# Package 'microbiomeMarker'

May 13, 2024

Title microbiome biomarker analysis toolkit

**Version** 1.10.0

**Description** To date, a number of methods have been developed for microbiome marker discovery based on metagenomic profiles, e.g. LEfSe. However, all of these methods have its own advantages and disadvantages, and none of them is considered standard or universal. Moreover, different programs or softwares may be development using different programming languages, even in different operating systems. Here, we have developed an all-in-one R package microbiomeMarker that integrates commonly used differential analysis methods as well as three machine learning-based approaches, including Logistic regression, Random forest, and Support vector machine, to facilitate the identification of microbiome markers.

# License GPL-3

biocViews Metagenomics, Microbiome, DifferentialExpression

URL https://github.com/yiluheihei/microbiomeMarker

BugReports https://github.com/yiluheihei/microbiomeMarker/issues

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microbiomeMarker-package

microbiomeMarker: A package for microbiome biomarker discovery

# Description

The microboimeMarker package provides several methods to identify micribome biomarker, such as lefse, deseq2.

abundances

Extract taxa abundances

# Description

Extract taxa abundances from phyloseq objects.

```
abundances(object, transform = c("identity", "log10", "log10p"), norm = FALSE)
## S4 method for signature 'otu_table'
abundances(object, transform = c("identity", "log10", "log10p"), norm = FALSE)
## S4 method for signature 'phyloseq'
abundances(object, transform = c("identity", "log10", "log10p"), norm = FALSE)
## S4 method for signature 'microbiomeMarker'
abundances(object, transform = c("identity", "log10", "log10p"))
```

#### Arguments

object	otu_table, phyloseq, or microbiomeMarker.
transform	transformation to apply, the options inclulde:
	• "identity", return the original data without any transformation.
	<ul> <li>"log10", the transformation is log10(object), and if the data contains zeros the transformation is log10(1 + object).</li> </ul>
	<ul> <li>"log10p", the transformation is log10(1 + object).</li> </ul>
norm	logical, indicating whether or not to return the normalized taxa abundances.

# Value

abundance matrix with taxa in rows and samples in columns.

#### See Also

otu\_table, phyloseq, microbiomeMarker, transform\_abundances

#### Examples

```
data(caporaso)
abd <- abundances(caporaso)</pre>
```

aggregate\_taxa Aggregate Taxa

#### Description

Summarize phyloseq data into a higher phylogenetic level.

#### Usage

```
aggregate_taxa(x, level, verbose = FALSE)
```

#### assign-otu\_table

#### Arguments

х	phyloseq-class object
level	Summarization level (from rank_names(pseq))
verbose	verbose

#### Details

This provides a convenient way to aggregate phyloseq OTUs (or other taxa) when the phylogenetic tree is missing. Calculates the sum of OTU abundances over all OTUs that map to the same higher-level group. Removes ambiguous levels from the taxonomy table. Returns a phyloseq object with the summarized abundances.

#### Value

Summarized phyloseq object

#### Author(s)

Contact: Leo Lahti <microbiome-admin@googlegroups.com>

#### References

```
See citation('microbiome')
```

#### Examples

```
data(caporaso)
caporaso_phylum <- aggregate_taxa(caporaso, "Phylum")</pre>
```

assign-otu\_table Assign a new OTU table

# Description

Assign a new OTU table in microbiomeMarker object

#### Usage

```
## S4 replacement method for signature 'microbiomeMarker,otu_table'
otu_table(x) <- value</pre>
```

## S4 replacement method for signature 'microbiomeMarker,phyloseq'
otu\_table(x) <- value</pre>

```
## S4 replacement method for signature 'microbiomeMarker,microbiomeMarker'
otu_table(x) <- value</pre>
```

# Arguments

х	microbiomeMarker
value	<pre>otu_table, phyloseq, or microbiomeMarker</pre>

#### Value

a microbiomeMarker object.

compare_DA	Comparing the results of differential analysis methods by Empirical
	power and False Discovery Rate

# Description

Calculating power, false discovery rates, false positive rates and auc ( area under the receiver operating characteristic (ROC) curve) for various DA methods.

# Usage

```
compare_DA(
   ps,
   group,
   taxa_rank = "none",
   methods,
   args = list(),
   n_rep = 20,
   effect_size = 5,
   k = NULL,
   relative = TRUE,
   BPPARAM = BiocParallel::SnowParam(progressbar = TRUE)
)
```

# Arguments

ps, group, taxa\_rank

	main arguments of all differential analysis methods. ps: a phyloseq::phyloseq object; group, character, the variable to set the group, must be one of the var of the sample metadata; taxa_rank: character, taxonomic rank, please not that since the abundance table is spiked in the lowest level, only taxa_rank = "none" is allowed.
methods	character vector, differential analysis methods to be compared, available meth- ods are "aldex", "ancom", "ancombc", "deseq2", "edger", "lefse", "limma_voom", "metagenomeseq", "simple_stat".
args	named list, which used to set the extra arguments of the differential analysis methods, so the names must be contained in methods. For more see details below.

# confounder

n_rep	integer, number of times to run the differential analyses.
effect_size	numeric, the effect size for the spike-ins. Default 5.
k	numeric vector of length 3, number of features to spike in each tertile (lower, mid, upper), e.g. $k=c(5,10,15)$ means 5 features spiked in low abundance tertile, 10 features spiked in mid abundance tertile and 15 features spiked in high abundance tertile. Default NULL, which will spike 2 percent of the total amount of features in each tertile (a total of 6 percent), but minimum c(5,5,5).
relative	logical, whether rescale the total number of individuals observed for each sample to the original level after spike-in. Default TRUE.
BPPARAM	BiocParallel::BiocParallelParam instance defining the parallel back-end.

# Details

To make this function support for different arguments for a certain DA method args allows list of list of list e.g. args = list(lefse = list(list(norm = "CPM"), list(norm = "TSS"))), which specify to compare the different norm arguments for lefse analysis.

For taxa\_rank, only taxa\_rank = "none" is supported, if this argument is not "none", it will be forced to "none" internally.

#### Value

an compareDA object, which contains a two-length list of:

- metrics: data.frame, FPR, AUC and spike detection rate for each run.
- mm: differential analysis results.

confounder

Confounder analysis

#### Description

Confounding variables may mask the actual differential features. This function utilizes constrained correspondence analysis (CCA) to measure the confounding factors.

# Usage

```
confounder(
   ps,
   target_var,
   norm = "none",
   confounders = NULL,
   permutations = 999,
   ...
)
```

#### Arguments

ps	a phyloseq::phyloseq object.
target_var	character, the variable of interest
norm	norm the methods used to normalize the microbial abundance data. See normalize() for more details.
confounders	the confounding variables to be measured, if NULL, all variables in the meta data will be analyzed.
permutations	the number of permutations, see vegan::anova.cca().
	extra arguments passed to vegan::anova.cca().

#### Value

a data.frame contains three variables: confounder, pseudo-F and p value.

#### Examples

```
data(caporaso)
confounder(caporaso, "SampleType", confounders = "ReportedAntibioticUsage")
```

data-caporaso

16S rRNA data from "Moving pictures of the human microbiome"

# Description

16S read counts and phylogenetic tree file of 34 Illumina samples derived from Moving Pictures of the Human Microbiome (Caporaso et al.) Group label: gut, left palm, right palm, and tongue - indicating different sampled body sites.

# Format

a phyloseq::phyloseq object

#### Author(s)

Yang Cao

# Source

Data was downloaded from https://www.microbiomeanalyst.ca

# References

Caporaso, et al. Moving pictures of the human microbiome. Genome Biol 12, R50 (2011). https://doi.org/10.1186/gb-2011-12-5-r50 data-cid\_ying

#### Description

Data from a cohort of 94 Bone Marrow Transplant patients previously published on in CID

#### Format

a phyloseq::phyloseq object

#### Author(s)

Yang Cao

# Source

https://github.com/ying14/yingtools2/tree/master/data

# References

Ying, et al. Intestinal Domination and the Risk of Bacteremia in Patients Undergoing Allogeneic Hematopoietic Stem Cell Transplantation, Clinical Infectious Diseases, Volume 55, Issue 7, 1 October 2012, Pages 905–914,

https://academic.oup.com/cid/article/55/7/905/428203

data-ecam	Data from Early Childhood Antibiotics and the Microbiome (ECAM)
	study

#### Description

The data from a subset of the Early Childhood Antibiotics and the Microbiome (ECAM) study, which tracked the microbiome composition and development of 43 infants in the United States from birth to 2 years of age, identifying microbiome associations with antibiotic exposure, delivery mode, and diet.

# Format

a phyloseq::phyloseq object

# References

Bokulich, Nicholas A., et al. "Antibiotics, birth mode, and diet shape microbiome maturation during early life." Science translational medicine 8.343 (2016): 343ra82-343ra82.

https://github.com/FrederickHuangLin/ANCOM/tree/master/data

data-enterotypes\_arumugam

Enterotypes data of 39 samples

#### Description

The data contains 22 European metagenomes from Danish, French, Italian, and Spanish individuals, and 13 Japanese and 4 American.

#### Format

a phyloseq::phyloseq object

# Author(s)

Yang Cao

# References

Arumugam, Manimozhiyan, et al. Enterotypes of the human gut microbiome. nature 473.7346 (2011): 174-180.

data-kostic\_crc Data from a study on colorectal cancer (kostic 2012)

#### Description

The data from a study on colorectal cancer. Samples that had no DIAGNOSIS attribute assigned and with less than 500 reads (counts) were removed, and 191 samples remains (91 healthy and 86 Tumors).

## Format

a phyloseq::phyloseq object

#### Author(s)

Yang Cao

#### References

Kostic et al. Genomic analysis identifies association of Fusobacterium with colorectal carcinoma. Genome research, 2012, 22(2), 292-298.

data-oxygen

Oxygen availability 16S dataset, of which taxa table has been summarized for python lefse input

## Description

A small subset of the HMP 16S dataset for finding biomarkers characterizing different level of oxygen availability in different bodysites

#### Format

a phyloseq::phyloseq object

#### Author(s)

Yang Cao

# Source

http://huttenhower.sph.harvard.edu/webfm\_send/129

data-pediatric\_ibd IBD stool samples

# Description

43 pediatric IBD stool samples obtained from the Integrative Human Microbiome Project Consortium (iHMP). Group label: CD and Controls.

# Format

a phyloseq::phyloseq object

# Author(s)

Yang Cao

#### Source

https://www.microbiomeanalyst.ca/MicrobiomeAnalyst/resources

data-spontaneous\_colitis

This is a sample data from lefse python script, a 16S dataset for studying the characteristics of the fecal microbiota in a mouse model of spontaneous colitis.

# Description

The dataset contains 30 abundance profiles (obtained processing the 16S reads with RDP) belonging to 10 rag2 (control) and 20 truc (case) mice.

#### Format

a phyloseq::phyloseq object

# Author(s)

Yang Cao

# Source

http://www.huttenhower.org/webfm\_send/73

extract\_posthoc\_res Extract results from a posthoc test

#### Description

This function extracts the results of posthoc test.

# Usage

```
extract_posthoc_res(object, features = NULL)
```

#### Arguments

object	a postHocTest object.
features	either NULL extracts results of all features, or a character vector to specify the test results of which features are extracted.

#### Value

a IRanges::SimpleDFrameList object.

# Examples

```
require(IRanges)
pht <- postHocTest(</pre>
    result = DataFrameList(
        featureA = DataFrame(
            comparisons = c("group2-group1",
                "group3-group1",
                "group3-group2"),
            diff_mean = runif(3),
            pvalue = rep(0.01, 3),
            ci_{lower} = rep(0.01, 3),
            ci_upper = rep(0.011, 3)
        ),
        featureB = DataFrame(
            comparisons = c("group2-group1",
                "group3-group1",
                "group3-group2"),
            diff_mean = runif(3),
            pvalue = rep(0.01, 3),
            ci_lower = rep(0.01, 3),
            ci_upper = rep(0.011, 3)
        )
   ),
    abundance = data.frame(
        featureA = runif(3),
        featureB = runif(3),
        group = c("group1", "group2", "grou3")
   )
)
extract_posthoc_res(pht, "featureA")[[1]]
```

get\_treedata\_phyloseq Generate tree data from phyloseq object

# Description

Generate tree data from phyloseq object

#### Usage

```
get_treedata_phyloseq(ps, sep = "|")
```

#### Arguments

ps	a phyloseq::phyloseq object
sep	character, separate between different levels of taxa, default

# Value

a tidytree::treedata object

#### Author(s)

Yang Cao

import\_dada2 Import ful

#### Import function to read the the output of dada2 as phyloseq object

# Description

Import the output of dada2 into phyloseq object

#### Usage

```
import_dada2(
   seq_tab,
   tax_tab = NULL,
   sam_tab = NULL,
   phy_tree = NULL,
   keep_taxa_rows = TRUE
)
```

# Arguments

seq_tab	matrix-like, ASV table, the output of dada2::removeBimeraDenovo.
tax_tab	matrix, taxonomy table, the output of dada2::assignTaxonomy or dada2::addSpecies
sam_tab	data.frame or phyloseq::sample_data, sample data
phy_tree	ape::phylo class or character represents the path of the tree file
keep_taxa_rows	logical, whether keep taxa in rows or not in the otu_table of the returned phyloseq object, default TRUE.

