correlation-based network. If so, the return res_cor_p list will be NULL.

'bray' 1-B, where B is Bray-Curtis dissimilarity; based on `vegan::vegdist` function

'pearson' Pearson correlation; If `use_WGCNA_pearson_spearman` and `use_NetCoMi_pearson_spearman` are both FALSE, use the function `cor.test` in R; `use_WGCNA_pearson_spearman` = TRUE invoke `corAndPvalue` function of WGCNA package; `use_NetCoMi_pearson_spearman` = TRUE invoke `netConstruct` function of NetCoMi package

'spearman' Spearman correlation; other details are same with the 'pearson' option

'sparcc' SparCC algorithm (Friedman & Alm, PLoS Comp Biol, 2012, <doi:10.1371/journal.pcbi.1002687>); use NetCoMi package when `use_sparcc_method` = "NetCoMi"; use SpiecEasi package when `use_sparcc_method` = "SpiecEasi" and require SpiecEasi to be installed from Github (<https://github.com/zdk123/SpiecEasi>)

'bicor' Calculate biweight midcorrelation efficiently for matrices based on `WGCNA::bicor` function; This option can invoke `netConstruct` function of NetCoMi package; Make sure WGCNA and NetCoMi packages are both installed

'cclasso' Correlation inference of Composition data through Lasso method based on `netConstruct` function of NetCoMi package; for details, see `NetCoMi::cclasso` function

'ccrepe' Calculates compositionality-corrected p-values and q-values for compositional data using an arbitrary distance metric based on `NetCoMi::netConstruct` function; also see `NetCoMi::ccrepe` function

`use_WGCNA_pearson_spearman` default FALSE; whether use WGCNA package to calculate correlation when `cor_method` = "pearson" or "spearman".

`use_NetCoMi_pearson_spearman` default FALSE; whether use NetCoMi package to calculate correlation when `cor_method` = "pearson" or "spearman". The important difference between NetCoMi and others is the features of zero handling and data normalization; See <doi: 10.1093/bib/bbaa290>.

`use_sparcc_method` default `c("NetCoMi", "SpiecEasi")[1]`; use NetCoMi package or SpiecEasi package to perform SparCC when `cor_method` = "sparcc".

`taxa_level` default "OTU"; taxonomic rank; 'OTU' denotes using feature abundance table; other available options should be one of the colnames of `tax_table` of input dataset.

`filter_thres` default 0; the relative abundance threshold.

`nThreads` default 1; the CPU thread number; available when `use_WGCNA_pearson_spearman` = TRUE or `use_sparcc_method` = "SpiecEasi".

`SparCC_simu_num` default 100; SparCC simulation number for bootstrap when `use_sparcc_method` = "SpiecEasi".

`env_cols` default NULL; numeric or character vector to select the column names of environmental data in `dataset$sample_table`; the environmental data can be used in the correlation network (as the nodes) or FlashWeave network.

`add_data` default NULL; provide environmental variable table additionally instead of `env_cols` parameter; rownames must be sample names.

... parameters pass to `NetCoMi::netConstruct` for other operations, such as zero handling and/or data normalization when `cor_method` and other parameters refer to `NetCoMi` package.

Returns: `res_cor_p` list with the correlation (association) matrix and p value matrix. Note that when `cor_method` and other parameters refer to `NetCoMi` package, the p value table are all zero as the significant associations have been selected.

Examples:

```
\donttest{
data(dataset)
# for correlation network
t1 <- trans_network$new(dataset = dataset, cor_method = "pearson",
taxa_level = "OTU", filter_thres = 0.0002)
# for non-correlation network
t1 <- trans_network$new(dataset = dataset, cor_method = NULL)
}
```

Method `cal_network()`: Construct network based on the `igraph` package or `SpiecEasi` package or `julia FlashWeave` package or `beemStatic` package.

Usage:

```
trans_network$cal_network(
  network_method = c("COR", "SpiecEasi", "gcode", "FlashWeave", "beemStatic")[1],
  COR_p_thres = 0.01,
  COR_p_adjust = "fdr",
  COR_return_padjust = FALSE,
  COR_weight = TRUE,
  COR_cut = 0.6,
  COR_optimization = FALSE,
  COR_optimization_low_high = c(0.01, 0.8),
  COR_optimization_seq = 0.01,
  SpiecEasi_method = "mb",
  FlashWeave_tempdir = NULL,
  FlashWeave_meta_data = FALSE,
  FlashWeave_other_para = "alpha=0.01,sensitive=true,heterogeneous=true",
  FlashWeave_gml = NULL,
  beemStatic_t_strength = 0.001,
  beemStatic_t_stab = 0.8,
  add_taxa_name = "Phylum",
  delete_unlinked_nodes = TRUE,
  username_rawtaxa_notOTU = FALSE,
  ...
)
```

Arguments:

`network_method` default "COR"; "COR", "SpiecEasi", "gcode", "FlashWeave" or "beemStatic";
`network_method` = NULL means skipping the network construction for the customized use.
 The option details:

'COR' correlation-based network; use the correlation and p value matrices in `res_cor_p` list stored in the object; See Deng et al. (2012) <doi:10.1186/1471-2105-13-113> for other details

'SpiecEasi' SpiecEasi network; relies on algorithms of sparse neighborhood and inverse covariance selection; belong to the category of conditional dependence and graphical models; see <https://github.com/zdk123/SpiecEasi> for installing the R package; see Kurtz et al. (2015) <doi:10.1371/journal.pcbi.1004226> for the algorithm details

'gcodat' hypothesize the logistic normal distribution of microbiome data; use penalized maximum likelihood method to estimate the sparse structure of inverse covariance for latent normal variables to address the high dimensionality of the microbiome data; belong to the category of conditional dependence and graphical models; depend on the R NetCoMi package <https://github.com/stefpeschel/NetCoMi>; see FANG et al. (2017) <doi:10.1089/cmb.2017.0054> for the algorithm details

'FlashWeave' FlashWeave network; Local-to-global learning framework; belong to the category of conditional dependence and graphical models; good performance on heterogeneous datasets to find direct associations among taxa; see <https://github.com/meringlab/FlashWeave.jl> for installing julia language and FlashWeave package; julia must be in the computer system env path, otherwise the program can not find it; see Tackmann et al. (2019) <doi:10.1016/j.cels.2019.08.002> for the algorithm details

'beemStatic' beemStatic network; extend generalized Lotka-Volterra model to cases of cross-sectional datasets to infer interaction among taxa based on expectation-maximization algorithm; see <https://github.com/CSB5/BEEM-static> for installing the R package; see Li et al. (2021) <doi:10.1371/journal.pcbi.1009343> for the algorithm details

COR_p_thres default 0.01; the p value threshold for the correlation-based network.

COR_p_adjust default "fdr"; p value adjustment method, see method parameter of p.adjust function for available options, in which COR_p_adjust = "none" means giving up the p value adjustment.

COR_return_padjust default FALSE; Whether to return the adjusted p-value matrix and store it in the object (named res_cor_p\$p.adjust).

COR_weight default TRUE; whether use correlation coefficient as the weight of edges; FALSE represents weight = 1 for all edges.

COR_cut default 0.6; correlation coefficient threshold for the correlation network.

COR_optimization default FALSE; whether use random matrix theory (RMT) based method to determine the correlation coefficient; see <https://doi.org/10.1186/1471-2105-13-113>

COR_optimization_low_high default c(0.01, 0.8); the low and high value threshold used for the RMT optimization; only useful when COR_optimization = TRUE.

COR_optimization_seq default 0.01; the interval of correlation coefficient used for RMT optimization; only useful when COR_optimization = TRUE.

SpiecEasi_method default "mb"; either 'glasso' or 'mb'; see spiec.easi function in package SpiecEasi and <https://github.com/zdk123/SpiecEasi>.

FlashWeave_tempdir default NULL; The temporary directory used to save the temporary files for running FlashWeave; If not assigned, use the system user temp.

FlashWeave_meta_data default FALSE; whether use env data for the optimization, If TRUE, the function automatically find the env_data in the object and generate a file for meta_data_path parameter of FlashWeave package.

