Package ‘benchdamic’

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Type Package

Title Benchmark of differential abundance methods on microbiome data

Version 1.5.2

Description Starting from a microbiome dataset (16S or WMS with absolute count values) it is possible to perform several analysis to assess the performances of many differential abundance detection methods. A basic and standardized version of the main differential abundance analysis methods is supplied but the user can also add his method to the benchmark. The analyses focus on 4 main aspects: i) the goodness of fit of each method’s distributional assumptions on the observed count data, ii) the ability to control the false discovery rate, iii) the within and between method concordances, iv) the truthfulness of the findings if any apriori knowledge is given. Several graphical functions are available for result visualization.

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addKnowledge

Description

Add a priori knowledge for each feature tested by a method.

Usage

addKnowledge(method, priorKnowledge, enrichmentCol, namesCol = NULL)

Arguments

- **method**: Output of differential abundance detection method in which DA information is extracted by the getDA function.
- **priorKnowledge**: data.frame (with feature names as row.names) containing feature level meta-data.
- **enrichmentCol**: name of the column containing information for enrichment analysis.
- **namesCol**: name of the column containing new names for features (default namesCol = NULL).

Value

A data.frame with a new column containing information for enrichment analysis.

See Also

createEnrichment.

Examples

data("ps_plaque_16S")
data("microbial_metabolism")

# Extract genera from the phyloseq tax_table slot
genera <- phyloseq::tax_table(ps_plaque_16S)[, "GENUS"]
# Genera as rownames of microbial_metabolism data.frame
rownames(microbial_metabolism) <- microbial_metabolism$Genus
# Match OTUs to their metabolism
priorInfo <- data.frame(genera,
    "Type" = microbial_metabolism[genera, "Type"])
# Unmatched genera becomes "Unknown"
unknown_metabolism <- is.na(priorInfo$Type)
priorInfo[unknown_metabolism, "Type"] <- "Unknown"
priorInfo$Type <- factor(priorInfo$Type)
# Add a more informative names column
priorInfo[, "newNames"] <- paste0(rownames(priorInfo), priorInfo[, "GENUS"])

# DA Analysis

# Make sure the subject ID variable is a factor
phyloseq::sample_data(ps_plaque_16S)[, "RSID"] <- as.factor(
  phyloseq::sample_data(ps_plaque_16S)[["RSID"]])

# Add scaling factors
ps_plaque_16S <- norm_edgeR(object = ps_plaque_16S, method = "TMM")

# DA analysis
da.limma <- DA_limma(
  object = ps_plaque_16S,
  design = ~ 1 + RSID + HMP_BODY_SUBSITE,
  coef = "HMP_BODY_SUBSITESupragingival Plaque",
  norm = "TMM"
)

DA <- getDA(method = da.limma, slot = "pValMat", colName = "adjP",
            type = "pvalue", direction = "logFC", threshold_pvalue = 0.05,
            threshold_logfc = 1, top = NULL)

# Add a priori information
DA_info <- addKnowledge(method = DA, priorKnowledge = priorInfo,
                          enrichmentCol = "Type", namesCol = "newNames")

---

**Description**

Compute the area between the bisector and the concordance curve.

**Usage**

areaCAT(concordance, plotIt = FALSE)

**Arguments**

- **concordance** A long format data.frame produced by `createConcordance` function.
- **plotIt** Plot the concordance (default `plotIt = FALSE`).

**Value**

A long format data.frame object with several columns:

- comparison which indicates the comparison number;
- n_features which indicates the total number of taxa in the comparison dataset;
- method1 which contains the first method name;
- method2 which contains the first method name;
• rank;
• concordance which is defined as the cardinality of the intersection of the top rank elements of each list, divided by rank, i.e., \( \frac{L_{\text{1:rank}} \cap M_{\text{1:rank}}}{\text{rank}} \), where L and M represent the lists of the extracted statistics of method1 and method2 respectively;
• heightOver which is the distance between the bisector and the concordance value;
• areaOver which is the cumulative sum of the heightOver value.