# Details

The output of the dada2 pipeline is a feature table of amplicon sequence variants (an ASV table): A matrix with rows corresponding to samples and columns to ASVs, in which the value of each entry is the number of times that ASV was observed in that sample. This table is analogous to the traditional OTU table. Conveniently, taxa names are saved as ASV1, ASV2, ..., in the returned phyloseq object.

# Value

phyloseq::phyloseq object hold the taxonomy info, sample metadata, number of reads per ASV.

#### import\_picrust2

#### Examples

```
seq_tab <- readRDS(system.file("extdata", "dada2_seqtab.rds",
    package = "microbiomeMarker"
))
tax_tab <- readRDS(system.file("extdata", "dada2_taxtab.rds",
    package = "microbiomeMarker"
))
sam_tab <- read.table(system.file("extdata", "dada2_samdata.txt",
    package = "microbiomeMarker"
), sep = "\t", header = TRUE, row.names = 1)
ps <- import_dada2(seq_tab = seq_tab, tax_tab = tax_tab, sam_tab = sam_tab)
ps</pre>
```

import\_picrust2 Import function to read the output of picrust2 as phyloseq object

#### Description

Import the output of picrust2 into phyloseq object

#### Usage

```
import_picrust2(
  feature_tab,
  sam_tab = NULL,
  trait = c("PATHWAY", "COG", "EC", "KO", "PFAM", "TIGRFAM", "PHENO")
)
```

#### Arguments

feature_tab	character, file path of the prediction abundance table of functional feature.
sam_tab	character, file path of the sample meta data.
trait	character, options are picrust2 function traits (including "COG", "EC", "KO" "PFAM", "TIGRFAM", and "PHENO") and "PATHWAY".

#### Details

**PICRUSt2** is a software for predicting abundances of functional profiles based on marker gene sequencing data. The functional profiles can be predicted from the taxonomic profiles using PI-CRUSt2. "Function" usually refers to gene families such as KEGG orthologs and Enzyme Classification numbers, but predictions can be made for any arbitrary trait.

In the phyloseq object, the predicted function abundance profile is stored in otu\_table slot. And the functional trait is saved in tax\_table slot, if the descriptions of function features is not added to the predicted table, tax\_table will have only one rank Picrust\_trait to represent the function feature id, or if the descriptions are added one more rank Picrust\_description will be added to represent the description of function feature.

# Value

phyloseq::phyloseq object.

# Examples

```
sam_tab <- system.file(
    "extdata", "picrust2_metadata.tsv",
    package = "microbiomeMarker")
feature_tab <- system.file(
    "extdata", "path_abun_unstrat_descrip.tsv.gz",
    package = "microbiomeMarker")
ps <- import_picrust2(feature_tab, sam_tab, trait = "PATHWAY")
ps</pre>
```

import\_qiime2 Import function

Import function to read the the output of dada2 as phyloseq object

# Description

Import the qiime2 artifacts, including feature table, taxonomic table, phylogenetic tree, representative sequence and sample metadata into phyloseq object.

#### Usage

```
import_qiime2(
   otu_qza,
   taxa_qza = NULL,
   sam_tab = NULL,
   refseq_qza = NULL,
   tree_qza = NULL
)
```

# Arguments

otu_qza	character, file path of the feature table from qiime2.
taxa_qza	character, file path of the taxonomic table from qiime2, default NULL.
sam_tab	character, file path of the sample metadata in tsv format, default NULL.
refseq_qza	character, file path of the representative sequences from qiime2, default NULL.
tree_qza	character, file path of the phylogenetic tree from qiime2, default NULL.

## Value

phyloseq::phyloseq object.

#### marker\_table

#### Examples

```
otuqza_file <- system.file(</pre>
    "extdata", "table.qza",
    package = "microbiomeMarker"
)
taxaqza_file <- system.file(</pre>
    "extdata", "taxonomy.qza",
package = "microbiomeMarker"
)
sample_file <- system.file(</pre>
    "extdata", "sample-metadata.tsv",
    package = "microbiomeMarker"
)
treeqza_file <- system.file(</pre>
    "extdata", "tree.qza",
    package = "microbiomeMarker"
)
ps <- import_qiime2(</pre>
    otu_qza = otuqza_file, taxa_qza = taxaqza_file,
    sam_tab = sample_file, tree_qza = treeqza_file
)
ps
```

marker\_table Build or access the marker\_table

# Description

This is the recommended function for both building and accessing microbiome marker table (marker\_table).

#### Usage

```
marker_table(object)
## S4 method for signature 'data.frame'
marker_table(object)
```

```
## S4 method for signature 'microbiomeMarker'
marker_table(object)
```

# Arguments

object	an object among the set of classes defined by the microbiomeMarker package
	that contain marker_table

#### Value

a marker\_table object.

# Examples

```
data(enterotypes_arumugam)
mm <- run_limma_voom(
    enterotypes_arumugam,
    "Enterotype",
    contrast = c("Enterotype 3", "Enterotype 2"),
    pvalue_cutoff = 0.05,
    p_adjust = "fdr"
)
marker_table(mm)</pre>
```

marker\_table-class The S4 class for storing microbiome marker information

#### Description

This Class is inherit from data.frame. Rows represent the microbiome markers and variables represents feature of the marker.

# Fields

names, row.names a character vector, inherited from the input data.frame

.data a list, each element corresponding the each column of the input data.frame

.S3Class character, the S3 class marker\_table inherited from: "data.frame"

#### Author(s)

Yang Cao

marker\_table<- Assign marker\_table to object</pre>

# Description

This function replace the marker\_table slot of object with value.

#### Usage

```
marker_table(object) <- value</pre>
```

#### Arguments

object	a microbiomeMarker object to modify.
value	new value to replace the marker_table slot of object. Either a marker_table-class,
	a data.frame that can be coerced into marker_table-class.

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#### microbiomeMarker

# Value

a microbiomeMarker object.

#### Examples

```
data(enterotypes_arumugam)
mm <- run_limma_voom(
    enterotypes_arumugam,
    "Enterotype",
    contrast = c("Enterotype 3", "Enterotype 2"),
    pvalue_cutoff = 0.1,
    p_adjust = "fdr"
)
mm_marker <- marker_table(mm)
mm_marker
marker_table(mm) <- mm_marker[1:2, ]
marker_table(mm)</pre>
```

microbiomeMarker Build microbiomeMarker-class objects

# Description

This the constructor to build the microbiomeMarker object, don't use the new() constructor.

# Usage

```
microbiomeMarker(
  marker_table = NULL,
  norm_method = NULL,
  diff_method = NULL,
   ...
)
```

#### Arguments

marker_table	a marker_table object differtial analysis.
norm_method	character, method used to normalize the input phyloseq object.
diff_method	character, method used for microbiome marker identification.
	arguments passed to phyloseq::phyloseq()

# Value

a microbiomeMarker object.

#### See Also

phyloseq::phyloseq()

#### Examples

```
microbiomeMarker(
   marker_table = marker_table(data.frame(
        feature = c("speciesA", "speciesB"),
        enrich_group = c("groupA", "groupB"),
        ef_{logFC} = c(-2, 2),
        pvalue = c(0.01, 0.01),
        padj = c(0.01, 0.01),
        row.names = c("marker1", "marker2")
   )),
   norm_method = "TSS",
   diff_method = "DESeq2",
   otu_table = otu_table(matrix(
       c(4, 1, 1, 4),
        nrow = 2, byrow = TRUE,
        dimnames = list(c("speciesA", "speciesB"), c("sample1", "sample2"))
   ),
   taxa_are_rows = TRUE
   ),
    tax_table = tax_table(matrix(
        c("speciesA", "speciesB"),
        nrow = 2,
        dimnames = list(c("speciesA", "speciesB"), "Species")
   )),
    sam_data = sample_data(data.frame(
        group = c("groupA", "groupB"),
        row.names = c("sample1", "sample2")
   ))
)
```

microbiomeMarker-class

The main class for microbiomeMarker data

#### Description

microbiomeMarker-class is inherited from the phyloseq::phyloseq by adding a custom slot microbiome\_marker to save the differential analysis results. And it provides a seamless interface with **phyloseq**, which makes **microbiomeMarker** simple and easy to use. For more details on see the document of phyloseq::phyloseq.

#### Usage

```
## S4 method for signature 'microbiomeMarker'
show(object)
```

#### Arguments

object a microbiomeMarker-class object

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# nmarker

# Value

a microbiomeMarker object.

#### Slots

```
marker_table a data.frame, a marker_table object.
norm_method character, method used to normalize the input phyloseq object.
diff_method character, method used for marker identification.
```

# See Also

phyloseq::phyloseq,marker\_table,summarize\_taxa()

nmarker

Get the number of microbiome markers

#### Description

Get the number of microbiome markers

#### Usage

```
nmarker(object)
```

## S4 method for signature 'microbiomeMarker'
nmarker(object)

## S4 method for signature 'marker\_table'
nmarker(object)

#### Arguments

object a microbiomeMarker or marker\_table object

#### Value

an integer, the number of microbiome markers

#### Examples

```
mt <- marker_table(data.frame(
    feature = c("speciesA", "speciesB"),
    enrich_group = c("groupA", "groupB"),
    ef_logFC = c(-2, 2),
    pvalue = c(0.01, 0.01),
    padj = c(0.01, 0.01),
    row.names = c("marker1", "marker2")
))
nmarker(mt)</pre>
```

Normalize the microbial abundance data

#### Description

It is critical to normalize the feature table to eliminate any bias due to differences in the sampling sequencing depth. This function implements six widely-used normalization methods for microbial compositional data.

For rarefying, reads in the different samples are randomly removed until the same predefined number has been reached, to assure all samples have the same library size. Rarefying normalization method is the standard in microbial ecology. Please note that the authors of phyloseq do not advocate using this rarefying a normalization procedure, despite its recent popularity

TSS simply transforms the feature table into relative abundance by dividing the number of total reads of each sample.

CSS is based on the assumption that the count distributions in each sample are equivalent for low abundant genes up to a certain threshold. Only the segment of each sample's count distribution that is relatively invariant across samples is scaled by CSS

RLE assumes most features are not differential and uses the relative abundances to calculate the normalization factor.

TMM calculates the normalization factor using a robust statistics based on the assumption that most features are not differential and should, in average, be equal between the samples. The TMM scaling factor is calculated as the weighted mean of log-ratios between each pair of samples, after excluding the highest count OTUs and OTUs with the largest log-fold change.

In CLR, the log-ratios are computed relative to the geometric mean of all features.

norm\_cpm: This normalization method is from the original LEfSe algorithm, recommended when very low values are present (as shown in the LEfSe galaxy).

# Usage

```
## S4 method for signature 'phyloseq'
normalize(object, method = "TSS", ...)
## S4 method for signature 'otu_table'
normalize(object, method = "TSS", ...)
## S4 method for signature 'data.frame'
normalize(object, method = "TSS", ...)
## S4 method for signature 'matrix'
normalize(object, method = "TSS", ...)
norm_rarefy(
    object,
```

```
size = min(sample_sums(object)),
  rng_seed = FALSE,
  replace = TRUE,
  trim_otus = TRUE,
  verbose = TRUE
)
norm_tss(object)
norm_css(object, sl = 1000)
norm_rle(
  object,
  locfunc = stats::median,
  type = c("poscounts", "ratio"),
  geo_means = NULL,
  control_genes = NULL
)
norm_tmm(
  object,
  ref_column = NULL,
  logratio_trim = 0.3,
  sum_trim = 0.05,
  do_weighting = TRUE,
  Acutoff = -1e+10
)
norm_clr(object)
norm_cpm(object)
```

# Arguments

object	a phyloseq::phyloseq or phyloseq::otu_table
method	the methods used to normalize the microbial abundance data. Options includes:
	• "none": do not normalize.
	• "rarefy": random subsampling counts to the smallest library size in the data set.
	• "TSS": total sum scaling, also referred to as "relative abundance", the abundances were normalized by dividing the corresponding sample library size.
	• "TMM": trimmed mean of m-values. First, a sample is chosen as reference. The scaling factor is then derived using a weighted trimmed mean over the differences of the log-transformed gene-count fold-change between the sample and the reference.
	• "RLE", relative log expression, RLE uses a pseudo-reference calculated using the geometric mean of the gene-specific abundances over all samples.

	<ul><li>The scaling factors are then calculated as the median of the gene counts ratios between the samples and the reference.</li><li>"CSS": cumulative sum scaling, calculates scaling factors as the cumulative sum of gene abundances up to a data-derived threshold.</li></ul>
	• "CLR": centered log-ratio normalization.
	• "CPM": pre-sample normalization of the sum of the values to 1e+06.
	other arguments passed to the corresponding normalization methods.
<pre>size, rng_seed,</pre>	replace, trim_otus, verbose
	extra arguments passed to phyloseq::rarefy_even_depth().
sl	The value to scale.
locfunc	a function to compute a location for a sample. By default, the median is used.
type	method for estimation: either "ratio" or "poscounts" (recommend).
geo_means	default NULL, which means the geometric means of the counts are used. A vector of geometric means from another count matrix can be provided for a "frozen" size factor calculation.
control_genes	default NULL, which means all taxa are used for size factor estimation, numeric or logical index vector specifying the taxa used for size factor estimation (e.g. core taxa).
ref_column	column to use as reference
logratio_trim	amount of trim to use on log-ratios
sum_trim	amount of trim to use on the combined absolute levels ("A" values)
do_weighting	whether to compute the weights or not
Acutoff	cutoff on "A" values to use before trimming

# Value

the same class with object.