FlashWeave_other_para default "alpha=0.01,sensitive=true,heterogeneous=true"; the parameters passed to julia FlashWeave package; user can change the parameters or add more according to FlashWeave help document; An exception is meta_data_path parameter as it is generated based on the data inside the object, see FlashWeave_meta_data parameter for the description.

FlashWeave_gml default NULL; The path of FlashWeave output gml file for customized usage.

This parameter is provided for some customized needs. For instance, it can be cumbersome to input bacterial and fungal abundances as separate input files for network analysis using the above parameter. Users can run FlashWeave on their own, and then provide the resulting gml file to this parameter, which allows them to continue using other functions.

beemStatic_t_strength default 0.001; for network_method = "beemStatic"; the threshold used to limit the number of interactions (strength); same with the t_strength parameter in showInteraction function of beemStatic package.

beemStatic_t_stab default 0.8; for network_method = "beemStatic"; the threshold used to limit the number of interactions (stability); same with the t_stab parameter in showInteraction function of beemStatic package.

add_taxa_name default "Phylum"; one or more taxonomic rank name; used to add taxonomic rank name to network node properties.

delete_unlinked_nodes default TRUE; whether delete the nodes without any link.

username_rawtaxa_notOTU default FALSE; whether use OTU name as representatives of taxa when taxa_level != "OTU". Default FALSE means using taxonomic information of taxa_level instead of OTU name.

... parameters pass to SpiecEasi::spiec.easi when network_method = "SpiecEasi"; pass to NetCoMi::netConstruct when network_method = "gcoda"; pass to beemStatic::func.EM when network_method = "beemStatic".

Returns: res_network stored in object.

Examples:

```
\dontrun{
# for correlation network
t1 <- trans_network$new(dataset = dataset, cor_method = "pearson",
taxa_level = "OTU", filter_thres = 0.001)
t1$cal_network(COR_p_thres = 0.05, COR_cut = 0.6)
t1 <- trans_network$new(dataset = dataset, cor_method = NULL, filter_thres = 0.003)
t1$cal_network(network_method = "SpiecEasi", SpiecEasi_method = "mb")
t1 <- trans_network$new(dataset = dataset, cor_method = NULL, filter_thres = 0.005)
t1$cal_network(network_method = "beemStatic")
t1 <- trans_network$new(dataset = dataset, cor_method = NULL, filter_thres = 0.001)
t1$cal_network(network_method = "FlashWeave")
}
```

Method cal_module(): Calculate network modules and add module names to the network node properties.

Usage:

```
trans_network$cal_module(
  method = "cluster_fast_greedy",
  module_name_prefix = "M"
)
```

Arguments:

method default "cluster_fast_greedy"; the method used to find the optimal community structure of a graph; the following are available functions (options) from igraph package: "cluster_fast_greedy", "cluster_walktrap", "cluster_edge_betweenness",

```
"cluster_infomap", "cluster_label_prop", "cluster_leading_eigen",
"cluster_louvain", "cluster_spinglass", "cluster_optimal".
```

For the details of these functions, please see the help document, such as `help(cluster_fast_greedy)`;

Note that the default "cluster_fast_greedy" method can not be applied to directed network. If directed network is provided, the function can automatically switch the default method from "cluster_fast_greedy" to "cluster_walktrap".

`module_name_prefix` default "M"; the prefix of module names; module names are made of the `module_name_prefix` and numbers; numbers are assigned according to the sorting result of node numbers in modules with decreasing trend.

Returns: `res_network` with modules, stored in object.

Examples:

```
\donttest{
t1 <- trans_network$new(dataset = dataset, cor_method = "pearson",
taxa_level = "OTU", filter_thres = 0.0002)
t1$cal_network(COR_p_thres = 0.01, COR_cut = 0.6)
t1$cal_module(method = "cluster_fast_greedy")
}
```

Method `save_network()`: Save network as gexf style, which can be opened by Gephi (<https://gephi.org/>).

Usage:

```
trans_network$save_network(filepath = "network.gexf", ...)
```

Arguments:

`filepath` default "network.gexf"; file path to save the network.

... parameters pass to gexf function of rgexf package except for nodes, edges, edgesLabel, edgesWeight, nodesAtt, edgesAtt and meta.

Returns: None

Examples:

```
\dontrun{
t1$save_network(filepath = "network.gexf")
}
```

Method `cal_network_attr()`: Calculate network properties.

Usage:

```
trans_network$cal_network_attr()
```

Returns: `res_network_attr` stored in object.

Examples:

```
\donttest{
t1$cal_network_attr()
}
```

Method `get_node_table()`: Get the node property table. The properties include the node names, modules allocation, degree, betweenness, abundance, taxonomy, within-module connectivity (zi) and among-module connectivity (Pi) <doi:10.1186/1471-2105-13-113; 10.1016/j.geoderma.2022.115866>.

Usage:

```
trans_network$get_node_table(node_roles = TRUE)
```

Arguments:

`node_roles` default TRUE; whether calculate the node roles <doi:10.1038/nature03288; 10.1186/1471-2105-13-113>. The role of node i is characterized by its within-module connectivity (z_i) and among-module connectivity (P_i) as follows

$$z_i = \frac{k_{ib} - \bar{k}_b}{\sigma_{k_b}}$$

$$P_i = 1 - \sum_{c=1}^{N_M} \left(\frac{k_{ic}}{k_i} \right)^2$$

where k_{ib} is the number of links of node i to other nodes in its module b , $\bar{k}_b$ and σ_{k_b} are the average and standard deviation of within-module connectivity, respectively over all the nodes in module b , k_i is the number of links of node i in the whole network, k_{ic} is the number of links from node i to nodes in module c , and N_M is the number of modules in the network.

Returns: `res_node_table` in object; Abundance expressed as a percentage; `betweenness centrality`: betweenness centrality; `betweenness centrality`: closeness centrality; `eigenvector centrality`: eigenvector centrality; `z`: within-module connectivity; `p`: among-module connectivity.

Examples:

```
\donttest{
t1$get_node_table(node_roles = TRUE)
}
```

Method `get_edge_table()`: Get the edge property table, including connected nodes, label and weight.

Usage:

```
trans_network$get_edge_table()
```

Returns: `res_edge_table` in object.

Examples:

```
\donttest{
t1$get_edge_table()
}
```

Method `get_adjacency_matrix()`: Get the adjacency matrix from the network graph.

Usage:

```
trans_network$get_adjacency_matrix(...)
```

Arguments:

... parameters passed to `as_adjacency_matrix` function of `igraph` package.

Returns: `res_adjacency_matrix` in object.

Examples:

```
\donttest{
t1$get_adjacency_matrix(attr = "weight")
}
```

Method `plot_network()`: Plot the network based on a series of methods from other packages, such as `igraph`, `ggraph` and `networkD3`. The `networkD3` package provides dynamic network. It is especially useful for a glimpse of the whole network structure and finding the interested nodes and edges in a large network. In contrast, the `igraph` and `ggraph` methods are suitable for relatively small network.

Usage:

```
trans_network$plot_network(
  method = c("igraph", "ggraph", "networkD3")[1],
  node_label = "name",
  node_color = NULL,
  ggraph_layout = "fr",
  ggraph_node_size = 2,
  ggraph_node_text = TRUE,
  ggraph_text_color = NULL,
  ggraph_text_size = 3,
  networkD3_node_legend = TRUE,
  networkD3_zoom = TRUE,
  ...
)
```

Arguments:

`method` default "igraph"; The available options:

'igraph' call `plot.igraph` function in `igraph` package for a static network; see `plot.igraph` for the parameters

'ggraph' call `ggraph` function in `ggraph` package for a static network

'networkD3' use `forceNetwork` function in `networkD3` package for a dynamic network; see `forceNetwork` function for the parameters

`node_label` default "name"; node label shown in the plot for `method = "ggraph"` or `method = "networkD3"`; Please see the column names of `object$res_node_table`, which is the returned table of function `object$get_node_table`; User can select other column names in `res_node_table`.

`node_color` default NULL; node color assignment for `method = "ggraph"` or `method = "networkD3"`; Select a column name of `object$res_node_table`, such as "module".

`ggraph_layout` default "fr"; for `method = "ggraph"`; see `layout` parameter of `create_layout` function in `ggraph` package.

`ggraph_node_size` default 2; for `method = "ggraph"`; the node size.