See Also

`createConcordance` and `plotConcordance`

Examples

data(ps_plaque_16S)

# Balanced design for dependent samples
my_splits <- createSplits(
  object = ps_plaque_16S, varName = "HMP_BODY_SUBSITE",
  balanced = TRUE, paired = "RSID", N = 10 # N = 100 suggested
)

# Make sure the subject ID variable is a factor
phyloseq::sample_data(ps_plaque_16S)[, "RSID"] <- as.factor(
  phyloseq::sample_data(ps_plaque_16S)[["RSID"]]
)

# Initialize some limma based methods
my_limma <- set_limma(design = ~ RSID + HMP_BODY_SUBSITE,
  coef = "HMP_BODY_SUBSITE\text{Supragingival Plaque}",
  norm = c("TMM", "CSS"))

# Set the normalization methods according to the DA methods
my_norm <- setNormalizations(fun = c("norm_edgeR", "norm_CSS"),
  method = c("TMM", "CSS"))

# Run methods on split datasets
results <- runSplits(split_list = my_splits, method_list = my_limma,
  normalization_list = my_norm, object = ps_plaque_16S)

# Concordance for p-values
concordance_pvalues <- createConcordance(
  object = results, slot = "pValMat", colName = "rawP", type = "pvalue"
)

# Add area over the concordance curve
concordance_area <- areaCAT(concordance = concordance_pvalues)
Description

For the i top-ranked members of each list, concordance is defined as \( \frac{\text{length}(\text{intersect}(\text{vec1}[1:i], \text{vec2}[1:i])))}{i} \).

Usage

\[
\text{CAT}(\text{vec1}, \text{vec2}, \text{maxrank} = \text{min}(\text{length}(\text{vec1}), \text{length}(\text{vec2})))
\]

Arguments

- vec1, vec2: Two numeric vectors, for computing concordance. If these are numeric vectors with names, the numeric values will be used for sorting and the names will be used for calculating concordance. Otherwise, they are assumed to be already-ranked vectors, and the values themselves will be used for calculating concordance.
- maxrank: Optionally specify the maximum size of top-ranked items that you want to plot.

Value

A data.frame with two columns: rank containing the length of the top lists and concordance which is the fraction in common that the two provided lists have in the top rank items.

See Also

\text{createConcordance}.

Examples

\[
\text{vec1} \leftarrow \text{c("A" = 10, "B" = 5, "C" = 20, "D" = 15)}
\]
\[
\text{vec2} \leftarrow \text{c("A" = 1, "B" = 2, "C" = 3, "D" = 4)}
\]
\[
\text{CAT}(\text{vec1}, \text{vec2})
\]

Description

Check if the normalization function’s name and the method’s name to compute normalization/scaling factors are correctly matched.

Usage

\[
\text{checkNormalization}(\text{fun}, \text{method, ...})
\]
Arguments

fun       a character with the name of normalization function (e.g. "norm_edgeR", "norm_DESeq2", "norm_CSS"...).
method    a character with the normalization method (e.g. "TMM", "upperquartile"... if the fun is "norm_edgeR").
...       other arguments if needed (e.g. for norm_edgeR normalizations).

Value

a list object containing the normalization method and its parameters.

See Also

setNormalizations, norm_edgeR, norm_DESeq2, norm_CSS, norm_TSS

Examples

# Check if TMM normalization belong to "norm_edgeR"
check_TMM_normalization <- checkNormalization(fun = "norm_edgeR",
                                               method = "TMM")

createColors

Description

Produce a qualitative set of colors.

Usage

createColors(variable)

Arguments

variable       character vector or factor variable.

Value

A named vector containing the color codes.

Examples

# Given qualitative variable
cond <- factor(c("A", "A", "B", "B", "C", "D"),
               levels = c("A", "B", "C", "D"))

# Associate a color to each level (or unique value, if not a factor)
cond_colors <- createColors(cond)
createConcordance

Description

Compute the between and within method concordances comparing the lists of extracted statistics from the outputs of the differential abundance detection methods.

Usage

```r
createConcordance(object, slot = "pValMat", colName = "rawP", type = "pvalue")
```

Arguments

- **object**: Output of differential abundance detection methods. `pValMat`, `statInfo` matrices, and method's name must be present (See vignette for detailed information).
- **slot**: A character vector with 1 or number-of-methods-times repeats of the slot names where to extract values for each method (default `slot = "pValMat"`).
- **colName**: A character vector with 1 or number-of-methods-times repeats of the column name of the slot where to extract values for each method (default `colName = "rawP"`).
- **type**: A character vector with 1 or number-of-methods-times repeats of the value type of the column selected where to extract values for each method. Two values are possible: "pvalue" or "logfc" (default `type = "pvalue"`).

Value

A long format `data.frame` object with several columns:

- **comparison** which indicates the comparison number;
- **n_features** which indicates the total number of taxa in the comparison dataset;
- **method1** which contains the first method name;
- **method2** which contains the first method name;
- **rank**;
- **concordance** which is defined as the cardinality of the intersection of the top rank elements of each list, divided by rank, i.e., \((L_{1:rank} \cap M_{1:rank})/rank\), where L and M represent the lists of the extracted statistics of method1 and method2 respectively (averaged values between subset1 and subset2).