# See Also

```
edgeR::calcNormFactors(),DESeq2::estimateSizeFactorsForMatrix(),metagenomeSeq::cumNorm()
phyloseq::rarefy_even_depth()
metagenomeSeq::calcNormFactors()
DESeq2::estimateSizeFactorsForMatrix()
edgeR::calcNormFactors()
```

# Examples

```
data(caporaso)
normalize(caporaso, "TSS")
```

phyloseq2DESeq2

# Description

This function convert [phyloseq::phyloseq-class] to [DESeq2::DESeqDataSet-class], which can then be tested using

# Usage

```
phyloseq2DESeq2(ps, design, ...)
```

# Arguments

ps	the [phyloseq::phyloseq-class] object to convert, which must have a [phyloseq::sample_data()'] component.
design	a formula or matrix, the formula expresses how the counts for each gene de- pend on the variables in colData. Many R formula are valid, including designs with multiple variables, e.g., ~ group + condition. This argument is passed to DESeq2::DESeqDataSetFromMatrix().
	additional arguments passed to DESeq2::DESeqDataSetFromMatrix(), Most users will not need to pass any additional arguments here.

# Value

a DESeq2::DESeqDataSet object.

# See Also

DESeq2::DESeqDataSetFromMatrix(),DESeq2::DESeq()

# Examples

```
data(caporaso)
phyloseq2DESeq2(caporaso, ~SampleType)
```

phyloseq2edgeR Convert phyloseq data to edgeR DGEL:	∟ist <i>object</i>
---	--------------------

# Description

This function convert phyloseq::phyloseq object to edgeR::DGEList object, can then can be used to perform differential analysis using the methods in edgeR.

# Usage

```
phyloseq2edgeR(ps, ...)
```

#### Arguments

ps	a phyloseq::phyloseq object.
	optional, additional named arguments passed to edgeR::DGEList(). Most users
	will not need to pass any additional arguments here.

# Value

A edgeR::DGEList object.

# Examples

data(caporaso)
dge <- phyloseq2edgeR(caporaso)</pre>

phyloseq2metagenomeSeq

Convert phyloseq data to MetagenomeSeq MRexperiment object

# Description

The phyloseq data is converted to the relevant metagenomeSeq::MRexperiment object, which can then be tested in the zero-inflated mixture model framework in the metagenomeSeq package.

#### Usage

```
phyloseq2metagenomeSeq(ps, ...)
```

```
otu_table2metagenomeSeq(ps, ...)
```

#### Arguments

ps	<pre>phyloseq::phyloseq object for phyloseq2metagenomeSeq(), or phyloseq::otu_table</pre>
	<pre>object for otu_table2metagenomeseq().</pre>
	optional, additional named arguments passed to metagenomeSeq::newMRexperiment().
	Most users will not need to pass any additional arguments here.

# Value

A metagenomeSeq::MRexperiment object.

#### See Also

```
metagenomeSeq::fitTimeSeries(), metagenomeSeq::fitLogNormal(),metagenomeSeq::fitZig(),
metagenomeSeq::MRtable(),metagenomeSeq::MRfulltable()
```

#### Examples

data(caporaso)
phyloseq2metagenomeSeq(caporaso)

plot.compareDA Plotting DA comparing result

# Description

Plotting DA comparing result

# Usage

```
## S3 method for class 'compareDA'
plot(x, sort = c("score", "auc", "fpr", "power"), ...)
```

# Arguments

х	an compareDA object, output from compare_DA().
sort	character string specifying sort method. Possibilities are "score" which is calculated as $(auc - 0.5) * power - fdr$ , "auc" for area under the ROC curve, "fpr" for false positive rate, "power" for empirical power.
	extra arguments, just ignore it.

# Value

a ggplot2::ggplot object containing 4 subplots: "auc", "fdr", "power"and "score" plot.

<pre>plot_abundance</pre>	plot the abundances of markers	
---------------------------	--------------------------------	--

# Description

plot the abundances of markers

# Usage

```
plot_abundance(mm, label_level = 1, max_label_len = 60, markers = NULL, group)
```

# Arguments

mm	a microbiomeMarker object
label_level	integer, number of label levels to be displayed, default 1, 0 means display the full name of the feature
<pre>max_label_len</pre>	integer, maximum number of characters of feature label, default 60
markers	character vector, markers to display, default NULL, indicating plot all markers.
group	character, the variable to set the group

#### Value

a ggplot2::ggplot object.

#### Examples

```
data(enterotypes_arumugam)
mm <- run_limma_voom(
    enterotypes_arumugam,
    "Enterotype",
    contrast = c("Enterotype 3", "Enterotype 2"),
    pvalue_cutoff = 0.01,
    p_adjust = "none"
)
plot_abundance(mm, group = "Enterotype")</pre>
```

plot\_cladogram

plot cladogram of micobiomeMaker results

#### Description

plot cladogram of micobiomeMaker results

#### Usage

```
plot_cladogram(
    mm,
    color,
    only_marker = FALSE,
    branch_size = 0.2,
    alpha = 0.2,
    node_size_scale = 1,
    node_size_offset = 1,
    clade_label_level = 4,
    clade_label_font_size = 4,
    annotation_shape = 22,
    annotation_shape_size = 5,
    group_legend_param = list(),
    marker_legend_param = list()
```

```
)
```

#### Arguments

mma microbiomeMarker objectcolora color vector, used to highlight the clades of microbiome biomarker. The values<br/>will be matched in order (usually alphabetical) with the groups. If this is a named<br/>vector, then the colors will be matched based on the names instead.

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only_marker	logical, whether show all the features or only markers in the cladogram, default FALSE.
branch_size	numeric, size of branch, default 0.2
alpha	alpha parameter for shading, default 0.2
<pre>node_size_scal</pre>	e
	<pre>the parameter 'a' controlling node size: node_size=a*log(relative_abundance) + b</pre>
node_size_offs	et
	<pre>the parameter 'b' controlling node size: node_size=a*log(relative_abundance) + b</pre>
clade_label_level	
	max level of taxa used to label the clade, other level of taxa will be shown on the side.
clade_label_font_size	
	font size of the clade label, default 4.
annotation_sha	pe
	shape used for annotation, default 22
annotation_shape_size	
	size used for annotation shape, default 5
group_legend_p	aram, marker_legend_param a list specifying extra parameters of group legend and marker legend, such as direction ( the direction of the guide), nrow (the desired number of rows of legends). See ggplot2::guide_legend() for more details.

# Value

a ggtree object

# Author(s)

Chenhao Li, Guangchuang Yu, Chenghao Zhu, Yang Cao

# References

This function is modified from clada.anno from microbiomeViz.

# See Also

ggtree::ggtree()

# Examples

```
data(kostic_crc)
kostic_crc_small <- phyloseq::subset_taxa(
    kostic_crc,
    Phylum %in% c("Firmicutes")
)
mm_lefse <- run_lefse(
    kostic_crc_small,</pre>
```

```
wilcoxon_cutoff = 0.01,
group = "DIAGNOSIS",
kw_cutoff = 0.01,
multigrp_strat = TRUE,
lda_cutoff = 4
)
plot_cladogram(mm_lefse, color = c("darkgreen", "red"))
```

plot\_ef\_bar

bar and dot plot of effect size of microbiomeMarker data

#### Description

bar and dot plot of effect size microbiomeMarker data. This function returns a ggplot2 object that can be saved or further customized using **ggplot2** package.

# Usage

```
plot_ef_bar(mm, label_level = 1, max_label_len = 60, markers = NULL)
plot_ef_dot(mm, label_level = 1, max_label_len = 60, markers = NULL)
```

# Arguments

mm	a microbiomeMarker object
label_level	integer, number of label levels to be displayed, default 1, 0 means display the full name of the feature $% \left( 1,1,1,1,1,1,1,1,1,1,1,1,1,1,1,1,1,1,1,$
<pre>max_label_len</pre>	integer, maximum number of characters of feature label, default 60
markers	character vector, markers to display, default NULL, indicating plot all markers.

#### Value

a ggplot project

# Examples

```
data(enterotypes_arumugam)
mm <- run_limma_voom(
    enterotypes_arumugam,
    "Enterotype",
    contrast = c("Enterotype 3", "Enterotype 2"),
    pvalue_cutoff = 0.01,
    p_adjust = "none"
)
plot_ef_bar(mm)</pre>
```

plot\_heatmap

# Description

Display the microbiome marker using heatmap, in which rows represents the marker and columns represents the samples.

# Usage

```
plot_heatmap(
    mm,
    transform = c("log10", "log10p", "identity"),
    cluster_marker = FALSE,
    cluster_sample = FALSE,
    markers = NULL,
    label_level = 1,
    max_label_len = 60,
    sample_label = FALSE,
    scale_by_row = FALSE,
    annotation_col = NULL,
    group,
    ...
)
```

# Arguments

mm	a microbiomeMarker object
transform	transformation to apply, for more details see transform_abundances():
	• "identity", return the original data without any transformation.
	• "log10", the transformation is log10(object), and if the data contains zeros the transformation is log10(1 + object).
	<ul> <li>"log10p", the transformation is log10(1 + object).</li> </ul>
cluster_marker	,cluster_sample
	logical, controls whether to perform clustering in markers (rows) and samples (cols), default FALSE.
markers	character vector, markers to display, default NULL, indicating plot all markers.
label_level	integer, number of label levels to be displayed, default 1, 0 means display the full name of the feature
<pre>max_label_len</pre>	integer, maximum number of characters of feature label, default 60
<pre>sample_label</pre>	logical, controls whether to show the sample labels in the heatmap, default FALSE.
<pre>scale_by_row</pre>	logical, controls whether to scale the heatmap by the row (marker) values, default FALSE.

annotation_col	assign colors for the top annotation using a named vector, passed to col in ComplexHeatmap::HeatmapAnnotation().
group	character, the variable to set the group
	extra arguments passed to ComplexHeatmap::Heatmap().

# Value

a ComplexHeatmap::Heatmap object.

# See Also

transform\_abundances,ComplexHeatmap::Heatmap()

# Examples

```
data(kostic_crc)
kostic_crc_small <- phyloseq::subset_taxa(
    kostic_crc,
    Phylum %in% c("Firmicutes")
)
mm_lefse <- run_lefse(
    kostic_crc_small,
    wilcoxon_cutoff = 0.01,
    group = "DIAGNOSIS",
    kw_cutoff = 0.01,
    multigrp_strat = TRUE,
    lda_cutoff = 4
)
plot_heatmap(mm_lefse, group = "DIAGNOSIS")</pre>
```

plot\_postHocTest postHocTest plot

#### Description

Visualize the result of post-hoc test using ggplot2

#### Usage

```
plot_postHocTest(pht, feature, step_increase = 0.12)
```

#### Arguments

pht	a postHocTest object
feature	character, to plot the post-toc test result of this feature
step_increase	numeric vector with the increase in fraction of total height for every additional comparison to minimize overlap, default 0.12.

#### plot\_sl\_roc

#### Value

a ggplot object

#### Examples

```
data(enterotypes_arumugam)
ps <- phyloseq::subset_samples(
    enterotypes_arumugam,
    Enterotype %in% c("Enterotype 3", "Enterotype 2", "Enterotype 1")
) %>%
    phyloseq::subset_taxa(Phylum == "Bacteroidetes")
pht <- run_posthoc_test(ps, group = "Enterotype")
plot_postHocTest(pht, feature = "p_Bacteroidetes|g_Alistipes")</pre>
```

plot\_sl\_roc ROC curve of microbiome marker from supervised learning methods

#### Description

Show the ROC curve of the microbiome marker calculated by run\_sl.

#### Usage

```
plot_sl_roc(mm, group, nfolds = 3, nrepeats = 3, tune_length = 5, ...)
```

#### Arguments

mm a microbiomeMarker object.
group, nfolds, nrepeats, tune\_length, ...
same with the run\_sl().

#### Value

a ggplot2::ggplot object.

#### See Also

run\_sl()

# Examples

```
data(enterotypes_arumugam)
# small example phyloseq object for test
ps_s <- phyloseq::subset_taxa(
    enterotypes_arumugam,
    Phylum %in% c("Firmicutes", "Bacteroidetes")
)
set.seed(2021)</pre>
```

```
mm <- run_sl(
    ps_s,
    group = "Gender",
    taxa_rank = "Genus",
    nfolds = 2,
    nrepeats = 1,
    top_n = 15,
    norm = "TSS",
    method = "LR",
)
plot_sl_roc(mm, group = "Gender")</pre>
```

postHocTest Build postHocTest object

#### Description

This function is used for create postHocTest object, and is only used for developers.