`ggraph_node_text` default TRUE; for `method = "ggraph"`; whether show the label text of nodes.

`ggraph_text_color` default NULL; for `method = "ggraph"`; a column name of `object$res_node_table` used to assign label text colors.

`ggraph_text_size` default 3; for `method = "ggraph"`; the node label text size.

`networkD3_node_legend` default TRUE; used for `method = "networkD3"`; logical value to enable node colour legends; Please see the `legend` parameter in `networkD3::forceNetwork` function.

`networkD3_zoom` default TRUE; used for `method = "networkD3"`; logical value to enable (TRUE) or disable (FALSE) zooming; Please see the `zoom` parameter in `networkD3::forceNetwork` function.

... parameters passed to plot.igraph function when method = "igraph" or forceNetwork function when method = "networkD3".

Returns: network plot.

Examples:

```
\donttest{
t1$plot_network(method = "igraph", layout = layout_with_kk)
t1$plot_network(method = "ggraph", node_color = "module")
t1$plot_network(method = "networkD3", node_color = "module")
}
```

Method cal_eigen(): Calculate eigengenes of modules, i.e. the first principal component based on PCA analysis, and the percentage of variance <doi:10.1186/1471-2105-13-113>.

Usage:

```
trans_network$cal_eigen()
```

Returns: res_eigen and res_eigen_expla in object.

Examples:

```
\donttest{
t1$cal_eigen()
}
```

Method plot_taxa_roles(): Plot the roles or metrics of nodes based on the res_node_table data (coming from function get_node_table) stored in the object.

Usage:

```
trans_network$plot_taxa_roles(
  use_type = c(1, 2)[1],
  roles_color_background = FALSE,
  roles_color_values = NULL,
  add_label = FALSE,
  add_label_group = c("Network hubs", "Module hubs", "Connectors"),
  add_label_text = "name",
  label_text_size = 4,
  label_text_color = "grey50",
  label_text_italic = FALSE,
  label_text_parse = FALSE,
  plot_module = FALSE,
  x_lim = c(0, 1),
  use_level = "Phylum",
  show_value = c("z", "p"),
  show_number = 1:10,
  plot_color = "Phylum",
  plot_shape = "taxa_roles",
  plot_size = "Abundance",
  color_values = RColorBrewer::brewer.pal(12, "Paired"),
  shape_values = c(16, 17, 7, 8, 15, 18, 11, 10, 12, 13, 9, 3, 4, 0, 1, 2, 14),
  ...
)
```

Arguments:

`use_type` default 1; 1 or 2; 1 represents taxa roles plot (node roles include Module hubs, Network hubs, Connectors and Peripherals <doi:10.1038/nature03288; 10.1186/1471-2105-13-113>). The 'p' column (Pi, among-module connectivity) in `res_node_table` table is used in x-axis. The 'z' column (Zi, within-module connectivity) is used in y-axis; 2 represents the layered plot with taxa as x axis and the index (e.g., Zi and Pi) as y axis. Please refer to `res_node_table` data stored in the object for the detailed information.

`roles_color_background` default FALSE; for `use_type=1`; TRUE: use background colors for each area; FALSE: use classic point colors.

`roles_color_values` default NULL; for `use_type=1`; color palette for background or points.

`add_label` default FALSE; for `use_type = 1`; whether add labels for the points.

`add_label_group` default c("Network hubs", "Module hubs", "Connectors"); If `add_label = TRUE`, which part in `taxa_roles` column is used to show labels; character vectors.

`add_label_text` default "name"; If `add_label = TRUE`; which column of `object$res_node_table` is used to label the text.

`label_text_size` default 4; The text size of the label.

`label_text_color` default "grey50"; The text color of the label.

`label_text_italic` default FALSE; whether use italic style for the label text.

`label_text_parse` default FALSE; whether parse the label text. See the `parse` parameter in `ggrepel::geom_text_repel` function.

`plot_module` default FALSE; for `use_type=1`; whether plot the modules information.

`x_lim` default c(0, 1); for `use_type=1`; x axis range when `roles_color_background = FALSE`.

`use_level` default "Phylum"; for `use_type=2`; used taxonomic level in x axis.

`show_value` default c("z", "p"); for `use_type=2`; indexes used in y axis. Please see `res_node_table` in the object for other available indexes.

`show_number` default 1:10; for `use_type=2`; showed number in x axis, sorting according to the nodes number.

`plot_color` default "Phylum"; for `use_type=2`; variable for color.

`plot_shape` default "taxa_roles"; for `use_type=2`; variable for shape.

`plot_size` default "Abundance"; for `use_type=2`; used for point size; a fixed number (e.g. 5) is also acceptable.

`color_values` default `RColorBrewer::brewer.pal(12, "Paired")`; for `use_type=2`; color vector.

`shape_values` default c(16, 17, 7, 8, 15, 18, 11, 10, 12, 13, 9, 3, 4, 0, 1, 2, 14); for `use_type=2`; shape vector, see `ggplot2` tutorial for the shape meaning.

... parameters pass to `geom_point` function of `ggplot2` package.

Returns: `ggplot`.

Examples:

```
\donttest{
t1$plot_taxa_roles(roles_color_background = FALSE)
}
```

Method `subset_network()`: Subset of the network.

Usage:

```
trans_network$subset_network(
  node = NULL,
  edge = NULL,
  rm_single = TRUE,
  node_alledges = FALSE,
  return_igraph = TRUE
)
```

Arguments:

`node` default NULL; provide the node names that will be used in the sub-network.

`edge` default NULL; provide the edge label or numbers that need to be remained. For the edge label, it should must be "+" or "-". For the numbers, they should fall within the range of rows in `res_edge_table` of the object.

`rm_single` default TRUE; whether remove the nodes without any edge in the sub-network. So this function can also be used to remove the nodes without any edge when `node` and `edge` are both NULL.

`node_alledges` default FALSE; whether remain the nodes and edges that related to the nodes provided in `node` parameter. When this parameter is set to TRUE, the network will filter based on edges rather than directly on nodes. The logic is that if at least one of the two nodes connected by an edge is included in the nodes provided by the `node` parameter, the edge will be retained. Otherwise, it will be filtered out. When this parameter is set to FALSE, the network will filter directly based on the `node` parameter. Any nodes not included in the `node` parameter will be filtered out.

`return_igraph` default TRUE; whether return the network with igraph format. If FALSE, return `trans_network` object.

Returns: a new network

Examples:

```
\donttest{
t1$subset_network(node = t1$res_node_table %>% base::subset(module == "M1") %>%
  rownames, rm_single = TRUE)
# return a sub network that contains all nodes of module M1
}
```

Method `cal_powerlaw()`: Fit degrees to a power law distribution. First, perform a bootstrapping hypothesis test to determine whether degrees follow a power law distribution. If the distribution follows power law, then fit degrees to power law distribution and return the parameters.

Usage:

```
trans_network$cal_powerlaw(...)
```

Arguments:

... parameters pass to `bootstrap_p` function in `powerLaw` package.

Returns: `res_powerlaw_p` and `res_powerlaw_fit`; see `powerLaw::bootstrap_p` function for the bootstrapping p value details; see `igraph::fit_power_law` function for the power law fit return details.

Examples:

```
\donttest{
t1$cal_powerlaw()
}
```

Method `cal_sum_links()`: This function is used to sum the links number from one taxa to another or in the same taxa, for example, at Phylum level. This is very useful to fast see how many nodes are connected between different taxa or within the taxa.

Usage:

```
trans_network$cal_sum_links(taxa_level = "Phylum")
```

Arguments:

`taxa_level` default "Phylum"; taxonomic rank.

Returns: `res_sum_links_pos` and `res_sum_links_neg` in object.

Examples:

```
\donttest{
t1$cal_sum_links(taxa_level = "Phylum")
}
```

Method `plot_sum_links()`: Plot the summed linkages among taxa.

Usage:

```
trans_network$plot_sum_links(
  plot_pos = TRUE,
  plot_num = NULL,
  color_values = RColorBrewer::brewer.pal(8, "Dark2"),
  method = c("chorddiag", "circlize")[1],
  ...
)
```

Arguments:

`plot_pos` default TRUE; If TRUE, plot the summed positive linkages; If FALSE, plot the summed negative linkages.

`plot_num` default NULL; number of taxa presented in the plot.