See Also

`extractStatistics` and `areaCAT`. 
Examples

    data(ps_plaque_16S)

    # Balanced design
    my_splits <- createSplits(
        object = ps_plaque_16S, varName = "HMP_BODY_SUBSITE", balanced = TRUE,
        paired = "RSID", N = 10 # N = 100 suggested
    )

    # Make sure the subject ID variable is a factor
    phyloseq::sample_data(ps_plaque_16S)[, "RSID"] <- as.factor(
        phyloseq::sample_data(ps_plaque_16S)[["RSID"]]
    )

    # Initialize some limma based methods
    my_limma <- set_limma(design = ~ RSID + HMP_BODY_SUBSITE,
        coef = "HMP_BODY_SUBSITESupragingival Plaque",
        norm = c("TMM", "CSS")
    )

    # Set the normalization methods according to the DA methods
    my_norm <- setNormalizations(fun = c("norm_edgeR", "norm_CSS"),
        method = c("TMM", "CSS")
    )

    # Run methods on split datasets
    results <- runSplits(split_list = my_splits, method_list = my_limma,
        normalization_list = my_norm, object = ps_plaque_16S)

    # Concordance for p-values
    concordance_pvalues <- createConcordance(
        object = results, slot = "pValMat", colName = "rawP", type = "pvalue"
    )

    # Concordance for log fold changes
    concordance_logfc <- createConcordance(
        object = results, slot = "statInfo", colName = "logFC", type = "logfc"
    )

    # Concordance for log fold changes in the first method and p-values in the
    # other
    concordance_logfc_pvalues <- createConcordance(
        object = results, slot = c("statInfo", "pValMat"),
        colName = c("logFC", "rawP"), type = c("logfc", "pvalue")
    )

Description

Create a data.frame object with several information to perform enrichment analysis.
Usage

```r
createEnrichment(
  object,
  priorKnowledge,
  enrichmentCol,
  namesCol = NULL,
  slot = "pValMat",
  colName = "adjP",
  type = "pvalue",
  direction = NULL,
  threshold_pvalue = 1,
  threshold_logfc = 0,
  top = NULL,
  alternative = "greater",
  verbose = FALSE
)
```

Arguments

- **object**: Output of differential abundance detection methods. `pValMat`, `statInfo` matrices, and method’s name must be present (See vignette for detailed information).
- **priorKnowledge**: data.frame (with feature names as row.names) containing feature level metadata.
- **enrichmentCol**: name of the column containing information for enrichment analysis.
- **namesCol**: name of the column containing new names for features (default `namesCol = NULL`).
- **slot**: A character vector with 1 or number-of-methods-times repeats of the slot names where to extract values for each method (default `slot = "pValMat"`).
- **colName**: A character vector with 1 or number-of-methods-times repeats of the column name of the slot where to extract values for each method (default `colName = "rawP"`).
- **type**: A character vector with 1 or number-of-methods-times repeats of the value type of the column selected where to extract values for each method. Two values are possible: "pvalue" or "logfc" (default `type = "pvalue"`).
- **direction**: A character vector with 1 or number-of-methods-times repeats of the `statInfo`'s column name containing information about the signs of differential abundance (usually log fold changes) for each method (default `direction = NULL`).
- **threshold_pvalue**: A single or a numeric vector of thresholds for p-values. If present, features with p-values lower than `threshold_pvalue` are considered differentially abundant. Set `threshold_pvalue = 1` to not filter by p-values.
- **threshold_logfc**: A single or a numeric vector of thresholds for log fold changes. If present, features with log fold change absolute values higher than `threshold_logfc` are considered differentially abundant. Set `threshold_logfc = 0` to not filter by log fold change values.
createEnrichment

top
If not null, the top number of features, ordered by p-values or log fold change values, are considered as differentially abundant (default top = NULL).

alternative
indicates the alternative hypothesis and must be one of "two.sided", "greater" or "less". You can specify just the initial letter. Only used in the 2 x 2 case.

verbose
Boolean to display the kind of extracted values (default verbose = FALSE).

Value
a list of objects for each method. Each list contains:

- data a data.frame object with DA directions, statistics, and feature names;
- tables a list of 2x2 contingency tables;
- tests the list of Fisher exact tests' p-values for each contingency table;
- summaries a list with the first element of each contingency table and its p-value (for graphical purposes);

See Also
addKnowledge, extractDA, and enrichmentTest.