# Usage

```
postHocTest(
  result,
  abundance,
  conf_level = 0.95,
  method = "tukey",
  method_str = paste("Posthoc multiple comparisons of means: ", method)
)
```

# Arguments

result	a IRanges::SimpleDFrameList object.
abundance	data.frame.
conf_level	numeric, confidence level.
method	character, method for posthoc test.
method_str	character, illustrates which method is used for posthoc test.

#### Value

a postHocTest object.

# Examples

```
require(IRanges)
pht <- postHocTest(
    result = DataFrameList(
        featureA = DataFrame(
            comparisons = c("group2-group1",</pre>
```

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```
"group3-group1",
                "group3-group2"),
            diff_mean = runif(3),
            pvalue = rep(0.01, 3),
            ci_lower = rep(0.01, 3),
            ci_upper = rep(0.011, 3)
        ),
        featureB = DataFrame(
            comparisons = c("group2-group1",
                "group3-group1",
                "group3-group2"),
            diff_mean = runif(3),
            pvalue = rep(0.01, 3),
            ci_lower = rep(0.01, 3),
            ci_upper = rep(0.011, 3)
        )
   ),
    abundance = data.frame(
        featureA = runif(3),
        featureB = runif(3),
        group = c("group1", "group2", "grou3")
   )
pht
```

postHocTest-class	The postHocTest Class, represents the result of post-hoc test result
	among multiple groups

# Description

)

The postHocTest Class, represents the result of post-hoc test result among multiple groups

# Usage

```
## S4 method for signature 'postHocTest'
show(object)
```

#### Arguments

object a postHocTest-class object

# Value

a postHocTest object.

result a IRanges::DataFrameList, each DataFrame consists of five variables:

- comparisons: character, specify which two groups to test (the group names are separated by "\_)
- diff\_mean: numeric, difference in mean abundances
- pvalue: numeric, p values
- ci\_lower and ci\_upper: numeric, lower and upper confidence interval of difference in mean abundances

abundance abundance of each feature in each group

conf\_level confidence level

method method used for post-hoc test

method\_str method illustration

# Author(s)

Yang Cao

reexports

Objects exported from other packages

#### Description

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

#### magrittr %>%

phyloseq import\_biom, import\_mothur, import\_qiime, nsamples, ntaxa, otu\_table, sample\_data, sample\_names, tax\_table, taxa\_names

run\_aldex

Perform differential analysis using ALDEx2

#### Description

Perform differential analysis using ALDEx2

# run\_aldex

# Usage

```
run_aldex(
    ps,
    group,
    taxa_rank = "all",
    transform = c("identity", "log10", "log10p"),
    norm = "none",
    norm_para = list(),
    method = c("t.test", "wilcox.test", "kruskal", "glm_anova"),
    p_adjust = c("none", "fdr", "bonferroni", "holm", "hochberg", "hommel", "BH", "BY"),
    pvalue_cutoff = 0.05,
    mc_samples = 128,
    denom = c("all", "iqlr", "zero", "lvha"),
    paired = FALSE
)
```

ps	a phyloseq::phyloseq object
group	character, the variable to set the group
taxa_rank	character to specify taxonomic rank to perform differential analysis on. Should be one of phyloseq::rank_names(phyloseq), or "all" means to summarize the taxa by the top taxa ranks (summarize_taxa(ps, level = rank_names(ps)[1])), or "none" means perform differential analysis on the original taxa (taxa_names(phyloseq), e.g., OTU or ASV).
transform	character, the methods used to transform the microbial abundance. See transform_abundances() for more details. The options include:
	<ul> <li>"identity", return the original data without any transformation (default).</li> <li>"log10", the transformation is log10(object), and if the data contains zeros the transformation is log10(1 + object).</li> <li>"log10p", the transformation is log10(1 + object).</li> </ul>
norm	the methods used to normalize the microbial abundance data. See normalize() for more details. Options include:
	<ul> <li>"none": do not normalize.</li> <li>"rarefy": random subsampling counts to the smallest library size in the data set.</li> <li>"TSS": total sum scaling, also referred to as "relative abundance", the abundances were normalized by dividing the corresponding sample library size.</li> <li>"TMM": trimmed mean of m-values. First, a sample is chosen as reference. The scaling factor is then derived using a weighted trimmed mean over the differences of the log-transformed gene-count fold-change between the sample and the reference.</li> <li>"RLE", relative log expression, RLE uses a pseudo-reference calculated using the geometric mean of the gene-specific abundances over all samples. The scaling factors are then calculated as the median of the gene counts ratios between the samples and the reference.</li> </ul>

	• "CSS": cumulative sum scaling, calculates scaling factors as the cumulative sum of gene abundances up to a data-derived threshold.
	• "CLR": centered log-ratio normalization.
	• "CPM": pre-sample normalization of the sum of the values to 1e+06.
norm_para	arguments passed to specific normalization methods
method	test method, options include: "t.test" and "wilcox.test" for two groups compari- son, "kruskal" and "glm_anova" for multiple groups comparison.
p_adjust	method for multiple test correction, default none, for more details see stats::p.adjust.
<pre>pvalue_cutoff</pre>	cutoff of p value, default 0.05.
<pre>mc_samples</pre>	integer, the number of Monte Carlo samples to use for underlying distributions estimation, 128 is usually sufficient.
denom	character string, specify which features used to as the denominator for the geo- metric mean calculation. Options are:
	• "all", with all features.
	• "iqlr", accounts for data with systematic variation and centers the features on the set features that have variance that is between the lower and upper quartile of variance.
	<ul> <li>"zero", a more extreme case where there are many non-zero features in one condition but many zeros in another. In this case the geometric mean of each group is calculated using the set of per-group non-zero features.</li> <li>"lvha", with house keeping features.</li> </ul>
paired	logical, whether to perform paired tests, only worked for method "t.test" and "wilcox.test".

# Value

a microbiomeMarker object.

# References

Fernandes, A.D., Reid, J.N., Macklaim, J.M. et al. Unifying the analysis of high-throughput sequencing datasets: characterizing RNA-seq, 16S rRNA gene sequencing and selective growth experiments by compositional data analysis. Microbiome 2, 15 (2014).

# See Also

```
ALDEx2::aldex()
```

# Examples

```
data(enterotypes_arumugam)
ps <- phyloseq::subset_samples(
    enterotypes_arumugam,
    Enterotype %in% c("Enterotype 3", "Enterotype 2")
)
run_aldex(ps, group = "Enterotype")</pre>
```

run\_ancom

# Description

Perform significant test by comparing the pairwise log ratios between all features.

# Usage

```
run_ancom(
    ps,
    group,
    confounders = character(0),
    taxa_rank = "all",
    transform = c("identity", "log10", "log10p"),
    norm = "TSS",
    norm_para = list(),
    p_adjust = c("none", "fdr", "bonferroni", "holm", "hochberg", "hommel", "BH", "BY"),
    pvalue_cutoff = 0.05,
    W_cutoff = 0.75
)
```

ps	a phyloseq-class object.
group	character, the variable to set the group.
confounders	character vector, the confounding variables to be adjusted. default character(0), indicating no confounding variable.
taxa_rank	character to specify taxonomic rank to perform differential analysis on. Should be one of phyloseq::rank_names(phyloseq), or "all" means to summarize the taxa by the top taxa ranks (summarize_taxa(ps, level = rank_names(ps)[1])), or "none" means perform differential analysis on the original taxa (taxa_names(phyloseq), e.g., OTU or ASV).
transform	character, the methods used to transform the microbial abundance. See transform_abundances() for more details. The options include:
	<ul> <li>"identity", return the original data without any transformation.</li> <li>"log10", the transformation is log10(object), and if the data contains zeros the transformation is log10(1 + object).</li> </ul>
	<ul> <li>"log10p", the transformation is log10(1 + object).</li> </ul>
norm	the methods used to normalize the microbial abundance data. See normalize() for more details. Options include:
	<ul><li> "none": do not normalize.</li><li> "rarefy": random subsampling counts to the smallest library size in the data set.</li></ul>

	<ul> <li>"TSS": total sum scaling, also referred to as "relative abundance", the abundances were normalized by dividing the corresponding sample library size.</li> <li>"TMM": trimmed mean of m-values. First, a sample is chosen as reference. The scaling factor is then derived using a weighted trimmed mean over the differences of the log-transformed gene-count fold-change between the sample and the reference.</li> <li>"RLE", relative log expression, RLE uses a pseudo-reference calculated using the geometric mean of the gene-specific abundances over all samples.</li> </ul>
	<ul><li>The scaling factors are then calculated as the median of the gene counts ratios between the samples and the reference.</li><li>"CSS": cumulative sum scaling, calculates scaling factors as the cumulative</li></ul>
	sum of gene abundances up to a data-derived threshold.
	• "CLR": centered log-ratio normalization.
	• "CPM": pre-sample normalization of the sum of the values to 1e+06.
norm_para	named list. other arguments passed to specific normalization methods. Most users will not need to pass any additional arguments here.
p_adjust	method for multiple test correction, default none, for more details see stats::p.adjust.
<pre>pvalue_cutoff</pre>	significance level for each of the statistical tests, default 0.05.
W_cutoff	lower bound for the proportion for the W-statistic, default 0.7.

## Details

In an experiment with only two treatments, this tests the following hypothesis for feature *i*:

$$H_{0i}: E(log(\mu_i^1)) = E(log(\mu_i^2))$$

where  $\mu_i^1$  and  $\mu_i^2$  are the mean abundances for feature *i* in the two groups.

The developers of this method recommend the following significance tests if there are 2 groups, use non-parametric Wilcoxon rank sum test stats::wilcox.test(). If there are more than 2 groups, use nonparametric stats::kruskal.test() or one-way ANOVA stats::aov().

#### Value

a microbiomeMarker object, in which the slot of marker\_table contains four variables:

- feature, significantly different features.
- enrich\_group, the class of the differential features enriched.
- effect\_size, differential means for two groups, or F statistic for more than two groups.
- W, the W-statistic, number of features that a single feature is tested to be significantly different against.

#### Author(s)

Huang Lin, Yang Cao

#### run\_ancombc

## References

Mandal et al. "Analysis of composition of microbiomes: a novel method for studying microbial composition", Microbial Ecology in Health & Disease, (2015), 26.

#### Examples

```
data(enterotypes_arumugam)
ps <- phyloseq::subset_samples(
    enterotypes_arumugam,
    Enterotype %in% c("Enterotype 3", "Enterotype 2")
)
run_ancom(ps, group = "Enterotype")</pre>
```

run\_ancombc

*Differential analysis of compositions of microbiomes with bias correction (ANCOM-BC).* 

#### Description

Differential abundance analysis for microbial absolute abundance data. This function is a wrapper of ANCOMBC::ancombc().

## Usage

```
run_ancombc(
  ps,
 group,
 confounders = character(0),
  contrast = NULL,
  taxa_rank = "all",
  transform = c("identity", "log10", "log10p"),
 norm = "none",
 norm_para = list(),
 p_adjust = c("none", "fdr", "bonferroni", "holm", "hochberg", "hommel", "BH", "BY"),
 prv_cut = 0.1,
 lib_cut = 0,
  struc_zero = FALSE,
 neg_{lb} = FALSE,
  tol = 1e-05,
 max_{iter} = 100,
 conserve = FALSE,
  pvalue_cutoff = 0.05
)
```

ps	a phyloseq::phyloseq object, which consists of a feature table, a sample meta- data and a taxonomy table.
group	the name of the group variable in metadata. Specifying group is required for detecting structural zeros and performing global test.
confounders	character vector, the confounding variables to be adjusted. default character(0), indicating no confounding variable.
contrast	this parameter only used for two groups comparison while there are multiple groups. For more please see the following details.
taxa_rank	character to specify taxonomic rank to perform differential analysis on. Should be one of phyloseq::rank_names(phyloseq), or "all" means to summarize the taxa by the top taxa ranks (summarize_taxa(ps, level = rank_names(ps)[1])), or "none" means perform differential analysis on the original taxa (taxa_names(phyloseq), e.g., OTU or ASV).
transform	character, the methods used to transform the microbial abundance. See transform_abundances() for more details. The options include:
	<ul> <li>"identity", return the original data without any transformation (default).</li> <li>"log10", the transformation is log10(object), and if the data contains zeros the transformation is log10(1 + object).</li> </ul>
	<ul> <li>"log10p", the transformation is log10(1 + object).</li> </ul>
norm	the methods used to normalize the microbial abundance data. See normalize() for more details. Options include:
	• "none": do not normalize.
	• "rarefy": random subsampling counts to the smallest library size in the data set.
	• "TSS": total sum scaling, also referred to as "relative abundance", the abun-
	dances were normalized by dividing the corresponding sample library size.
	• "TMM": trimmed mean of m-values. First, a sample is chosen as reference.
	The scaling factor is then derived using a weighted trimmed mean over
	the differences of the log-transformed gene-count fold-change between the sample and the reference.
	• "RLE", relative log expression, RLE uses a pseudo-reference calculated
	using the geometric mean of the gene-specific abundances over all samples.
	The scaling factors are then calculated as the median of the gene counts ratios between the samples and the reference.
	• "CSS": cumulative sum scaling, calculates scaling factors as the cumulative sum of gene abundances up to a data-derived threshold.
	"CLR": centered log-ratio normalization.
	• "CPM": pre-sample normalization of the sum of the values to 1e+06.
norm_para	named list. other arguments passed to specific normalization methods. Most users will not need to pass any additional arguments here.
p_adjust	method to adjust p-values by. Default is "holm". Options include "holm",
	"hochberg", "hommel", "bonferroni", "BH", "BY", "fdr", "none". See stats::p.adjust() for more details.

run\_ancombc

prv_cut	a numerical fraction between 0 and 1. Taxa with prevalences less than $prv_cut$ will be excluded in the analysis. Default is 0.10.
lib_cut	a numerical threshold for filtering samples based on library sizes. Samples with library sizes less than lib_cut will be excluded in the analysis. Default is 0, i.e. do not filter any sample.
struc_zero	whether to detect structural zeros. Default is FALSE.
neg_lb	whether to classify a taxon as a structural zero in the corresponding study group using its asymptotic lower bound. Default is FALSE.
tol	the iteration convergence tolerance for the E-M algorithm. Default is 1e-05.
max_iter	the maximum number of iterations for the E-M algorithm. Default is 100.
conserve	whether to use a conservative variance estimate of the test statistic. It is recom- mended if the sample size is small and/or the number of differentially abundant taxa is believed to be large. Default is FALSE.
<pre>pvalue_cutoff</pre>	level of significance. Default is 0.05.

## Details

contrast must be a two length character or NULL (default). It is only required to set manually for two groups comparison when there are multiple groups. The order determines the direction of comparison, the first element is used to specify the reference group (control). This means that, the first element is the denominator for the fold change, and the second element is used as baseline (numerator for fold change). Otherwise, users do required to concern this parameter (set as default NULL), and if there are two groups, the first level of groups will set as the reference group; if there are multiple groups, it will perform an ANOVA-like testing to find markers which difference in any of the groups.

## Value

a microbiomeMarker object.

## References

Lin, Huang, and Shyamal Das Peddada. "Analysis of compositions of microbiomes with bias correction." Nature communications 11.1 (2020): 1-11.

## See Also

ANCOMBC::ancombc

#### Examples