`color_values` default `RColorBrewer::brewer.pal(8, "Dark2")`; colors palette for taxa.

`method` default `c("chorddiag", "circlize")[1]`; `chorddiag` package <<https://github.com/mattflor/chorddiag>> or `circlize` package.

... pass to `chorddiag::chorddiag` function when `method = "chorddiag"` or `circlize::chordDiagram` function when `method = "circlize"`. Note that for `circlize::chordDiagram` function, `keep.diagonal`, `symmetric` and `self.link` parameters have been fixed to fit the input data.

Returns: please see the invoked function.

Examples:

```
\dontrun{
test1$plot_sum_links(method = "chorddiag", plot_pos = TRUE, plot_num = 10)
test1$plot_sum_links(method = "circlize", transparency = 0.2,
  annotationTrackHeight = circlize::mm_h(c(5, 5)))
}
```

Method `random_network()`: Generate random networks, compare them with the empirical network and get the p value of topological properties. The generation of random graph is based on the `erdos.renyi.game` function of `igraph` package. The numbers of vertices and edges in the random graph are same with the empirical network stored in the object.

Usage:

```
trans_network$random_network(runs = 100, output_sim = FALSE)
```

Arguments:

runs default 100; simulation number of random network.

output_sim default FALSE; whether output each simulated network result.

Returns: a data.frame with the following components:

Observed Topological properties of empirical network

Mean_sim Mean of properties of simulated networks

SD_sim SD of properties of simulated networks

p_value Significance, i.e. p values

When output_sim = TRUE, the columns from the five to the last are each simulated result.

Examples:

```
\dontrun{
t1$random_network(runs = 100)
}
```

Method trans_comm(): Transform classified features to community-like microtable object for further analysis, such as module-taxa table.

Usage:

```
trans_network$trans_comm(use_col = "module", abundance = TRUE)
```

Arguments:

use_col default "module"; which column to use as the 'community'; must be one of the name of res_node_table from function get_node_table.

abundance default TRUE; whether sum abundance of taxa. TRUE: sum the abundance for a taxon across all samples; FALSE: sum the frequency for a taxon across all samples.

Returns: a new [microtable](#) class.

Examples:

```
\donttest{
t2 <- t1$trans_comm(use_col = "module")
}
```

Method print(): Print the trans_network object.

Usage:

```
trans_network$print()
```

Method clone(): The objects of this class are cloneable with this method.

Usage:

```
trans_network$clone(deep = FALSE)
```

Arguments:

deep Whether to make a deep clone.

Examples

```
## -----
## Method `trans_network$new`
## -----

data(dataset)
# for correlation network
t1 <- trans_network$new(dataset = dataset, cor_method = "pearson",
  taxa_level = "OTU", filter_thres = 0.0002)
# for non-correlation network
t1 <- trans_network$new(dataset = dataset, cor_method = NULL)

## -----
## Method `trans_network$cal_network`
## -----

## Not run:
# for correlation network
t1 <- trans_network$new(dataset = dataset, cor_method = "pearson",
  taxa_level = "OTU", filter_thres = 0.001)
t1$cal_network(COR_p_thres = 0.05, COR_cut = 0.6)
t1 <- trans_network$new(dataset = dataset, cor_method = NULL, filter_thres = 0.003)
t1$cal_network(network_method = "SpiecEasi", SpiecEasi_method = "mb")
t1 <- trans_network$new(dataset = dataset, cor_method = NULL, filter_thres = 0.005)
t1$cal_network(network_method = "beemStatic")
t1 <- trans_network$new(dataset = dataset, cor_method = NULL, filter_thres = 0.001)
t1$cal_network(network_method = "FlashWeave")

## End(Not run)

## -----
## Method `trans_network$cal_module`
## -----

t1 <- trans_network$new(dataset = dataset, cor_method = "pearson",
  taxa_level = "OTU", filter_thres = 0.0002)
t1$cal_network(COR_p_thres = 0.01, COR_cut = 0.6)
t1$cal_module(method = "cluster_fast_greedy")

## -----
## Method `trans_network$save_network`
## -----

## Not run:
t1$save_network(filepath = "network.gexf")

## End(Not run)
```

```

## -----
## Method `trans_network$cal_network_attr`
## -----

t1$cal_network_attr()

## -----
## Method `trans_network$get_node_table`
## -----

t1$get_node_table(node_roles = TRUE)

## -----
## Method `trans_network$get_edge_table`
## -----

t1$get_edge_table()

## -----
## Method `trans_network$get_adjacency_matrix`
## -----

t1$get_adjacency_matrix(attr = "weight")

## -----
## Method `trans_network$plot_network`
## -----

t1$plot_network(method = "igraph", layout = layout_with_kk)
t1$plot_network(method = "ggraph", node_color = "module")
t1$plot_network(method = "networkD3", node_color = "module")

## -----
## Method `trans_network$cal_eigen`
## -----

t1$cal_eigen()

## -----
## Method `trans_network$plot_taxa_roles`
## -----

```

```

t1$plot_taxa_roles(roles_color_background = FALSE)

## -----
## Method `trans_network$subset_network`
## -----

t1$subset_network(node = t1$res_node_table %>% base::subset(module == "M1") %>%
  rownames, rm_single = TRUE)
# return a sub network that contains all nodes of module M1

## -----
## Method `trans_network$scal_powerlaw`
## -----

t1$scal_powerlaw()

## -----
## Method `trans_network$scal_sum_links`
## -----

t1$scal_sum_links(taxa_level = "Phylum")

## -----
## Method `trans_network$plot_sum_links`
## -----

## Not run:
test1$plot_sum_links(method = "chorddiag", plot_pos = TRUE, plot_num = 10)
test1$plot_sum_links(method = "circlize", transparency = 0.2,
  annotationTrackHeight = circlize::mm_h(c(5, 5)))

## End(Not run)

## -----
## Method `trans_network$random_network`
## -----

## Not run:
t1$random_network(runs = 100)

## End(Not run)

## -----
## Method `trans_network$trans_comm`

```

```
## -----

t2 <- t1$trans_comm(use_col = "module")
```

trans_norm

Feature abundance normalization/transformation.

Description

Feature abundance normalization/transformation for a microtable object or data.frame object.

Methods

Public methods:

- `trans_norm$new()`
- `trans_norm$norm()`
- `trans_norm$clone()`

Method new(): Get a transposed abundance table if the input is microtable object. In the table, rows are samples, and columns are features. This can make the further operations same with the traditional ecological methods.

Usage:

```
trans_norm$new(dataset = NULL)
```

Arguments:

dataset the `microtable` object or `data.frame` object. If it is `data.frame` object, please make sure that rows are samples, and columns are features.

Returns: data_table, stored in the object.

Examples:

```
library(microeco)
data(dataset)
t1 <- trans_norm$new(dataset = dataset)
```

Method norm(): Normalization/transformation methods.

Usage:

```
trans_norm$norm(
  method = "rarefy",
  sample.size = NULL,
  rngseed = 123,
  replace = TRUE,
  pseudocount = 1,
  intersect.no = 10,
  ct.min = 1,
  condition = NULL,
```

```

MARGIN = NULL,
logbase = 2,
CSS_p = NULL,
...
)

```

Arguments:

method default "rarefy"; See the following available options.

Methods for normalization:

- "rarefy": classic rarefaction based on R sample function.
- "SRS": scaling with ranked subsampling method based on the SRS package provided by Lukas Beule and Petr Karlovsky (2020) <doi:10.7717/peerj.9593>.
- "clr": Centered log-ratio normalization <ISBN:978-0-412-28060-3> <doi: 10.3389/fmicb.2017.02224>. It is defined:

$$clr_{ki} = \log \frac{x_{ki}}{g(x_i)}$$

where x_{ki} is the abundance of k th feature in sample i , $g(x_i)$ is the geometric mean of abundances for sample i . A pseudocount need to be added to deal with the zero. For more information, please see the 'clr' method in decostand function of vegan package.

- "rclr": Robust centered log-ratio normalization <doi:10.1128/msystems.00016-19>. It is defined:

$$rclr_{ki} = \log \frac{x_{ki}}{g(x_i > 0)}$$

where x_{ki} is the abundance of k th feature in sample i , $g(x_i > 0)$ is the geometric mean of abundances (> 0) for sample i . In rclr, zero values are kept as zeroes, and not taken into account.