Examples

data("ps_plaque_16S")
data("microbial_metabolism")

# Extract genera from the phyloseq tax_table slot
genera <- phyloseq::tax_table(ps_plaque_16S)[, "GENUS"]
# Genera as rownames of microbial_metabolism data.frame
rownames(microbial_metabolism) <- microbial_metabolism$Genus
# Match OTUs to their metabolism
priorInfo <- data.frame(genera,
    "Type" = microbial_metabolism[genera, "Type"])
# Unmatched genera becomes "Unknown"
unknown_metabolism <- is.na(priorInfo$Type)
priorInfo[unknown_metabolism, "Type"] <- "Unknown"
priorInfo$Type <- factor(priorInfo$Type)
# Add a more informative names column
priorInfo[, "newNames"] <- paste0(rownames(priorInfo), priorInfo[, "GENUS"])

# Add some normalization/scaling factors to the phyloseq object
my_norm <- setNormalizations(fun = c("norm_edgeR", "norm_CSS"),
    method = c("TMM", "CSS"))
ps_plaque_16S <- runNormalizations(normalization_list = my_norm,
    object = ps_plaque_16S)

# Initialize some limma based methods
my_limma <- set_limma(design = ~ 1 + RSID + HMP_BODY_SUBSITE,
    coef = "HMP_BODY_SUBSITESupragingival Plaque",
    norm = c("TMM", "CSS"))
# Make sure the subject ID variable is a factor
phyloseq::sample_data(ps_plaque_16S)[, "RSID"] <- as.factor(
  phyloseq::sample_data(ps_plaque_16S)[["RSID"]])

# Perform DA analysis
Plaque_16S_DA <- runDA(method_list = my_limma, object = ps_plaque_16S)

# Enrichment analysis
enrichment <- createEnrichment(object = Plaque_16S_DA,
  priorKnowledge = priorInfo, enrichmentCol = "Type", namesCol = "GENUS",
  slot = "pValMat", colName = "adjP", type = "pvalue", direction = "logFC",
  threshold_pvalue = 0.1, threshold_logfc = 1, top = 10, verbose = TRUE)

---

createMocks

---

### Description

Given the number of samples of the dataset from which the mocks should be created, this function produces a data.frame object with as many rows as the number of mocks and as many columns as the number of samples. If an odd number of samples is given, the lower even integer will be considered in order to obtain a balanced design for the mocks.

### Usage

```r
createMocks(nsamples, N = 1000)
```

### Arguments

- `nsamples` an integer representing the total number of samples.
- `N` number of mock comparison to generate.

### Value

a data.frame containing \(N\) rows and \(nsamples\) columns (if even). Each cell of the data frame contains the "grp1" or "grp2" characters which represent the mock groups pattern.

### Examples

```r
# Generate the pattern for 100 mock comparisons for an experiment with 30
# samples
mocks <- createMocks(nsamples = 30, N = 100)
head(mocks)
```
createPositives

Description

Inspect the list of p-values or/and log fold changes from the output of the differential abundance detection methods and count the True Positives (TP) and the False Positives (FP).

Usage

createPositives(
  object,
  priorKnowledge,
  enrichmentCol,
  namesCol = NULL,
  slot = "pValMat",
  colName = "adjP",
  type = "pvalue",
  direction = NULL,
  threshold_pvalue = 1,
  threshold_logfc = 0,
  top = NULL,
  alternative = "greater",
  verbose = FALSE,
  TP,
  FP
)

Arguments

object Output of differential abundance detection methods. pValMat, statInfo matrices, and method's name must be present (See vignette for detailed information).

priorKnowledge data.frame (with feature names as row.names) containing feature level metadata.

enrichmentCol name of the column containing information for enrichment analysis.

namesCol name of the column containing new names for features (default namesCol = NULL).

slot A character vector with 1 or number-of-methods-times repeats of the slot names where to extract values for each method (default slot = "pValMat").

colName A character vector with 1 or number-of-methods-times repeats of the column name of the slot where to extract values for each method (default colName = "rawP").

type A character vector with 1 or number-of-methods-times repeats of the value type of the column selected where to extract values for each method. Two values are possible: "pvalue" or "logfc" (default type = "pvalue").
createPositives

A character vector with 1 or number-of-methods-times repeats of the statInfo's column name containing information about the signs of differential abundance (usually log fold changes) for each method (default direction = NULL).

threshold_pvalue

A single or a numeric vector of thresholds for p-values. If present, features with p-values lower than threshold_pvalue are considered differentially abundant. Set threshold_pvalue = 1 to not filter by p-values.

threshold_logfc

A single or a numeric vector of thresholds for log fold changes. If present, features with log fold change absolute values higher than threshold_logfc are considered differentially abundant. Set threshold_logfc = 0 to not filter by log fold change values.

top

If not null, the top number of features, ordered by p-values or log fold change values, are considered as differentially abundant (default top = NULL).

alternative

indicates the alternative hypothesis and must be one of "two.sided", "greater" or "less". You can specify just the initial letter. Only used in the 2 × 2 case.

verbose

Boolean to display the kind of extracted values (default verbose = FALSE).