```
data(enterotypes_arumugam)
ps <- phyloseq::subset_samples(
    enterotypes_arumugam,
    Enterotype %in% c("Enterotype 3", "Enterotype 2")
)
run_ancombc(ps, group = "Enterotype")</pre>
```

run\_deseq2

#### Description

Differential expression analysis based on the Negative Binomial distribution using DESeq2.

# Usage

```
run_deseq2(
 ps,
  group,
  confounders = character(0),
  contrast = NULL,
  taxa_rank = "all",
  norm = "RLE",
 norm_para = list(),
  transform = c("identity", "log10", "log10p"),
  fitType = c("parametric", "local", "mean", "glmGamPoi"),
  sfType = "poscounts",
 betaPrior = FALSE,
 modelMatrixType,
 useT = FALSE,
 minmu = ifelse(fitType == "glmGamPoi", 1e-06, 0.5),
 p_adjust = c("none", "fdr", "bonferroni", "holm", "hochberg", "hommel", "BH", "BY"),
 pvalue_cutoff = 0.05,
  . . .
)
```

ps	a phyloseq::phyloseq object.
group	character, the variable to set the group, must be one of the var of the sample metadata.
confounders	character vector, the confounding variables to be adjusted. default character(0), indicating no confounding variable.
contrast	this parameter only used for two groups comparison while there are multiple groups. For more please see the following details.
taxa_rank	character to specify taxonomic rank to perform differential analysis on. Should be one of phyloseq::rank_names(phyloseq), or "all" means to summarize the taxa by the top taxa ranks (summarize_taxa(ps, level = rank_names(ps)[1])), or "none" means perform differential analysis on the original taxa (taxa_names(phyloseq), e.g., OTU or ASV).
norm	the methods used to normalize the microbial abundance data. See normalize() for more details. Options include:

	• "none": do not normalize.
	• "rarefy": random subsampling counts to the smallest library size in the data set.
	• "TMM": trimmed mean of m-values. First, a sample is chosen as reference. The scaling factor is then derived using a weighted trimmed mean over the differences of the log-transformed gene-count fold-change between the sample and the reference.
	• "RLE", relative log expression, RLE uses a pseudo-reference calculated using the geometric mean of the gene-specific abundances over all samples. The scaling factors are then calculated as the median of the gene counts ratios between the samples and the reference.
	<ul> <li>"CSS": cumulative sum scaling, calculates scaling factors as the cumulative sum of gene abundances up to a data-derived threshold.</li> </ul>
norm_para	arguments passed to specific normalization methods. Most users will not need to pass any additional arguments here.
transform	character, the methods used to transform the microbial abundance. See transform_abundances() for more details. The options include:
	<ul> <li>"identity", return the original data without any transformation (default).</li> <li>"log10", the transformation is log10(object), and if the data contains zeros the transformation is log10(1 + object).</li> <li>"log10p", the transformation is log10(1 + object).</li> </ul>
fitType,sfType	e, betaPrior, modelMatrixType, useT, minmu
	these seven parameters are inherited form DESeq2::DESeq().
	<ul> <li>fitType, either "parametric", "local", "mean", or "glmGamPoi" for the type of fitting of dispersions to the mean intensity.</li> </ul>
	<ul> <li>sfType, either "ratio", "poscounts", or "iterate" for the type of size factor estimation. We recommend to use "poscounts".</li> </ul>
	<ul> <li>betaPrior, whether or not to put a zero-mean normal prior on the non- intercept coefficients.</li> </ul>
	<ul> <li>modelMatrixType, either "standard" or "expanded", which describe how the model matrix,</li> </ul>
	<ul> <li>useT, logical, where Wald statistics are assumed to follow a standard Nor- mal.</li> </ul>
	• minmu, lower bound on the estimated count for fitting gene-wise dispersion.
	For more details, see DESeq2::DESeq(). Most users will not need to set this arguments (just use the defaults).
p_adjust	method for multiple test correction, default none, for more details see stats::p.adjust.
<pre>pvalue_cutoff</pre>	pvalue_cutoff numeric, p value cutoff, default 0.05.
	extra parameters passed to DESeq2::DESeq().

# Details

**Note**: DESeq2 requires the input is raw counts (un-normalized counts), as only the counts values allow assessing the measurement precision correctly. For more details see the vignette of DESeq2 (vignette("DESeq2")).

Thus, this function only supports "none", "rarefy", "RLE", "CSS", and "TMM" normalization methods. We strongly recommend using the "RLE" method (default normalization method in the DE-Seq2 package). The other normalization methods are used for expert users and comparisons among different normalization methods.

For two groups comparison, this function utilizes the Wald test (defined by DESeq2::nbinomWaldTest()) for hypothesis testing. A Wald test statistic is computed along with a probability (p-value) that a test statistic at least as extreme as the observed value were selected at random. contrasts are used to specify which two groups to compare. The order of the names determines the direction of fold change that is reported.

Likelihood ratio test (LRT) is used to identify the genes that significantly changed across all the different levels for multiple groups comparisons. The LRT identified the significant features by comparing the full model to the reduced model. It is testing whether a feature removed in the reduced model explains a significant variation in the data.

contrast must be a two length character or NULL (default). It is only required to set manually for two groups comparison when there are multiple groups. The order determines the direction of comparison, the first element is used to specify the reference group (control). This means that, the first element is the denominator for the fold change, and the second element is used as baseline (numerator for fold change). Otherwise, users do required to concern this parameter (set as default NULL), and if there are two groups, the first level of groups will set as the reference group; if there are multiple groups, it will perform an ANOVA-like testing to find markers which difference in any of the groups.

## Value

a microbiomeMarker object.

# References

Love, Michael I., Wolfgang Huber, and Simon Anders. "Moderated estimation of fold change and dispersion for RNA-seq data with DESeq2." Genome biology 15.12 (2014): 1-21.

#### See Also

DESeq2::results(),DESeq2::DESeq()

#### Examples

```
data(enterotypes_arumugam)
ps <- phyloseq::subset_samples(
    enterotypes_arumugam,
    Enterotype %in% c("Enterotype 3", "Enterotype 2")) %>%
    phyloseq::subset_taxa(Phylum %in% c("Firmicutes"))
run_deseq2(ps, group = "Enterotype")
```

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run\_edger

# Description

Differential expression analysis based on the Negative Binomial distribution using edgeR.

# Usage

```
run_edger(
    ps,
    group,
    confounders = character(0),
    contrast = NULL,
    taxa_rank = "all",
    method = c("LRT", "QLFT"),
    transform = c("identity", "log10", "log10p"),
    norm = "TMM",
    norm_para = list(),
    disp_para = list(),
    p_adjust = c("none", "fdr", "bonferroni", "holm", "hochberg", "hommel", "BH", "BY"),
    pvalue_cutoff = 0.05,
    ....
)
```

ps	ps a phyloseq::phyloseq object.
group	character, the variable to set the group, must be one of the var of the sample metadata.
confounders	character vector, the confounding variables to be adjusted. default character(0), indicating no confounding variable.
contrast	this parameter only used for two groups comparison while there are multiple groups. For more please see the following details.
taxa_rank	character to specify taxonomic rank to perform differential analysis on. Should be one of phyloseq::rank_names(phyloseq), or "all" means to summarize the taxa by the top taxa ranks (summarize_taxa(ps, level = rank_names(ps)[1])), or "none" means perform differential analysis on the original taxa (taxa_names(phyloseq), e.g., OTU or ASV).
method	character, used for differential analysis, please see details below for more info.
transform	character, the methods used to transform the microbial abundance. See transform_abundances() for more details. The options include:
	<ul> <li>"identity", return the original data without any transformation (default).</li> <li>"log10", the transformation is log10(object), and if the data contains zeros the transformation is log10(1 + object).</li> </ul>

	<ul> <li>"log10p", the transformation is log10(1 + object).</li> </ul>
norm	the methods used to normalize the microbial abundance data. See normalize() for more details. Options include:
	• "none": do not normalize.
	• "rarefy": random subsampling counts to the smallest library size in the data set.
	• "TSS": total sum scaling, also referred to as "relative abundance", the abundances were normalized by dividing the corresponding sample library size.
	• "TMM": trimmed mean of m-values. First, a sample is chosen as reference. The scaling factor is then derived using a weighted trimmed mean over the differences of the log-transformed gene-count fold-change between the sample and the reference.
	• "RLE", relative log expression, RLE uses a pseudo-reference calculated using the geometric mean of the gene-specific abundances over all samples. The scaling factors are then calculated as the median of the gene counts ratios between the samples and the reference.
	<ul> <li>"CSS": cumulative sum scaling, calculates scaling factors as the cumulative sum of gene abundances up to a data-derived threshold.</li> </ul>
	"CLR": centered log-ratio normalization.
	• "CPM": pre-sample normalization of the sum of the values to 1e+06.
norm_para	arguments passed to specific normalization methods. Most users will not need to pass any additional arguments here.
disp_para	additional arguments passed to edgeR::estimateDisp() used for dispersions estimation. Most users will not need to pass any additional arguments here.
p_adjust	method for multiple test correction, default none, for more details see stats::p.adjust.
pvalue_cutoff	numeric, p value cutoff, default 0.05
	extra arguments passed to the model. See edgeR::glmQLFit() and edgeR::glmFit() for more details.

#### Details

Note that edgeR is designed to work with actual counts. This means that transformation is not required in any way before inputting them to edgeR.

There are two test methods for differential analysis in **edgeR**, likelihood ratio test (LRT) and quasilikelihood F-test (QLFT). The QLFT method is recommended as it allows stricter error rate control by accounting for the uncertainty in dispersion estimation.

contrast must be a two length character or NULL (default). It is only required to set manually for two groups comparison when there are multiple groups. The order determines the direction of comparison, the first element is used to specify the reference group (control). This means that, the first element is the denominator for the fold change, and the second element is used as baseline (numerator for fold change). Otherwise, users do required to concern this parameter (set as default NULL), and if there are two groups, the first level of groups will set as the reference group; if there are multiple groups, it will perform an ANOVA-like testing to find markers which difference in any of the groups.

# run\_lefse

## Value

a microbiomeMarker object.

## Author(s)

Yang Cao

# References

Robinson, Mark D., and Alicia Oshlack. "A scaling normalization method for differential expression analysis of RNA-seq data." Genome biology 11.3 (2010): 1-9.

Robinson, Mark D., Davis J. McCarthy, and Gordon K. Smyth. "edgeR: a Bioconductor package for differential expression analysis of digital gene expression data." Bioinformatics 26.1 (2010): 139-140.

## See Also

```
edgeR::glmFit(),edgeR::glmQLFit(),edgeR::estimateDisp() ,normalize()
```

#### Examples

```
data(enterotypes_arumugam)
ps <- phyloseq::subset_samples(
    enterotypes_arumugam,
    Enterotype %in% c("Enterotype 3", "Enterotype 2")
)
run_edger(ps, group = "Enterotype")</pre>
```

run\_lefse

Liner discriminant analysis (LDA) effect size (LEFSe) analysis

#### Description

Perform Metagenomic LEFSe analysis based on phyloseq object.

## Usage