- "GMPR": Geometric mean of pairwise ratios <doi: 10.7717/peerj.4600>. For a given sample i , the size factor s_i is defined:

$$s_i = \left(\prod_{j=1}^n \text{Median}_{k|c_{ki}c_{kj} \neq 0} \left\{ \frac{c_{ki}}{c_{kj}} \right\} \right)^{1/n}$$

where k denotes all the features, and n denotes all the samples. For sample i , $GMPR = \frac{x_i}{s_i}$, where x_i is the feature abundances of sample i .

- "CSS": Cumulative sum scaling normalization based on the metagenomeSeq package <doi:10.1038/nmeth.2658>. For a given sample j , the scaling factor s_j^l is defined:

$$s_j^l = \sum_{i|c_{ij} \leq q_j^l} c_{ij}$$

where q_j^l is the l th quantile of sample j , that is, in sample j there are l features with counts smaller than q_j^l . c_{ij} denotes the count (abundance) of feature i in sample j . For $l = 0.95m$ (feature number), q_j^l corresponds to the 95th percentile of the count distribution for sample j . Normalized counts $\tilde{c}_{ij} = (\frac{c_{ij}}{s_j^l})(N)$, where N is an appropriately chosen normalization constant.

- "TSS": Total sum scaling. Abundance is divided by the sequencing depth. For a given sample j , normalized counts is defined:

$$\tilde{c}_{ij} = \frac{c_{ij}}{\sum_{i=1}^{N_j} c_{ij}}$$

where c_{ij} is the counts of feature i in sample j , and N_j is the feature number of sample j .

- "eBay": Empirical Bayes approach to normalization <10.1186/s12859-020-03552-z>. The implemented method is not tree-related. In the output, the sum of each sample is 1.
- "TMM": Trimmed mean of M-values method based on the normLibSizes function of edgeR package <doi: 10.1186/gb-2010-11-3-r25>.
- "DESeq2": Median ratio of gene counts relative to geometric mean per gene based on the DESeq function of DESeq2 package <doi: 10.1186/s13059-014-0550-8>. This option can invoke the trans_diff class and extract the normalized data from the original result. Note that either group or formula should be provided. The scaling factor is defined:

$$s_j = \text{Median}_i \frac{c_{ij}}{(\prod_{j=1}^n c_{ij})^{1/n}}$$

where c_{ij} is the counts of feature i in sample j , and n is the total sample number.

- "Wrench": Group-wise and sample-wise compositional bias factor <doi: 10.1186/s12864-018-5160-5>. Note that condition parameter is necessary to be passed to condition parameter in wrench function of Wrench package. As the input data must be microtable object, so the input condition parameter can be a column name of sample_table. The scaling factor is defined:

$$s_j = \frac{1}{p} \sum_{ij} W_{ij} \frac{X_{ij}}{\bar{X}_i}$$

where X_{ij} represents the relative abundance (proportion) for feature i in sample j , $\bar{X}_i$ is the average proportion of feature i across the dataset, W_{ij} represents a weight specific to each technique, and p is the feature number in sample.

- "RLE": Relative log expression.

Methods based on decostand function of vegan package:

- "total": divide by margin total (default MARGIN = 1, i.e. rows - samples).
- "max": divide by margin maximum (default MARGIN = 2, i.e. columns - features).
- "normalize": make margin sum of squares equal to one (default MARGIN = 1).
- "range": standardize values into range 0...1 (default MARGIN = 2). If all values are constant, they will be transformed to 0.
- "standardize": scale x to zero mean and unit variance (default MARGIN = 2).
- "pa": scale x to presence/absence scale (0/1).
- "log": logarithmic transformation.

Other methods for transformation:

- "AST": Arc sine square root transformation.

sample.size default NULL; library size for rarefaction when method = "rarefy" or "SRS". If not provided, use the minimum number across all samples. For "SRS" method, this parameter is passed to Cmin parameter of SRS function of SRS package.

rngseed default 123; random seed. Available when method = "rarefy" or "SRS".

replace default TRUE; see [sample](#) for the random sampling; Available when method = "rarefy".

pseudocount default 1; add pseudocount for those features with 0 abundance when method = "clr".

intersect.no default 10; the intersecting taxa number between paired sample for method = "GMPR".

ct.min default 1; the minimum number of counts required to calculate ratios for method = "GMPR".

condition default NULL; Only available when method = "Wrench". This parameter is passed to the condition parameter of wrench function in Wrench package. It must be a column name of sample_table or a vector with same length of samples.

MARGIN default NULL; 1 = samples, and 2 = features of abundance table; only available when method comes from decostand function of vegan package. If MARGIN is NULL, use the default value in decostand function.

logbase default 2; The logarithm base.

CSS_p default NULL; the pth quantile passed to the p argument of the cumNorm function when method = "CSS". Default NULL means the default cumNormStatFast function is used to calculate the quantile.

... parameters pass to vegan::decostand, or edgeR::normLibSizes when method = "TMM" or "RLE", or trans_diff class when method = "DESeq2", or wrench function of Wrench package when method = "Wrench".

Returns: new microtable object or data.frame object.

Examples:

```
newdataset <- t1$norm(method = "clr")
newdataset <- t1$norm(method = "log")
```

Method clone(): The objects of this class are cloneable with this method.

Usage:

```
trans_norm$clone(deep = FALSE)
```

Arguments:

deep Whether to make a deep clone.

Examples

```
## -----
## Method `trans_norm$new`
## -----

library(microeco)
data(dataset)
t1 <- trans_norm$new(dataset = dataset)

## -----
## Method `trans_norm$norm`
## -----

newdataset <- t1$norm(method = "clr")
newdataset <- t1$norm(method = "log")
```

trans_nullmodel	Create trans_nullmodel object for null model related analysis.
-----------------	--

Description

This class is a wrapper for a series of null model related approaches, including the mantel correlogram analysis of phylogenetic signal, beta nearest taxon index (betaNTI), beta net relatedness index (betaNRI), NTI, NRI and RCbray (Raup–Crick Bray–Curtis) calculations. See <doi:10.1111/j.1600-0587.2010.06548.x; 10.1890/ES10-00117.1; 10.1038/ismej.2013.93; 10.1038/s41598-017-17736-w> for the algorithms and applications.

Methods

Public methods:

- `trans_nullmodel$new()`
- `trans_nullmodel$cal_mantel_corr()`
- `trans_nullmodel$plot_mantel_corr()`
- `trans_nullmodel$cal_betampd()`
- `trans_nullmodel$cal_betamntd()`
- `trans_nullmodel$cal_ses_betampd()`
- `trans_nullmodel$cal_ses_betamntd()`
- `trans_nullmodel$cal_rcbray()`
- `trans_nullmodel$cal_process()`
- `trans_nullmodel$cal_NRI()`
- `trans_nullmodel$cal_NTI()`
- `trans_nullmodel$cal_Cscore()`
- `trans_nullmodel$cal_NST()`
- `trans_nullmodel$cal_NST_test()`
- `trans_nullmodel$cal_NST_convert()`
- `trans_nullmodel$clone()`

Method new():

Usage:

```
trans_nullmodel$new(
  dataset = NULL,
  filter_thres = 0,
  taxa_number = NULL,
  group = NULL,
  select_group = NULL,
  env_cols = NULL,
  add_data = NULL,
  complete_na = FALSE
)
```

Arguments:

dataset the object of `microtable` Class.
 filter_thres default 0; the relative abundance threshold.
 taxa_number default NULL; how many taxa the user want to keep, if provided, filter_thres parameter will be forcible invalid.
 group default NULL; which column name in sample_table is selected as the group for the following selection.
 select_group default NULL; one or more elements in group, used to select samples.
 env_cols default NULL; number or name vector to select the environmental data in dataset\$sample_table.
 add_data default NULL; provide environmental data table additionally.
 complete_na default FALSE; whether fill the NA in environmental data based on the method in mice package.

Returns: data_comm and data_tree in object.

Examples:

```
data(dataset)
data(env_data_16S)
t1 <- trans_nullmodel$new(dataset, filter_thres = 0.0005, add_data = env_data_16S)
```

Method `cal_mantel_corr()`: Calculate mantel correlogram.