TP

A list of length-2 vectors. The entries in the vector are the direction ("UP Abundant", "DOWN Abundant", or "non-DA") in the first position, and the level of the enrichment variable (enrichmentCol) which is expected in that direction, in the second position.

FP

A list of length-2 vectors. The entries in the vector are the direction ("UP Abundant", "DOWN Abundant", or "non-DA") in the first position, and the level of the enrichment variable (enrichmentCol) which is not expected in that direction, in the second position.

Value

A data.frame object which contains the number of TPs and FPs features for each method and for each threshold of the top argument.

See Also

getPositives, plotPositives.

Examples

data("ps_plaque_16S")
data("microbial_metabolism")

# Extract genera from the phyloseq tax_table slot
genera <- phyloseq::tax_table(ps_plaque_16S)[, "GENUS"]
# Genera as rownames of microbial_metabolism data.frame
rownames(microbial_metabolism) <- microbial_metabolism$Genus
# Match OTUs to their metabolism
priorInfo <- data.frame(genera,
    "Type" = microbial_metabolism[genera, "Type"])
# Unmatched genera becomes "Unknown"
unknown_metabolism <- is.na(priorInfo$Type)
priorInfo[unknown_metabolism, "Type"] <- "Unknown"
priorInfo$Type <- factor(priorInfo$Type)

# Add a more informative names column
priorInfo[, "newNames"] <- paste0(rownames(priorInfo), priorInfo[, "GENUS"])

# Add some normalization/scaling factors to the phyloseq object
my_norm <- setNormalizations(fun = c("norm_edgeR", "norm_CSS"),
                              method = c("TMM", "CSS"))
ps_plaque_16S <- runNormalizations(normalization_list = my_norm,
                                    object = ps_plaque_16S)

# Initialize some limma based methods
my_limma <- set_limma(design = ~ 1 + RSID + HMP_BODY_SUBSITE,
                       coef = "HMP_BODY_SUBSITESupragingival Plaque",
                       norm = c("TMM", "CSS"))

# Make sure the subject ID variable is a factor
phyloseq::sample_data(ps_plaque_16S)[, "RSID"] <- as.factor(
  phyloseq::sample_data(ps_plaque_16S)[["RSID"]]
)

# Perform DA analysis
Plaque_16S_DA <- runDA(method_list = my_limma, object = ps_plaque_16S)

# Count TPs and FPs, from the top 1 to the top 20 features.
# As direction is supplied, features are ordered by "logFC" absolute values.
positives <- createPositives(object = Plaque_16S_DA,
                              priorKnowledge = priorInfo, enrichmentCol = "Type",
                              namesCol = "newNames", slot = "pValMat", colName = "rawP",
                              type = "pvalue", direction = "logFC", threshold_pvalue = 1,
                              threshold_logfc = 0, top = 1:20, alternative = "greater",
                              verbose = FALSE,
                              TP = list(c("DOWN Abundant", "Anaerobic"), c("UP Abundant", "Aerobic")),
                              FP = list(c("DOWN Abundant", "Aerobic"), c("UP Abundant", "Anaerobic")))

# Plot the TP-FP differences for each threshold
plotPositives(positives = positives)

---

createSplits

Description

Given a phyloseq or TreeSummarizedExperiment object from which the random splits should be created, this function produces a list of 2 data.frame objects: Subset1 and Subset2 with as many rows as the number of splits and as many columns as the half of the number of samples.

Usage

createSplits
  object,
createSplits

```r
assay_name = "counts",
varName = NULL,
paired = NULL,
balanced = TRUE,
N = 1000
)
```

### Arguments

- **object**
  - a phyloseq or TreeSummarizedExperiment object.
- **assay_name**
  - the name of the assay to extract from the TreeSummarizedExperiment object (default `assayName = "counts"`). Not used if the input object is a phyloseq.
- **varName**
  - name of a factor variable of interest.
- **paired**
  - name of the unique subject identifier variable. If specified, paired samples will remain in the same split. (default = NULL).
- **balanced**
  - If TRUE a balanced design will be created for the splits (default balanced = TRUE).
- **N**
  - number of splits to generate.