```
run_lefse(
    ps,
    group,
    subgroup = NULL,
    taxa_rank = "all",
    transform = c("identity", "log10", "log10p"),
    norm = "CPM",
    norm_para = list(),
    kw_cutoff = 0.05,
    lda_cutoff = 2,
```

```
bootstrap_n = 30,
bootstrap_fraction = 2/3,
wilcoxon_cutoff = 0.05,
multigrp_strat = FALSE,
strict = c("0", "1", "2"),
sample_min = 10,
only_same_subgrp = FALSE,
curv = FALSE
)
```

# Arguments

ps	a phyloseq-class object
group	character, the column name to set the group
subgroup	character, the column name to set the subgroup
taxa_rank	character to specify taxonomic rank to perform differential analysis on. Should be one of phyloseq::rank_names(phyloseq), or "all" means to summarize the taxa by the top taxa ranks (summarize_taxa(ps, level = rank_names(ps)[1])), or "none" means perform differential analysis on the original taxa (taxa_names(phyloseq), e.g., OTU or ASV).
transform	character, the methods used to transform the microbial abundance. See transform_abundances() for more details. The options include:
	<ul> <li>"identity", return the original data without any transformation (default).</li> <li>"log10", the transformation is log10(object), and if the data contains zeros the transformation is log10(1 + object).</li> <li>"log10p", the transformation is log10(1 + object).</li> </ul>
norm	the methods used to normalize the microbial abundance data. See normalize() for more details. Options include:
	<ul> <li>"none": do not normalize.</li> <li>"rarefy": random subsampling counts to the smallest library size in the data set.</li> <li>"TSS": total sum scaling, also referred to as "relative abundance", the abundances were normalized by dividing the corresponding sample library size.</li> <li>"TMM": trimmed mean of m-values. First, a sample is chosen as reference. The scaling factor is then derived using a weighted trimmed mean over the differences of the log-transformed gene-count fold-change between the sample and the reference.</li> <li>"RLE", relative log expression, RLE uses a pseudo-reference calculated using the geometric mean of the gene-specific abundances over all samples. The scaling factors are then calculated as the median of the gene counts ratios between the samples and the reference.</li> <li>"CSS": cumulative sum scaling, calculates scaling factors as the cumulative sum of gene abundances up to a data-derived threshold.</li> <li>"CLR": pre-sample normalization of the sum of the values to 1e+06.</li> </ul>

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# run\_lefse

norm_para	named list. other arguments passed to specific normalization methods. Most users will not need to pass any additional arguments here.	
kw_cutoff	numeric, p value cutoff of kw test, default 0.05	
lda_cutoff	numeric, lda score cutoff, default 2	
bootstrap_n	integer, the number of bootstrap iteration for LDA, default 30	
bootstrap_frac	tion	
	numeric, the subsampling fraction value for each bootstrap iteration, default 2/3	
wilcoxon_cutof	f	
	numeric, p value cutoff of wilcoxon test, default 0.05	
multigrp_strat	logical, for multiple group tasks, whether the test is performed in a one-against one (more strict) or in a one-against all setting, default FALSE.	
strict	multiple testing options, 0 for no correction (default), 1 for independent com- parisons, 2 for independent comparison.	
sample_min	integer, minimum number of samples per subclass for performing wilcoxon test, default 10	
only_same_subgrp		
	logical, whether perform the wilcoxon test only among the subgroups with the same name, default FALSE	
curv	logical, whether perform the wilcoxon test using the Curtis's approach, defalt FALSE	

#### Value

a microbiomeMarker object, in which the slot of marker\_table contains four variables:

- feature, significantly different features.
- enrich\_group, the class of the differential features enriched.
- 1da, logarithmic LDA score (effect size)
- pvalue, p value of kw test.

# Author(s)

Yang Cao

# References

Segata, Nicola, et al. Metagenomic biomarker discovery and explanation. Genome biology 12.6 (2011): R60.

# See Also

normalize

# Examples

```
data(kostic_crc)
kostic_crc_small <- phyloseq::subset_taxa(
    kostic_crc,
    Phylum == "Firmicutes"
)
mm_lefse <- run_lefse(
    kostic_crc_small,
    wilcoxon_cutoff = 0.01,
    group = "DIAGNOSIS",
    kw_cutoff = 0.01,
    multigrp_strat = TRUE,
    lda_cutoff = 4
)</pre>
```

run\_limma\_voom Differential analysis using limma-voom

#### Description

Differential analysis using limma-voom

#### Usage

```
run_limma_voom(
    ps,
    group,
    confounders = character(0),
    contrast = NULL,
    taxa_rank = "all",
    transform = c("identity", "log10", "log10p"),
    norm = "none",
    norm_para = list(),
    voom_span = 0.5,
    p_adjust = c("none", "fdr", "bonferroni", "holm", "hochberg", "hommel", "BH", "BY"),
    pvalue_cutoff = 0.05,
    ...
)
```

#### Arguments

ps	ps a phyloseq::phyloseq object.
group	character, the variable to set the group, must be one of the var of the sample metadata.
confounders	character vector, the confounding variables to be adjusted. default $character(0)$ , indicating no confounding variable.

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contrast	this parameter only used for two groups comparison while there are multiple
contrast	groups. For more please see the following details.
taxa_rank	character to specify taxonomic rank to perform differential analysis on. Should be one of phyloseq::rank_names(phyloseq), or "all" means to summarize the taxa by the top taxa ranks (summarize_taxa(ps, level = rank_names(ps)[1])), or "none" means perform differential analysis on the original taxa (taxa_names(phyloseq), e.g., OTU or ASV).
transform	character, the methods used to transform the microbial abundance. See transform_abundances() for more details. The options include:
	<ul> <li>"identity", return the original data without any transformation (default).</li> <li>"log10", the transformation is log10(object), and if the data contains zeros the transformation is log10(1 + object).</li> <li>"log10p", the transformation is log10(1 + object).</li> </ul>
norm	the methods used to normalize the microbial abundance data. See normalize() for more details. Options include:
	• "none": do not normalize.
	• "rarefy": random subsampling counts to the smallest library size in the data set.
	• "TMM": trimmed mean of m-values. First, a sample is chosen as reference.
	The scaling factor is then derived using a weighted trimmed mean over
	the differences of the log-transformed gene-count fold-change between the sample and the reference.
	• "RLE", relative log expression, RLE uses a pseudo-reference calculated
	using the geometric mean of the gene-specific abundances over all samples.
	The scaling factors are then calculated as the median of the gene counts ratios between the samples and the reference.
	<ul> <li>"CSS": cumulative sum scaling, calculates scaling factors as the cumulative sum of gene abundances up to a data-derived threshold.</li> </ul>
norm_para	arguments passed to specific normalization methods. Most users will not need to pass any additional arguments here.
voom_span	width of the smoothing window used for the lowess mean-variance trend for limma::voom(). Expressed as a proportion between 0 and 1.
p_adjust	method for multiple test correction, default none, for more details see stats::p.adjust.
<pre>pvalue_cutoff</pre>	cutoff of p value, default 0.05.
	extra arguments passed to limma::eBayes().

## Details

contrast must be a two length character or NULL (default). It is only required to set manually for two groups comparison when there are multiple groups. The order determines the direction of comparison, the first element is used to specify the reference group (control). This means that, the first element is the denominator for the fold change, and the second element is used as baseline (numerator for fold change). Otherwise, users do required to concern this parameter (set as default NULL), and if there are two groups, the first level of groups will set as the reference group; if there are multiple groups, it will perform an ANOVA-like testing to find markers which difference in any of the groups.

## Value

a microbiomeMarker object.

#### References

Law, C. W., Chen, Y., Shi, W., & Smyth, G. K. (2014). voom: Precision weights unlock linear model analysis tools for RNA-seq read counts. Genome biology, 15(2), 1-17.

#### Examples

```
data(enterotypes_arumugam)
mm <- run_limma_voom(
    enterotypes_arumugam,
    "Enterotype",
    contrast = c("Enterotype 3", "Enterotype 2"),
    pvalue_cutoff = 0.01,
    p_adjust = "none"
)
mm</pre>
```

run\_marker

## Description

run\_marker is a wrapper of all differential analysis functions.

#### Usage

```
run_marker(
    ps,
    group,
    da_method = c("lefse", "simple_t", "simple_welch", "simple_white", "simple_kruskal",
    "simple_anova", "edger", "deseq2", "metagenomeseq", "ancom", "ancombc", "aldex",
    "limma_voom", "sl_lr", "sl_rf", "sl_svm"),
    taxa_rank = "all",
    transform = c("identity", "log10", "log10p"),
    norm = "none",
    norm_para = list(),
    p_adjust = c("none", "fdr", "bonferroni", "holm", "hochberg", "hommel", "BH", "BY"),
    pvalue_cutoff = 0.05,
    ...
)
```

# run\_marker

ps	a phyloseq::phyloseq object
group	character, the variable to set the group
da_method	<ul> <li>character to specify the differential analysis method. The options include:</li> <li>"lefse", linear discriminant analysis (LDA) effect size (LEfSe) method, for more details see run_lefse().</li> </ul>
	<ul> <li>"simple_t", "simple_welch", "simple_white", "simple_kruskal", and "simple_anova", simple statistic methods; "simple_t", "simple_welch" and "simple_white" for two groups comparison; "simple_kruskal", and "simple_anova" for multiple groups comparison. For more details see run_simple_stat().</li> <li>"edger", see run_edger().</li> <li>"deseq2", see run_deseq2().</li> <li>"metagenomeseq", differential expression analysis based on the Zero-inflated Log-Normal mixture model or Zero-inflated Gaussian mixture model using metagenomeSeq, see run_metagenomeseq().</li> <li>"ancom", see run_ancom().</li> <li>"ancombc", differential analysis of compositions of microbiomes with bias correction, see run_ancombc().</li> <li>"aldex", see run_aldex().</li> <li>"limma_voom", see run_limma_voom().</li> <li>"sl_hr", "sl_rf", and "sl_svm", there supervised leaning (SL) methods: lo-</li> </ul>
	gistic regression (lr), random forest (rf), or support vector machine (svm). For more details see run_sl().
taxa_rank	character to specify taxonomic rank to perform differential analysis on. Should be one of phyloseq::rank_names(phyloseq), or "all" means to summarize the taxa by the top taxa ranks (summarize_taxa(ps, level = rank_names(ps)[1])), or "none" means perform differential analysis on the original taxa (taxa_names(phyloseq), e.g., OTU or ASV).
transform	<ul> <li>character, the methods used to transform the microbial abundance. See transform_abundances() for more details. The options include:</li> <li>"identity", return the original data without any transformation (default).</li> <li>"log10", the transformation is log10(object), and if the data contains zeros the transformation is log10(1 + object).</li> <li>"log10p", the transformation is log10(1 + object).</li> </ul>
norm	the methods used to normalize the microbial abundance data. See normalize() for more details. Options include:
	<ul> <li>"none": do not normalize.</li> <li>"rarefy": random subsampling counts to the smallest library size in the data set.</li> <li>"TSS": total sum scaling, also referred to as "relative abundance", the abundances were normalized by dividing the corresponding sample library size.</li> <li>"TMM": trimmed mean of m-values. First, a sample is chosen as reference. The scaling factor is then derived using a weighted trimmed mean over the differences of the log-transformed gene-count fold-change between the sample and the reference.</li> </ul>

	• "RLE", relative log expression, RLE uses a pseudo-reference calculated using the geometric mean of the gene-specific abundances over all samples. The scaling factors are then calculated as the median of the gene counts ratios between the samples and the reference.
	• "CSS": cumulative sum scaling, calculates scaling factors as the cumulative sum of gene abundances up to a data-derived threshold.
	• "CLR": centered log-ratio normalization.
	• "CPM": pre-sample normalization of the sum of the values to 1e+06.
norm_para	arguments passed to specific normalization methods
p_adjust	method for multiple test correction, default none, for more details see stats::p.adjust.
<pre>pvalue_cutoff</pre>	numeric, p value cutoff, default 0.05.
	extra arguments passed to the corresponding differential analysis functions, e.g. run_lefse().

## Details

This function is only a wrapper of all differential analysis functions, We recommend to use the corresponding function, since it has a better default arguments setting.

#### Value

a microbiomeMarker object.

#### See Also

```
run_lefse(),run_simple_stat(),run_test_two_groups(),run_test_multiple_groups(),run_edger(),run_deseq2
run_metagenomeseq,run_ancom(),run_ancombc(),run_aldex(), run_limma_voom(),run_sl()
```

run\_metagenomeseq metagenomeSeq differential analysis

# Description

Differential expression analysis based on the Zero-inflated Log-Normal mixture model or Zero-inflated Gaussian mixture model using metagenomeSeq.

# Usage

```
run_metagenomeseq(
   ps,
   group,
   confounders = character(0),
   contrast = NULL,
   taxa_rank = "all",
   transform = c("identity", "log10", "log10p"),
   norm = "CSS",
```