Usage:

```
trans_nullmodel$cal_mantel_corr(
  use_env = NULL,
  break.pts = seq(0, 1, 0.02),
  cutoff = FALSE,
  ...
)
```

Arguments:

use_env default NULL; numeric or character vector to select env_data; if provide multiple variables or NULL, use PCA (principal component analysis) to reduce dimensionality.
 break.pts default seq(0, 1, 0.02); see break.pts parameter in mantel.correlog of vegan package.
 cutoff default FALSE; see cutoff parameter in mantel.correlog.
 ... parameters pass to mantel.correlog function in vegan package.

Returns: res_mantel_corr in object.

Examples:

```
\dontrun{
t1$cal_mantel_corr(use_env = "pH")
}
```

Method `plot_mantel_corr()`: Plot mantel correlogram.

Usage:

```
trans_nullmodel$plot_mantel_corr(point_shape = 22, point_size = 3)
```

Arguments:

point_shape default 22; the number for selecting point shape type; see ggplot2 manual for the number meaning.

point_size default 3; the point size.

Returns: ggplot.

Examples:

```
\dontrun{
t1$plot_mantel_corr()
}
```

Method cal_betampd(): Calculate betaMPD (mean pairwise distance). Same with picante::comdist function, but faster.

Usage:

```
trans_nullmodel$cal_betampd(abundance.weighted = TRUE)
```

Arguments:

abundance.weighted default TRUE; whether use abundance-weighted method.

Returns: res_betampd in object.

Examples:

```
\donttest{
t1$cal_betampd(abundance.weighted = TRUE)
}
```

Method cal_betamntd(): Calculate betaMNTD (mean nearest taxon distance). Same with picante::comdistnt function, but faster.

Usage:

```
trans_nullmodel$cal_betamntd(
  abundance.weighted = TRUE,
  exclude.conspecifics = FALSE,
  use_iCAMP = FALSE,
  use_iCAMP_force = TRUE,
  iCAMP_tempdir = NULL,
  ...
)
```

Arguments:

abundance.weighted default TRUE; whether use abundance-weighted method.

exclude.conspecifics default FALSE; see exclude.conspecifics parameter in comdistnt function of picante package.

use_iCAMP default FALSE; whether use bmntd.big function of iCAMP package to calculate betaMNTD. This method can store the phylogenetic distance matrix on the disk to lower the memory spending and perform the calculation parallelly.

use_iCAMP_force default FALSE; whether use bmntd.big function of iCAMP package automatically when the feature number is large.

iCAMP_tempdir default NULL; the temporary directory used to place the large tree file; If NULL; use the system user tempdir.

... paremeters pass to iCAMP::pdist.big function.

Returns: res_betamntd in object.

Examples:

```
\donttest{
t1$cal_betamntd(abundance.weighted = TRUE)
}
```

Method cal_ses_betampd(): Calculate standardized effect size of betaMPD, i.e. beta net relatedness index (betaNRI).

Usage:

```
trans_nullmodel$cal_ses_betampd(
  runs = 1000,
  null.model = c("taxa.labels", "richness", "frequency", "sample.pool", "phylogeny.pool",
    "independentswap", "trialswap")[1],
  abundance.weighted = TRUE,
  iterations = 1000
)
```

Arguments:

runs default 1000; simulation runs.

null.model default "taxa.labels"; The available options include "taxa.labels", "richness", "frequency", "sample.pool", "phylogeny.pool", "independentswap" and "trialswap"; see null.model parameter of ses.mntd function in picante package for the algorithm details.

abundance.weighted default TRUE; whether use weighted abundance.

iterations default 1000; iteration number for part null models to perform; see iterations parameter of picante::randomizeMatrix function.

Returns: res_ses_betampd in object.

Examples:

```
\dontrun{
# only run 50 times for the example; default 1000
t1$cal_ses_betampd(runs = 50, abundance.weighted = TRUE)
}
```

Method cal_ses_betamntd(): Calculate standardized effect size of betaMNTD, i.e. beta nearest taxon index (betaNTI).

Usage:

```
trans_nullmodel$cal_ses_betamntd(
  runs = 1000,
  null.model = c("taxa.labels", "richness", "frequency", "sample.pool", "phylogeny.pool",
    "independentswap", "trialswap")[1],
  abundance.weighted = TRUE,
  exclude.conspecifics = FALSE,
  use_iCAMP = FALSE,
  use_iCAMP_force = TRUE,
  iCAMP_tempdir = NULL,
  nworker = 2,
  iterations = 1000
)
```

Arguments:

runs default 1000; simulation number of null model.

null.model default "taxa.labels"; The available options include "taxa.labels", "richness", "frequency", "sample.pool", "phylogeny.pool", "independentswap" and "trialswap"; see null.model parameter of ses.mntd function in picante package for the algorithm details.

abundance.weighted default TRUE; whether use abundance-weighted method.

exclude.conspecificity default FALSE; see comdistnt in picante package.

use_iCAMP default FALSE; whether use bmntd.big function of iCAMP package to calculate betaMNTD. This method can store the phylogenetic distance matrix on the disk to lower the memory spending and perform the calculation parallelly.

use_iCAMP_force default FALSE; whether to make use_iCAMP to be TRUE when the feature number is large.

iCAMP_tempdir default NULL; the temporary directory used to place the large tree file; If NULL; use the system user tempdir.

nworker default 2; the CPU thread number.

iterations default 1000; iteration number for part null models to perform; see iterations parameter of picante::randomizeMatrix function.

Returns: res_ses_betamntd in object.

Examples:

```
\dontrun{
# only run 50 times for the example; default 1000
t1$cal_ses_betamntd(runs = 50, abundance.weighted = TRUE, exclude.conspecificity = FALSE)
}
```

Method cal_rcbray(): Calculate Bray–Curtis-based Raup–Crick (RCbray) <doi: 10.1890/ES10-00117.1>.

Usage:

```
trans_nullmodel$cal_rcbray(
  runs = 1000,
  verbose = TRUE,
  null.model = "independentswap"
)
```

Arguments:

runs default 1000; simulation runs.

verbose default TRUE; whether show the calculation process message.

null.model default "independentswap"; see more available options in randomizeMatrix function of picante package.

Returns: res_rcbray in object.

Examples:

```
\dontrun{
# only run 50 times for the example; default 1000
t1$cal_rcbray(runs = 50)
}
```

Method `cal_process()`: Infer the ecological processes according to `ses.betaMNTD` (`betaNTI`)/`ses.betaMPD` (`betaNRI`) and `rcbray`.

Usage:

```
trans_nullmodel$cal_process(use_betamntd = TRUE, group = NULL)
```

Arguments:

`use_betamntd` default TRUE; whether use `ses.betaMNTD` (`betaNTI`); if False, use `ses.betaMPD` (`betaNRI`).

`group` default NULL; a column name in `sample_table` of `microtable` object. If provided, the analysis will be performed for each group instead of the whole.

Returns: `res_process` in object.

Examples:

```
\dontrun{
t1$cal_process(use_betamntd = TRUE)
}
```

Method `cal_NRI()`: Calculates Nearest Relative Index (NRI), equivalent to -1 times the standardized effect size of MPD.

Usage:

```
trans_nullmodel$cal_NRI(
  null.model = "taxa.labels",
  abundance.weighted = FALSE,
  runs = 999,
  ...
)
```

Arguments:

`null.model` default "taxa.labels"; Null model to use; see `null.model` parameter in `ses.mpd` function of `picante` package for available options.

`abundance.weighted` default FALSE; Should mean nearest relative distances for each species be weighted by species abundance?

`runs` default 999; Number of randomizations.

... parameters pass to `ses.mpd` function in `picante` package.

Returns: `res_NRI` in object, equivalent to -1 times `ses.mpd`.

Examples:

```
\donttest{
# only run 50 times for the example; default 999
t1$cal_NRI(null.model = "taxa.labels", abundance.weighted = FALSE, runs = 50)
}
```

Method `cal_NTI()`: Calculates Nearest Taxon Index (NTI), equivalent to -1 times the standardized effect size of MNTD.