### Value

A list of 2 `data.frame` objects: `Subset1` and `Subset2` containing `N` rows and half of the total number of samples columns. Each cell contains a unique sample identifier.

### Examples

```r
data(ps_plaque_16S)
set.seed(123)

# Balanced design for repeated measures

# Balanced design for independent samples
splits_df <- createSplits(
  object = ps_plaque_16S, varName = "HMP_BODY_SUBSITE", balanced = TRUE, N = 100
)

# Unbalanced design
splits_df <- createSplits(
  object = ps_plaque_16S, varName = "HMP_BODY_SUBSITE", balanced = FALSE, N = 100
)
```
createTIEC

Description

Extract the list of p-values from the outputs of the differential abundance detection methods to compute several statistics to study the ability to control the type I error and the p-values distribution.

Usage

createTIEC(object)

Arguments

object Output of the differential abundance tests on mock comparisons. Must follow a specific structure with comparison, method, matrix of p-values, and method's name (See vignette for detailed information).

Value

A list of data.frames:

- df_pval 5 columns per number_of_features x methods x comparisons rows data.frame. The four columns are called Comparison, Method, variable (containing the feature names), pval, and padj;
- df_FPR 5 columns per methods x comparisons rows data.frame. For each set of method and comparison, the proportion of false positives, considering 3 thresholds (0.01, 0.05, 0.1) are reported;
- df_FDR 4 columns per methods rows data.frame. For each method, the average proportion of mock comparisons where false positives are found, considering 3 thresholds (0.01, 0.05, 0.1), are reported. Each value is an estimate of the nominal False Discovery Rate (FDR);
- df_QQ contains the coordinates to draw the QQ-plot to compare the mean observed p-value distribution across comparisons, with the theoretical uniform distribution;
- df_KS 5 columns and methods x comparisons rows data.frame. For each set of method and comparison, the Kolmogorov-Smirnov test statistics and p-values are reported in KS and KS_pval columns respectively.

See Also

createMocks

Examples

# Load some data
data(ps_stool_16S)

# Generate the patterns for 10 mock comparison for an experiment
# (N = 1000 is suggested)
mocks <- createMocks(nsamples = phyloseq::nsamples(ps_stool_16S), N = 10)
head(mocks)

# Add some normalization/scaling factors to the phyloseq object
my_norm <- setNormalizations(fun = c("norm_edgeR", "norm_CSS"),
                              method = c("TMM", "CSS"))
ps_stool_16S <- runNormalizations(normalization_list = my_norm,
                                  object = ps_stool_16S)

# Initialize some limma based methods
my_limma <- set_limma(design = ~ group, coef = 2,
                       norm = c("TMM", "CSS"))

# Run methods on mock datasets
results <- runMocks(mocks = mocks, method_list = my_limma,
                     object = ps_stool_16S)

# Prepare results for Type I Error Control
TIEC_summary <- createTIEC(results)

# Plot the results
plotFPR(df_FPR = TIEC_summary$df_FPR)
plotFDR(df_FDR = TIEC_summary$df_FDR)
plotQQ(df_QQ = TIEC_summary$df_QQ, zoom = c(0, 0.1))
plotKS(df_KS = TIEC_summary$df_KS)
plotLogP(df_QQ = TIEC_summary$df_QQ)

---

**Description**

Fast run for the ALDEx2’s differential abundance detection method. Support for Welch’s t, Wilcoxon, Kruskal-Wallace, Kruskal-Wallace glm ANOVA-like, and glm tests.

**Usage**

```r
DA_ALDEx2(
  object,
  assay_name = "counts",
  pseudo_count = FALSE,
  design = NULL,
  mc.samples = 128,
  test = c("t", "wilcox", "kw", "kw_glm", "glm"),
  paired.test = FALSE,
  denom = "all",
  contrast = NULL,
  verbose = TRUE
)
```
Arguments