```
norm_para = list(),
method = c("ZILN", "ZIG"),
p_adjust = c("none", "fdr", "bonferroni", "holm", "hochberg", "hommel", "BH", "BY"),
pvalue_cutoff = 0.05,
...
```

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ps	ps a phyloseq::phyloseq object.
group	character, the variable to set the group, must be one of the var of the sample metadata.
confounders	character vector, the confounding variables to be adjusted. default character(0), indicating no confounding variable.
contrast	this parameter only used for two groups comparison while there are multiple groups. For more please see the following details.
taxa_rank	character to specify taxonomic rank to perform differential analysis on. Should be one of phyloseq::rank_names(ps), or "all" means to summarize the taxa by the top taxa ranks (summarize_taxa(ps, level = rank_names(ps)[1])), or "none" means perform differential analysis on the original taxa(taxa_names(ps), e.g., OTU or ASV).
transform	character, the methods used to transform the microbial abundance. See transform_abundances() for more details. The options include:
	<ul> <li>"identity", return the original data without any transformation (default).</li> <li>"log10", the transformation is log10(object), and if the data contains zeros the transformation is log10(1 + object).</li> <li>"log10p", the transformation is log10(1 + object).</li> </ul>
norm	the methods used to normalize the microbial abundance data. See normalize() for more details. Options include:
	<ul><li> "none": do not normalize.</li><li> "rarefy": random subsampling counts to the smallest library size in the data</li></ul>
	<ul> <li>set.</li> <li>"TSS": total sum scaling, also referred to as "relative abundance", the abundances were normalized by dividing the corresponding sample library size.</li> <li>"TMM": trimmed mean of m-values. First, a sample is chosen as reference. The scaling factor is then derived using a weighted trimmed mean over the differences of the log-transformed gene-count fold-change between the sample and the reference.</li> <li>"RLE", relative log expression, RLE uses a pseudo-reference calculated using the geometric mean of the gene-specific abundances over all samples. The scaling factors are then calculated as the median of the gene counts ratios between the samples and the reference.</li> <li>"CSS": cumulative sum scaling, calculates scaling factors as the cumulative sum of gene abundances up to a data-derived threshold.</li> <li>"CLR": centered log-ratio normalization.</li> </ul>

	• "CPM": pre-sample normalization of the sum of the values to 1e+06.
norm_para	arguments passed to specific normalization methods.
method	character, which model used for differential analysis, "ZILN" (Zero-inflated Log-Normal mixture model)" or "ZIG" (Zero-inflated Gaussian mixture model). And the zero-inflated log-normal model is preferred due to the high sensitivity and low FDR.
p_adjust	method for multiple test correction, default none, for more details see stats::p.adjust.
<pre>pvalue_cutoff</pre>	numeric, p value cutoff, default 0.05
	<pre>extra arguments passed to the model. more details see metagenomeSeq::fitFeatureModel() and metagenomeSeq::fitZig(), e.g. control (can be setted using metagenomeSeq::zigControl()) for metagenomeSeq::fitZig().</pre>

#### Details

metagnomeSeq provides two differential analysis methods, zero-inflated log-normal mixture model (implemented in metagenomeSeq::fitFeatureModel()) and zero-inflated Gaussian mixture model (implemented in metagenomeSeq::fitZig()). We recommend fitFeatureModel over fitZig due to high sensitivity and low FDR. Both metagenomeSeq::fitFeatureModel() and metagenomeSeq::fitZig() require the abundance profiles before normalization.

For metagenomeSeq::fitZig(), the output column is the coefficient of interest, and logFC column in the output of metagenomeSeq::fitFeatureModel() is analogous to coefficient. Thus, logFC is really just the estimate the coefficient of interest in metagenomeSeq::fitFeatureModel(). For more details see these question Difference between fitFeatureModel and fitZIG in metagenomeSeq.

contrast must be a two length character or NULL (default). It is only required to set manually for two groups comparison when there are multiple groups. The order determines the direction of comparison, the first element is used to specify the reference group (control). This means that, the first element is the denominator for the fold change, and the second element is used as baseline (numerator for fold change). Otherwise, users do required to concern this parameter (set as default NULL), and if there are two groups, the first level of groups will set as the reference group; if there are multiple groups, it will perform an ANOVA-like testing to find markers which difference in any of the groups.

Of note, metagenomeSeq::fitFeatureModel() is not allows for multiple groups comparison.

#### Value

a microbiomeMarker object.

#### Author(s)

Yang Cao

#### References

Paulson, Joseph N., et al. "Differential abundance analysis for microbial marker-gene surveys." Nature methods 10.12 (2013): 1200-1202.

## run\_posthoc\_test

# Examples

```
data(enterotypes_arumugam)
ps <- phyloseq::subset_samples(
    enterotypes_arumugam,
    Enterotype %in% c("Enterotype 3", "Enterotype 2")
)
run_metagenomeseq(ps, group = "Enterotype")</pre>
```

run\_posthoc\_test Post hoc pairwise comparisons for multiple groups test.

## Description

Multiple group test, such as anova and Kruskal-Wallis rank sum test, can be used to uncover the significant feature among all groups. Post hoc tests are used to uncover specific mean differences between pair of groups.

#### Usage

```
run_posthoc_test(
    ps,
    group,
    transform = c("identity", "log10", "log10p"),
    norm = "TSS",
    norm_para = list(),
    conf_level = 0.95,
    method = c("tukey", "games_howell", "scheffe", "welch_uncorrected")
)
```

ps group	a phyloseq::phyloseq object character, the variable to set the group
transform	character, the methods used to transform the microbial abundance. See transform_abundances() for more details. The options include:
	<ul> <li>"identity", return the original data without any transformation (default).</li> <li>"log10", the transformation is log10(object), and if the data contains zeros the transformation is log10(1 + object).</li> <li>"log10p", the transformation is log10(1 + object).</li> </ul>
norm	the methods used to normalize the microbial abundance data. See normalize() for more details. Options include:
	<ul> <li>a integer, e.g. 1e6 (default), indicating pre-sample normalization of the sum of the values to 1e6.</li> <li>"none": do not normalize.</li> </ul>

	• "rarefy": random subsampling counts to the smallest library size in the data set.
	• "TSS": total sum scaling, also referred to as "relative abundance", the abundances were normalized by dividing the corresponding sample library size.
	• "TMM": trimmed mean of m-values. First, a sample is chosen as reference. The scaling factor is then derived using a weighted trimmed mean over the differences of the log-transformed gene-count fold-change between the sample and the reference.
	• "RLE", relative log expression, RLE uses a pseudo-reference calculated using the geometric mean of the gene-specific abundances over all samples. The scaling factors are then calculated as the median of the gene counts ratios between the samples and the reference.
	• "CSS": cumulative sum scaling, calculates scaling factors as the cumulative sum of gene abundances up to a data-derived threshold.
	• "CLR": centered log-ratio normalization.
norm_para	arguments passed to specific normalization methods
conf_level	confidence level, default 0.95
method	one of "tukey", "games_howell", "scheffe", "welch_uncorrected", defining the method for the pairwise comparisons. See details for more information.

# Value

a postHocTest object

# See Also

postHocTest, run\_test\_multiple\_groups()

# Examples

```
data(enterotypes_arumugam)
ps <- phyloseq::subset_samples(
    enterotypes_arumugam,
    Enterotype %in% c("Enterotype 3", "Enterotype 2", "Enterotype 1")
) %>%
    phyloseq::subset_taxa(Phylum == "Bacteroidetes")
pht <- run_posthoc_test(ps, group = "Enterotype")
pht</pre>
```

run\_simple\_stat Simple statistical analysis of metagenomic profiles

## Description

Perform simple statistical analysis of metagenomic profiles. This function is a wrapper of run\_test\_two\_groups and run\_test\_multiple\_groups.

# run\_simple\_stat

# Usage

```
run_simple_stat(
 ps,
 group,
  taxa_rank = "all",
  transform = c("identity", "log10", "log10p"),
 norm = "TSS",
 norm_para = list(),
 method = c("welch.test", "t.test", "white.test", "anova", "kruskal"),
 p_adjust = c("none", "fdr", "bonferroni", "holm", "hochberg", "hommel", "BH", "BY"),
 pvalue_cutoff = 0.05,
 diff_mean_cutoff = NULL,
 ratio_cutoff = NULL,
 eta_squared_cutoff = NULL,
  conf_level = 0.95,
 nperm = 1000,
  . . .
)
```

ps	a phyloseq::phyloseq object
group	character, the variable to set the group
taxa_rank	character to specify taxonomic rank to perform differential analysis on. Should be one of phyloseq::rank_names(phyloseq), or "all" means to summarize the taxa by the top taxa ranks (summarize_taxa(ps, level = rank_names(ps)[1])), or "none" means perform differential analysis on the original taxa (taxa_names(phyloseq), e.g., OTU or ASV).
transform	character, the methods used to transform the microbial abundance. See transform_abundances() for more details. The options include:
	<ul> <li>"identity", return the original data without any transformation (default).</li> <li>"log10", the transformation is log10(object), and if the data contains zeros the transformation is log10(1 + object).</li> <li>"log10p", the transformation is log10(1 + object).</li> </ul>
norm	the methods used to normalize the microbial abundance data. See normalize() for more details. Options include:
	<ul> <li>"none": do not normalize.</li> <li>"rarefy": random subsampling counts to the smallest library size in the data set.</li> <li>"TSS": total sum scaling, also referred to as "relative abundance", the abundances were normalized by dividing the corresponding sample library size.</li> <li>"TMM": trimmed mean of m-values. First, a sample is chosen as reference. The scaling factor is then derived using a weighted trimmed mean over the differences of the log-transformed gene-count fold-change between the sample and the reference.</li> </ul>

	<ul> <li>"RLE", relative log expression, RLE uses a pseudo-reference calculated using the geometric mean of the gene-specific abundances over all samples. The scaling factors are then calculated as the median of the gene counts ratios between the samples and the reference.</li> <li>"CSS": cumulative sum scaling, calculates scaling factors as the cumulative sum of gene abundances up to a data-derived threshold.</li> <li>"CLR": centered log-ratio normalization.</li> <li>"CPM": pre-sample normalization of the sum of the values to 1e+06.</li> </ul>
norm_para	arguments passed to specific normalization methods
method	test method, options include: "welch.test", "t.test" and "white.test" for two groups comparison, "anova"and "kruskal" for multiple groups comparison.
p_adjust	method for multiple test correction, default none, for more details see stats::p.adjust.
<pre>pvalue_cutoff</pre>	numeric, p value cutoff, default 0.05
diff_mean_cuto	ff, ratio_cutoff
	only used for two groups comparison, cutoff of different means and ratios, de- fault NULL which means no effect size filter.
eta_squared_cu	toff
	only used for multiple groups comparison, numeric, cutoff of effect size (eta squared) default NULL which means no effect size filter.
conf_level	only used for two groups comparison, numeric, confidence level of interval.
nperm	integer, only used for two groups comparison, number of permutations for white non parametric t test estimation
	only used for two groups comparison, extra arguments passed to t.test() or fisher.test().

# Value

a microbiomeMarker object.

# See Also

run\_test\_two\_groups(),run\_test\_multiple\_groups()

# Examples

```
data(enterotypes_arumugam)
ps <- phyloseq::subset_samples(
    enterotypes_arumugam,
    Enterotype %in% c("Enterotype 3", "Enterotype 2")
)
run_simple_stat(ps, group = "Enterotype")</pre>
```

run\_sl

# Description

Identify biomarkers using logistic regression, random forest, or support vector machine.

# Usage