Usage:

```
trans_nullmodel$cal_NTI(
  null.model = "taxa.labels",
  abundance.weighted = FALSE,
```

```

    runs = 999,
    ...
)

```

Arguments:

`null.model` default "taxa.labels"; Null model to use; see `null.model` parameter in `ses.mntd` function of `picante` package for available options.

`abundance.weighted` default FALSE; Should mean nearest taxon distances for each species be weighted by species abundance?

`runs` default 999; Number of randomizations.

... parameters pass to `ses.mntd` function in `picante` package.

Returns: `res_NTI` in object, equivalent to -1 times `ses.mntd`.

Examples:

```

\donttest{
# only run 50 times for the example; default 999
t1$cal_NTI(null.model = "taxa.labels", abundance.weighted = TRUE, runs = 50)
}

```

Method `cal_Cscore()`: Calculates the (normalised) mean number of checkerboard combinations (C-score) using `C.score` function in `bipartite` package.

Usage:

```
trans_nullmodel$cal_Cscore(by_group = NULL, ...)
```

Arguments:

`by_group` default NULL; one column name or number in `sample_table`; calculate C-score for different groups separately.

... parameters pass to `bipartite::C.score` function.

Returns: vector.

Examples:

```

\dontrun{
t1$cal_Cscore(normalise = FALSE)
t1$cal_Cscore(by_group = "Group", normalise = FALSE)
}

```

Method `cal_NST()`: Calculate normalized stochasticity ratio (NST) based on the `NST` package.

Usage:

```
trans_nullmodel$cal_NST(method = "tNST", group, ...)
```

Arguments:

`method` default "tNST"; 'tNST' or 'pNST'. See the help document of `tNST` or `pNST` function in `NST` package for more details.

`group` a colname of `sample_table` in `microtable` object; the function can select the data from `sample_table` to generate a one-column (n x 1) matrix and provide it to the `group` parameter of `tNST` or `pNST` function.

... parameters pass to `NST::tNST` or `NST::pNST` function; see the document of corresponding function for more details.

Returns: res_NST stored in the object.

Examples:

```
\dontrun{
t1$cal_NST(group = "Group", dist.method = "bray", output.rand = TRUE, SES = TRUE)
}
```

Method cal_NST_test(): Test the significance of NST difference between each pair of groups.

Usage:

```
trans_nullmodel$cal_NST_test(method = "nst.boot", ...)
```

Arguments:

method default "nst.boot"; "nst.boot" or "nst.panova"; see NST::nst.boot function or NST::nst.panova function for the details.

... parameters pass to NST::nst.boot when method = "nst.boot" or NST::nst.panova when method = "nst.panova".

Returns: list. See the Return part of NST::nst.boot function or NST::nst.panova function in NST package.

Examples:

```
\dontrun{
t1$cal_NST_test()
}
```

Method cal_NST_convert(): Convert NST paired long format table to symmetric matrix form.

Usage:

```
trans_nullmodel$cal_NST_convert(column = 10)
```

Arguments:

column default 10; which column is selected for the conversion. See the columns of res_NST\$index.pair stored in the object.

Returns: symmetric matrix.

Examples:

```
\dontrun{
t1$cal_NST_convert(column = 10)
}
```

Method clone(): The objects of this class are cloneable with this method.

Usage:

```
trans_nullmodel$clone(deep = FALSE)
```

Arguments:

deep Whether to make a deep clone.

Examples

```

## -----
## Method `trans_nullmodel$new`
## -----

data(dataset)
data(env_data_16S)
t1 <- trans_nullmodel$new(dataset, filter_thres = 0.0005, add_data = env_data_16S)

## -----
## Method `trans_nullmodel$cal_mantel_corr`
## -----

## Not run:
t1$cal_mantel_corr(use_env = "pH")

## End(Not run)

## -----
## Method `trans_nullmodel$plot_mantel_corr`
## -----

## Not run:
t1$plot_mantel_corr()

## End(Not run)

## -----
## Method `trans_nullmodel$cal_betampd`
## -----

t1$cal_betampd(abundance.weighted = TRUE)

## -----
## Method `trans_nullmodel$cal_betamntd`
## -----

t1$cal_betamntd(abundance.weighted = TRUE)

## -----
## Method `trans_nullmodel$cal_ses_betampd`
## -----

## Not run:
# only run 50 times for the example; default 1000
t1$cal_ses_betampd(runs = 50, abundance.weighted = TRUE)

## End(Not run)

```

```

## -----
## Method `trans_nullmodel$cal_ses_betamntd`
## -----

## Not run:
# only run 50 times for the example; default 1000
t1$cal_ses_betamntd(runs = 50, abundance.weighted = TRUE, exclude.conspecifics = FALSE)

## End(Not run)

## -----
## Method `trans_nullmodel$cal_rcbray`
## -----

## Not run:
# only run 50 times for the example; default 1000
t1$cal_rcbray(runs = 50)

## End(Not run)

## -----
## Method `trans_nullmodel$cal_process`
## -----

## Not run:
t1$cal_process(use_betamntd = TRUE)

## End(Not run)

## -----
## Method `trans_nullmodel$cal_NRI`
## -----

# only run 50 times for the example; default 999
t1$cal_NRI(null.model = "taxa.labels", abundance.weighted = FALSE, runs = 50)

## -----
## Method `trans_nullmodel$cal_NTI`
## -----

# only run 50 times for the example; default 999
t1$cal_NTI(null.model = "taxa.labels", abundance.weighted = TRUE, runs = 50)

## -----
## Method `trans_nullmodel$cal_Cscore`
## -----

## Not run:

```

```

t1$cal_Cscore(normalise = FALSE)
t1$cal_Cscore(by_group = "Group", normalise = FALSE)

## End(Not run)

## -----
## Method `trans_nullmodel$cal_NST`
## -----

## Not run:
t1$cal_NST(group = "Group", dist.method = "bray", output.rand = TRUE, SES = TRUE)

## End(Not run)

## -----
## Method `trans_nullmodel$cal_NST_test`
## -----

## Not run:
t1$cal_NST_test()

## End(Not run)

## -----
## Method `trans_nullmodel$cal_NST_convert`
## -----

## Not run:
t1$cal_NST_convert(column = 10)

## End(Not run)

```

trans_venn

Create trans_venn object for the Venn diagram, petal plot and UpSet plot.

Description

This class is a wrapper for a series of intersection analysis related methods, including 2- to 5-way venn diagram, more than 5-way petal or UpSet plot and intersection transformations based on David et al. (2012) <doi:10.1128/AEM.01459-12>.

Methods

Public methods:

- `trans_venn$new()`
- `trans_venn$plot_venn()`
- `trans_venn$plot_bar()`
- `trans_venn$trans_comm()`

- `trans_venn$print()`
- `trans_venn$clone()`

Method new():

Usage:

```
trans_venn$new(
  dataset,
  ratio = NULL,
  name_joint = "&",
  ratio_both_joint = "; "
)
```

Arguments:

dataset the object of `microtable` class or a matrix-like table (data.frame or matrix object). If dataset is a matrix-like table, features must be rows.

ratio default NULL; NULL, "numratio", "seqratio" or "both"; "numratio": calculate the percentage of feature number; "seqratio": calculate the percentage of feature abundance; "both": "numratio" + "seqratio"; NULL: no additional percentage (raw counts).

name_joint default "&"; the joint mark for generating multi-sample names.

ratio_both_joint default "; "; the joint mark for the "both" option in ratio parameter. If you need to insert a line break, you can use "\n".

Returns: data_details and data_summary stored in the object.

Examples:

```
\donttest{
data(dataset)
t1 <- dataset$merge_samples("Group")
t1 <- trans_venn$new(dataset = t1, ratio = "numratio")
}
```

Method plot_venn(): Plot venn diagram.

Usage:

```
trans_venn$plot_venn(
  color_circle = RColorBrewer::brewer.pal(8, "Dark2"),
  fill_color = TRUE,
  text_size = 4.5,
  text_name_size = 6,
  text_name_position = NULL,
  alpha = 0.3,
  linesize = 1.1,
  petal_plot = FALSE,
  petal_color = "#BEAED4",
  petal_color_center = "#BEBADA",
  petal_a = 4,
  petal_r = 1,
  petal_use_lim = c(-12, 12),
  petal_center_size = 40,
  petal_move_xy = 4,
```

```

    petal_move_k = 2.3,
    petal_move_k_count = 1.3,
    petal_text_move = 40,
    other_text_show = NULL,
    other_text_position = c(2, 2),
    other_text_size = 5
  )

```

Arguments:

`color_circle` default `RColorBrewer::brewer.pal(8, "Dark2")`; color palette.