<table>
<thead>
<tr>
<th>Argument</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>object</td>
<td>a phyloseq or TreeSummarizedExperiment object.</td>
</tr>
<tr>
<td>assay_name</td>
<td>the name of the assay to extract from the TreeSummarizedExperiment object (default assayName = &quot;counts&quot;). Not used if the input object is a phyloseq.</td>
</tr>
<tr>
<td>pseudo_count</td>
<td>add 1 to all counts if TRUE (default pseudo_count = FALSE).</td>
</tr>
<tr>
<td>design</td>
<td>a character with the name of a variable to group samples and compare them or a formula to compute a model.matrix (when test = &quot;glm&quot;).</td>
</tr>
<tr>
<td>mc.samples</td>
<td>an integer. The number of Monte Carlo samples to use when estimating the underlying distributions. Since we are estimating central tendencies, 128 is usually sufficient.</td>
</tr>
<tr>
<td>test</td>
<td>a character string. Indicates which tests to perform. &quot;t&quot; runs Welch’s t test while &quot;wilcox&quot; runs Wilcoxon test. &quot;kw&quot; runs Kruskal-Wallace test while &quot;kw_glm&quot; runs glm ANOVA-like test. &quot;glm&quot; runs a generalized linear model.</td>
</tr>
<tr>
<td>paired.test</td>
<td>A boolean. Toggles whether to do paired-sample tests. Applies to effect = TRUE and test = &quot;t&quot;.</td>
</tr>
<tr>
<td>denom</td>
<td>An any variable (all, iqlr, zero, lvha, median, user) indicating features to use as the denominator for the Geometric Mean calculation. The default &quot;all&quot; uses the geometric mean abundance of all features. Using &quot;median&quot; returns the median abundance of all features. Using &quot;iqlr&quot; uses the features that are between the first and third quartile of the variance of the clr values across all samples. Using &quot;zero&quot; uses the non-zero features in each group as the denominator. This approach is an extreme case where there are many nonzero features in one condition but many zeros in another. Using &quot;lvha&quot; uses features that have low variance (bottom quartile) and high relative abundance (top quartile in every sample). It is also possible to supply a vector of row indices to use as the denominator. Here, the experimentalist is determining a-priori which rows are thought to be invariant. In the case of RNA-seq, this could include ribosomal protein genes and and other house-keeping genes. This should be used with caution because the offsets may be different in the original data and in the data used by the function because features that are 0 in all samples are removed by aldex.clr.</td>
</tr>
<tr>
<td>contrast</td>
<td>character vector with exactly three elements: the name of a variable used in &quot;design&quot;, the name of the level of interest, and the name of the reference level. If &quot;kw&quot; or &quot;kw_glm&quot; as test, contrast vector is not used.</td>
</tr>
<tr>
<td>verbose</td>
<td>an optional logical value. If TRUE, information about the steps of the algorithm is printed. Default verbose = TRUE.</td>
</tr>
</tbody>
</table>

Value

A list object containing the matrix of p-values 'pValMat', the matrix of summary statistics for each tag 'statInfo', and a suggested ‘name’ of the final object considering the parameters passed to the function.

See Also

aldex for the Dirichlet-Multinomial model estimation. Several and more complex tests are present in the ALDEx2 framework.
Examples

```r
set.seed(1)
# Create a very simple phyloseq object
counts <- matrix(rnbinom(n = 300, size = 3, prob = 0.5), nrow = 50, ncol = 6)
metadata <- data.frame("Sample" = c("S1", "S2", "S3", "S4", "S5", "S6"),
                       "group" = as.factor(c("A", "A", "A", "B", "B", "B")))
ps <- phyloseq::phyloseq(phyloseq::otu_table(counts, taxa_are_rows = TRUE),
                        phyloseq::sample_data(metadata))
# Differential abundance with t test and denom defined by the user
DA_t <- DA_ALDEx2(ps, design = "group", test = "t", denom = c(1,2,3),
                   paired.test = FALSE, contrast = c("group", "B", "A"))
# Differential abundance with wilcox test and denom = "iqlr"
DA_w <- DA_ALDEx2(ps, design = "group", test = "wilcox", denom = "iqlr",
                   paired.test = FALSE, contrast = c("group", "B", "A"))
# Differential abundance with kw test and denom = "zero"
# mc.samples = 2 to speed up (128 suggested)
DA_kw <- DA_ALDEx2(ps, design = "group", test = "kw", denom = "zero",
                   mc.samples = 2)
# Differential abundance with kw.glm test and denom = "median"
DA_kw.glm <- DA_ALDEx2(ps, design = "group", test = "kw", denom = "median",
                       mc.samples = 2)
# Differential abundance with glm test and denom = "all"
DA_glm <- DA_ALDEx2(ps, design = ~ group, test = "glm", denom = "all",
                    mc.samples = 2, contrast = c("group", "B", "A"))
```

Description

Fast run for ANCOM and ANCOM-BC2 differential abundance detection methods.