```
run_sl(
    ps,
    group,
    taxa_rank = "all",
    transform = c("identity", "log10", "log10p"),
    norm = "none",
    norm_para = list(),
    nfolds = 3,
    nrepeats = 3,
    sampling = NULL,
    tune_length = 5,
    top_n = 10,
    method = c("LR", "RF", "SVM"),
    ...
)
```

ps	a phyloseq-class object.
group	character, the variable to set the group.
taxa_rank	character to specify taxonomic rank to perform differential analysis on. Should be one of phyloseq::rank_names(phyloseq), or "all" means to summarize the taxa by the top taxa ranks (summarize_taxa(ps, level = rank_names(ps)[1])), or "none" means perform differential analysis on the original taxa (taxa_names(phyloseq), e.g., OTU or ASV).
transform	character, the methods used to transform the microbial abundance. See transform_abundances() for more details. The options include:
	<ul> <li>"identity", return the original data without any transformation (default).</li> <li>"log10", the transformation is log10(object), and if the data contains zeros the transformation is log10(1 + object).</li> <li>"log10p", the transformation is log10(1 + object).</li> </ul>
norm	the methods used to normalize the microbial abundance data. See normalize() for more details. Options include:
	<ul><li> "none": do not normalize.</li><li> "rarefy": random subsampling counts to the smallest library size in the data set.</li></ul>

	<ul> <li>"TSS": total sum scaling, also referred to as "relative abundance", the abundances were normalized by dividing the corresponding sample library size.</li> <li>"TMM": trimmed mean of m-values. First, a sample is chosen as reference. The scaling factor is then derived using a weighted trimmed mean over the differences of the log-transformed gene-count fold-change between the sample and the reference.</li> <li>"RLE", relative log expression, RLE uses a pseudo-reference calculated using the geometric mean of the gene-specific abundances over all samples. The scaling factors are then calculated as the median of the gene counts ratios between the samples and the reference.</li> <li>"CSS": cumulative sum scaling, calculates scaling factors as the cumulative sum of gene abundances up to a data-derived threshold.</li> <li>"CLR": centered log-ratio normalization.</li> <li>"CPM": pre-sample normalization of the sum of the values to 1e+06.</li> </ul>
norm_para	named list. other arguments passed to specific normalization methods. Most users will not need to pass any additional arguments here.
nfolds	the number of splits in CV.
nrepeats	the number of complete sets of folds to compute.
sampling	a single character value describing the type of additional sampling that is con- ducted after resampling (usually to resolve class imbalances). Values are "none", "down", "up", "smote", or "rose". For more details see caret::trainControl().
tune_length	an integer denoting the amount of granularity in the tuning parameter grid. For more details see caret::train().
top_n	an integer denoting the top n features as the biomarker according the importance score.
method	supervised learning method, options are "LR" (logistic regression), "RF" (rando forest), or "SVM" (support vector machine).
	$extra \ arguments \ passed \ to \ the \ classification. \ e.g., \ importance \ for \ random \ Forest:: random \ Forest.$

# Details

Only support two groups comparison in the current version. And the marker was selected based on its importance score. Moreover, The hyper-parameters are selected automatically by a grid-search based method in the N-time K-fold cross-validation. Thus, the identified biomarker based can be biased due to model overfitting for small datasets (e.g., with less than 100 samples).

The argument top\_n is used to denote the number of markers based on the importance score. There is no rule or principle on how to select top\_n, however, usually it is very useful to try a different top\_n and compare the performance of the marker predictions for the testing data.

# Value

a microbiomeMarker object.

## Author(s)

Yang Cao

## See Also

```
caret::train(),caret::trainControl()
```

#### Examples

```
data(enterotypes_arumugam)
# small example phyloseq object for test
ps_small <- phyloseq::subset_taxa(</pre>
    enterotypes_arumugam,
    Phylum %in% c("Firmicutes", "Bacteroidetes")
)
set.seed(2021)
mm <- run_sl(</pre>
    ps_small,
    group = "Gender",
    taxa_rank = "Genus",
    nfolds = 2,
    nrepeats = 1,
    top_n = 15,
    norm = "TSS",
    method = "LR",
)
mm
```

```
run_test_multiple_groups
```

Statistical test for multiple groups

# Description

Statistical test for multiple groups

#### Usage

```
run_test_multiple_groups(
    ps,
    group,
    taxa_rank = "all",
    transform = c("identity", "log10", "log10p"),
    norm = "TSS",
    norm_para = list(),
    method = c("anova", "kruskal"),
    p_adjust = c("none", "fdr", "bonferroni", "holm", "hochberg", "hommel", "BH", "BY"),
    pvalue_cutoff = 0.05,
    effect_size_cutoff = NULL
)
```

# Arguments

ps	a phyloseq::phyloseq object
group	character, the variable to set the group
taxa_rank	character to specify taxonomic rank to perform differential analysis on. Should be one of phyloseq::rank_names(phyloseq), or "all" means to summarize the taxa by the top taxa ranks (summarize_taxa(ps, level = rank_names(ps)[1])), or "none" means perform differential analysis on the original taxa (taxa_names(phyloseq), e.g., OTU or ASV).
transform	character, the methods used to transform the microbial abundance. See transform_abundances() for more details. The options include:
	<ul> <li>"identity", return the original data without any transformation (default).</li> <li>"log10", the transformation is log10(object), and if the data contains zeros the transformation is log10(1 + object).</li> <li>"log10p", the transformation is log10(1 + object).</li> </ul>
norm	the methods used to normalize the microbial abundance data. See normalize() for more details. Options include:
	<ul> <li>"none": do not normalize.</li> <li>"rarefy": random subsampling counts to the smallest library size in the data set.</li> <li>"TSS": total sum scaling, also referred to as "relative abundance", the abundances were normalized by dividing the corresponding sample library size.</li> <li>"TMM": trimmed mean of m-values. First, a sample is chosen as reference. The scaling factor is then derived using a weighted trimmed mean over the differences of the log-transformed gene-count fold-change between the sample and the reference.</li> <li>"RLE", relative log expression, RLE uses a pseudo-reference calculated using the geometric mean of the gene-specific abundances over all samples. The scaling factors are then calculated as the median of the gene counts ratios between the samples and the reference.</li> <li>"CSS": cumulative sum scaling, calculates scaling factors as the cumulative sum of gene abundances up to a data-derived threshold.</li> <li>"CLR": centered log-ratio normalization.</li> <li>"CPM": pre-sample normalization of the sum of the values to 1e+06.</li> </ul>
norm_para	arguments passed to specific normalization methods
method	test method, must be one of "anova" or "kruskal"
p_adjust	method for multiple test correction, default none, for more details see stats::p.adjust.
<pre>pvalue_cutoff</pre>	numeric, p value cutoff, default 0.05.
effect_size_cut	
	numeric, cutoff of effect size default NULL which means no effect size filter. The eta squared is used to measure the effect size for anova/kruskal test.

# Value

a microbiomeMarker object.

run\_test\_two\_groups

## See Also

run\_posthoc\_test(),run\_test\_two\_groups(),run\_simple\_stat()

#### Examples

```
data(enterotypes_arumugam)
ps <- phyloseq::subset_samples(
    enterotypes_arumugam,
    Enterotype %in% c("Enterotype 3", "Enterotype 2", "Enterotype 1")
)
mm_anova <- run_test_multiple_groups(
    ps,
    group = "Enterotype",
    method = "anova"
)</pre>
```

run\_test\_two\_groups Statistical test between two groups

#### Description

Statistical test between two groups

## Usage

```
run_test_two_groups(
    ps,
    group,
    taxa_rank = "all",
    transform = c("identity", "log10", "log10p"),
    norm = "TSS",
    norm_para = list(),
    method = c("welch.test", "t.test", "white.test"),
    p_adjust = c("none", "fdr", "bonferroni", "holm", "hochberg", "hommel", "BH", "BY"),
    pvalue_cutoff = 0.05,
    diff_mean_cutoff = NULL,
    ratio_cutoff = NULL,
    conf_level = 0.95,
    nperm = 1000,
    ...
```

```
)
```

ps	a phyloseq::phyloseq object
group	character, the variable to set the group

taxa_rank	character to specify taxonomic rank to perform differential analysis on. Should be one of phyloseq::rank_names(phyloseq), or "all" means to summarize the taxa by the top taxa ranks (summarize_taxa(ps, level = rank_names(ps)[1])), or "none" means perform differential analysis on the original taxa (taxa_names(phyloseq), e.g., OTU or ASV).
transform	character, the methods used to transform the microbial abundance. See transform_abundances() for more details. The options include:
	• "identity", return the original data without any transformation (default).
	<ul> <li>"log10", the transformation is log10(object), and if the data contains zeros the transformation is log10(1 + object).</li> </ul>
	<ul> <li>"log10p", the transformation is log10(1 + object).</li> </ul>
norm	the methods used to normalize the microbial abundance data. See normalize() for more details. Options include:
	• "none": do not normalize.
	• "rarefy": random subsampling counts to the smallest library size in the data set.
	• "TSS": total sum scaling, also referred to as "relative abundance", the abun-
	dances were normalized by dividing the corresponding sample library size.
	• "TMM": trimmed mean of m-values. First, a sample is chosen as reference.
	The scaling factor is then derived using a weighted trimmed mean over the differences of the last transformed game event field shares between the
	the differences of the log-transformed gene-count fold-change between the sample and the reference.
	• "RLE", relative log expression, RLE uses a pseudo-reference calculated
	using the geometric mean of the gene-specific abundances over all samples.
	The scaling factors are then calculated as the median of the gene counts ratios between the samples and the reference.
	<ul> <li>"CSS": cumulative sum scaling, calculates scaling factors as the cumulative sum of gene abundances up to a data-derived threshold.</li> </ul>
	"CLR": centered log-ratio normalization.
	• "CPM": pre-sample normalization of the sum of the values to 1e+06.
norm_para	arguments passed to specific normalization methods
method	test method, must be one of "welch.test", "t.test" or "white.test"
p_adjust	method for multiple test correction, default none, for more details see stats::p.adjust.
pvalue_cutoff	numeric, p value cutoff, default 0.05
diff_mean_cutor	ff,ratio_cutoff
	cutoff of different means and ratios, default NULL which means no effect size filter.
conf_level	numeric, confidence level of interval.
nperm	integer, number of permutations for white non parametric t test estimation
	extra arguments passed to t.test() or fisher.test()

# Value

a microbiomeMarker object.

# subset\_marker

#### Author(s)

Yang Cao

## See Also

run\_test\_multiple\_groups(),run\_simple\_stat

#### Examples

```
data(enterotypes_arumugam)
mm_welch <- run_test_two_groups(
    enterotypes_arumugam,
    group = "Gender",
    method = "welch.test"
)
mm_welch</pre>
```

subset\_marker Subset microbiome markers

# Description

Subset markers based on an expression related to the columns and values within the marker\_table slot of mm.

#### Usage

```
subset_marker(mm, ...)
```

#### Arguments

mm	a microbiomeMarker or marker_table object.
	the subsetting expression passed to $\verb+base::subset()$ .

# Value

a subset object in the same class with mm.

#### Examples

```
data(enterotypes_arumugam)
mm <- run_limma_voom(
    enterotypes_arumugam,
    "Enterotype",
    contrast = c("Enterotype 3", "Enterotype 2"),
    pvalue_cutoff = 0.01,
    p_adjust = "none"
)
subset_marker(mm, pvalue < 0.005)</pre>
```

summarize\_taxa

#### Description

Provides summary information of the representation of a taxonomic levels within each sample.

#### Usage

summarize\_taxa(ps, level = rank\_names(ps)[1], absolute = TRUE, sep = "|")

#### Arguments

ps	a phyloseq-class object.
level	taxonomic level to summarize, default the top level rank of the ps.
absolute	logical, whether return the absolute abundance or relative abundance, default FALSE.
sep	a character string to separate the taxonomic levels.

#### Value

a phyloseq::phyloseq object, where each row represents a taxa, and each col represents the taxa abundance of each sample.

# Examples

```
data(enterotypes_arumugam)
summarize_taxa(enterotypes_arumugam)
```

summary.compareDA Summary differential analysis methods comparison results

#### Description

Summary differential analysis methods comparison results

#### Usage

```
## S3 method for class 'compareDA'
summary(
   object,
   sort = c("score", "auc", "fpr", "power"),
   boot = TRUE,
   boot_n = 1000L,
   prob = c(0.05, 0.95),
   ...
)
```

#### Arguments

object	an compareDA object, output from compare_DA().
sort	character string specifying sort method. Possibilities are "score" which is calculated as $(auc - 0.5) * power - fdr$ , "auc" for area under the ROC curve, "fpr" for false positive rate, "power" for empirical power.
boot	logical, whether use bootstrap for confidence limites of the score, default TRUE. Recommended to be TRUE unless n_rep is larger then 100 in compare_DA().
boot_n	integer, number of bootstraps, default 1000L.
prob	two length numeric vector, confidence limits for score, default c(0.05, 0.95).
	extra arguments affecting the summary produced.

# Value

a data.frame containing measurements for differential analysis methods:

- call: differential analysis commands.
- auc: area under curve of ROC.
- fpr: false positive rate
- power: empirical power.
- fdr: false discover7y rate.
- score: score which is calculated as (auc 0.5) \* power f dr.
- score\_\*: confidence limits of score.

transform\_abundances Transform the taxa abundances in otu\_table sample by sample

# Description

Transform the taxa abundances in otu\_table sample by sample, which means the counts of each sample will be transformed individually.

# Usage

```
transform_abundances(object, transform = c("identity", "log10", "log10p"))
```

# Arguments

object	<pre>otu_table, phyloseq, or microbiomeMarker.</pre>
transform	transformation to apply, the options include:
	• "identity", return the original data without any transformation.
	• "log10", the transformation is log10(object), and if the data contains ze-
	ros the transformation is $log10(1 + object)$ .

• "log10p", the transformation is log10(1 + object).

# Value

A object matches the class of argument object with the transformed otu\_table.

## See Also

abundances()

# Examples

```
data(oxygen)
x1 <- transform_abundances(oxygen)
head(otu_table(x1), 10)
x2 <- transform_abundances(oxygen, "log10")
head(otu_table(x2), 10)
x3 <- transform_abundances(oxygen, "log10p")
head(otu_table(x3), 10)</pre>
```

Ε

#### Extract marker\_table object

## Description

Operators acting on marker\_table to extract parts.

## Usage

## S4 method for signature 'marker\_table,ANY,ANY,ANY'
x[i, j, ..., drop = TRUE]

# Arguments

Х	a marker_table object.
i,j	elements to extract.
	<pre>see base::Extract().</pre>
drop	ignored now.

# Value

a marker\_table object.

# See Also

base::Extract()

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