`fill_color` default `TRUE`; whether fill the area color.

`text_size` default 4.5; text size in plot.

`text_name_size` default 6; name size in plot.

`text_name_position` default `NULL`; name position in plot.

`alpha` default .3; alpha for transparency.

`linesize` default 1.1; cycle line size.

`petal_plot` default `FALSE`; whether use petal plot.

`petal_color` default `"#BEAED4"`; color of the petals; If `petal_color` only has one color value, all the petals will be assigned with this color value. If `petal_color` has multiple colors, and the number of color values is smaller than the petal number, the function can append more colors automatically with the color interpolation.

`petal_color_center` default `"#BEBADA"`; color of the center in the petal plot.

`petal_a` default 4; the length of the ellipse.

`petal_r` default 1; scaling up the size of the ellipse.

`petal_use_lim` default `c(-12, 12)`; the width of the plot.

`petal_center_size` default 40; petal center circle size.

`petal_move_xy` default 4; the distance of text to circle.

`petal_move_k` default 2.3; the distance of title to circle.

`petal_move_k_count` default 1.3; the distance of data text to circle.

`petal_text_move` default 40; the distance between two data text.

`other_text_show` default `NULL`; other characters used to show in the plot.

`other_text_position` default `c(1, 1)`; the text position for text in `other_text_show`.

`other_text_size` default 5; the text size for text in `other_text_show`.

Returns: `ggplot`.

Examples:

```

\donttest{
t1$plot_venn()
}

```

Method `plot_bar()`: Plot the intersections using histogram, i.e. UpSet plot. Especially useful when samples > 5.

Usage:

```

trans_venn$plot_bar(
  left_plot = TRUE,

```

```

    sort_samples = FALSE,
    up_y_title = "Intersection size",
    up_y_title_size = 15,
    up_y_text_size = 8,
    up_bar_fill = "grey70",
    up_bar_width = 0.9,
    bottom_y_text_size = 12,
    bottom_height = 1,
    bottom_point_size = 3,
    bottom_point_color = "black",
    bottom_background_fill = "grey95",
    bottom_background_alpha = 1,
    bottom_line_width = 0.5,
    bottom_line_colour = "black",
    left_width = 0.3,
    left_bar_fill = "grey70",
    left_bar_alpha = 1,
    left_bar_width = 0.9,
    left_x_text_size = 10,
    left_background_fill = "white",
    left_background_alpha = 1
)

```

Arguments:

`left_plot` default TRUE; whether add the left bar plot to show the feature number of each sample.

`sort_samples` default FALSE; TRUE is used to sort samples according to the number of features in each sample. FALSE means the sample order is same with that in `sample_table` of the raw dataset.

`up_y_title` default "Intersection set"; y axis title of upper plot.

`up_y_title_size` default 15; y axis title size of upper plot.

`up_y_text_size` default 4; y axis text size of upper plot.

`up_bar_fill` default "grey70"; bar fill color of upper plot.

`up_bar_width` default 0.9; bar width of upper plot.

`bottom_y_text_size` default 12; y axis text size, i.e. sample name size, of bottom sample plot.

`bottom_height` default 1; bottom plot height relative to the upper bar plot. 1 represents the height of bottom plot is same with the upper bar plot.

`bottom_point_size` default 3; point size of bottom plot.

`bottom_point_color` default "black"; point color of bottom plot.

`bottom_background_fill` default "grey95"; fill color for the striped background in the bottom sample plot. If the parameter length is 1, use "white" to distinguish the color stripes. If the parameter length is greater than 1, use all provided colors.

`bottom_background_alpha` default 1; the color transparency for the parameter `bottom_background_fill`.

`bottom_line_width` default 0.5; the line width in the bottom plot.

`bottom_line_colour` default "black"; the line color in the bottom plot.

`left_width` default 0.3; left bar plot width relative to the right bottom plot.

`left_bar_fill` default "grey70"; fill color for the left bar plot presenting feature number.
`left_bar_alpha` default 1; the color transparency for the parameter `left_bar_fill`.
`left_bar_width` default 0.9; bar width of left plot.
`left_x_text_size` default 10; x axis text size of the left bar plot.
`left_background_fill` default "white"; fill color for the striped background in the left plot.
 If the parameter length is 1, use "white" to distinguish the color stripes. If the parameter length is greater than 1, use all provided colors.
`left_background_alpha` default 1; the color transparency for the parameter `left_background_fill`.

Returns: a ggplot2 object.

Examples:

```
\donttest{
t2 <- t1$plot_bar()
}
```

Method `trans_comm()`: Transform intersection result to community-like microtable object for further composition analysis.

Usage:

```
trans_venn$trans_comm(use_frequency = TRUE)
```

Arguments:

`use_frequency` default TRUE; whether only use OTUs occurrence frequency, i.e. presence/absence data; if FALSE, use abundance data.

Returns: a new `microtable` class.

Examples:

```
\donttest{
t2 <- t1$trans_comm(use_frequency = TRUE)
}
```

Method `print()`: Print the `trans_venn` object.

Usage:

```
trans_venn$print()
```

Method `clone()`: The objects of this class are cloneable with this method.

Usage:

```
trans_venn$clone(deep = FALSE)
```

Arguments:

`deep` Whether to make a deep clone.

Examples

```
## -----
## Method `trans_venn$new`
## -----
```

```
data(dataset)
t1 <- dataset$merge_samples("Group")
t1 <- trans_venn$new(dataset = t1, ratio = "numratio")

## -----
## Method `trans_venn$plot_venn`
## -----

t1$plot_venn()

## -----
## Method `trans_venn$plot_bar`
## -----

t2 <- t1$plot_bar()

## -----
## Method `trans_venn$trans_comm`
## -----

t2 <- t1$trans_comm(use_frequency = TRUE)
```

Index

* **Description**

microeco, [6](#)

* **datasets**

dataset, [3](#)

env_data_16S, [5](#)

fungi_func_FungalTraits, [5](#)

fungi_func_FUNGuild, [5](#)

otu_table_16S, [19](#)

otu_table_ITS, [19](#)

phylo_tree_16S, [20](#)

prok_func_FAPROTAX, [20](#)

prok_func_NJC19_list, [20](#)

sample_info_16S, [21](#)

sample_info_ITS, [21](#)

Tax4Fun2_KEGG, [21](#)

taxonomy_table_16S, [22](#)

taxonomy_table_ITS, [22](#)

aov, [36](#)

clone, [2](#)

data.frame, [23](#)

dataset, [3](#)

dropallfactors, [4](#)

env_data_16S, [5](#)

fungi_func_FungalTraits, [5](#)

fungi_func_FUNGuild, [5](#)

grepl, [14](#)

microeco, [6](#)

microtable, [6](#), [6](#), [24](#), [35](#), [42](#), [52](#), [62](#), [72](#), [88](#),
[96](#), [108](#), [112](#), [117](#), [127](#), [130](#)

otu_table_16S, [19](#)

otu_table_ITS, [19](#)

phylo_tree_16S, [20](#)

prok_func_FAPROTAX, [20](#)

prok_func_NJC19_list, [20](#)

sample, [115](#)

sample_info_16S, [21](#)

sample_info_ITS, [21](#)

Tax4Fun2_KEGG, [21](#)

taxonomy_table_16S, [22](#)

taxonomy_table_ITS, [22](#)

tidy_taxonomy, [22](#)

trans_abund, [6](#), [24](#)

trans_alpha, [6](#), [35](#), [48](#), [49](#), [67](#), [74](#)

trans_beta, [6](#), [41](#)

trans_classifier, [6](#), [51](#)

trans_diff, [6](#), [61](#)

trans_env, [6](#), [40](#), [69](#), [72](#)

trans_func, [6](#), [88](#)

trans_network, [6](#), [95](#)

trans_norm, [6](#), [10](#), [64](#), [112](#)

trans_nullmodel, [6](#), [116](#)

trans_venn, [6](#), [126](#)

write.table, [12](#), [14](#)