Usage

```r
DA_ANCOM(
  object,
  assay_name = "counts",
  pseudo_count = FALSE,
  fix_formula = NULL,
  adj_formula = NULL,
  rand_formula = NULL,
  lme_control = lme4::lmerControl(),
  contrast = NULL,
  p_adj_method = "BH",
  struc_zero = FALSE,
  BC = TRUE,
  n_c1 = 1,
  verbose = TRUE
)
```
Arguments

- **object**: a phyloseq or TreeSummarizedExperiment object.
- **assay_name**: the name of the assay to extract from the TreeSummarizedExperiment object (default assayName = "counts"). Not used if the input object is a phyloseq.
- **pseudo_count**: add 1 to all counts if TRUE (default pseudo_count = FALSE).
- **fix_formula**: Used when BC = TRUE (ANCOM-BC2). The character string expresses how the microbial absolute abundances for each taxon depend on the fixed effects in metadata.
- **adj_formula**: Used when BC = FALSE (ANCOM). The character string represents the formula for covariate adjustment. Default is NULL.
- **rand_formula**: Optionally used when BC = TRUE or BC = FALSE. The character string expresses how the microbial absolute abundances for each taxon depend on the random effects in metadata. ANCOMB and ANCOM-BC2 follows the lmerTest package in formulating the random effects. See ?lmerTest::lmer for more details. Default is rand_formula = NULL.
- **lme_control**: a list of control parameters for mixed model fitting. See ?lme4::lmerControl for details.
- **contrast**: character vector with exactly, three elements: a string indicating the name of factor whose levels are the conditions to be compared, the name of the level of interest, and the name of the other level.
- **p_adj_method**: character. method to adjust p-values. Default is "holm". Options include "holm", "hochberg", "hommel", "bonferroni", "BH", "BY", "fdr", "none". See ?stats::p.adjust for more details.
- **struc_zero**: logical. Whether to detect structural zeros based on group. Default is FALSE. See Details for a more comprehensive discussion on structural zeros.
- **BC**: boolean for ANCOM method to use. If TRUE the bias correction (ANCOM-BC2) is computed (default BC = TRUE). When BC = FALSE computational time may increase and p-values are not computed.
- **n_cl**: numeric. The number of nodes to be forked. For details, see ?parallel::makeCluster. Default is 1 (no parallel computing).
- **verbose**: an optional logical value. If TRUE, information about the steps of the algorithm is printed. Default verbose = TRUE.

Value

A list object containing the matrix of p-values `pValMat`, a matrix of summary statistics for each tag `statInfo`, and a suggested ‘name’ of the final object considering the parameters passed to the function. ANCOM (BC = FALSE) does not produce p-values but W statistics. Hence, `pValMat` matrix is filled with \( 1 - W / (nfeatures - 1) \) values which are not p-values. To find DA features a threshold on this statistic can be used (liberal < 0.4, < 0.3, < 0.2, < 0.1 conservative).

See Also

- ancombc for analysis of microbiome compositions with bias correction or without it ancom.
Examples

```r
set.seed(1)
# Create a very simple phyloseq object
counts <- matrix(rnbinom(n = 60, size = 3, prob = 0.5), nrow = 10, ncol = 6)
metadata <- data.frame("Sample" = c("S1", "S2", "S3", "S4", "S5", "S6"),
  "group" = as.factor(c("A", "A", "A", "B", "B", "B")))
ps <- phyloseq::phyloseq(phyloseq::otu_table(counts, taxa_are_rows = TRUE),
  phyloseq::sample_data(metadata))

# Differential abundance
DA_ANCOM(object = ps, pseudo_count = FALSE, fix_formula = "group", contrast =
c("group", "B", "A"), verbose = FALSE)
```

Description

Fast run for basic differential abundance detection methods such as wilcox and t tests.

Usage

```r
DA_basic(
  object, 
  assay_name = "counts",
  pseudo_count = FALSE,
  contrast = NULL,
  test = c("t", "wilcox"),
  paired = FALSE,
  verbose = TRUE
)
```

Arguments

- **object**: a phyloseq or TreeSummarizedExperiment object.
- **assay_name**: the name of the assay to extract from the TreeSummarizedExperiment object (default assayName = "counts"). Not used if the input object is a phyloseq.
- **pseudo_count**: add 1 to all counts if TRUE (default pseudo_count = FALSE).
- **contrast**: character vector with exactly, three elements: a string indicating the name of factor whose levels are the conditions to be compared, the name of the level of interest, and the name of the other level.
- **test**: name of the test to perform. Choose between "t" or "wilcox".
- **paired**: boolean. Choose whether the test is paired or not (default paired = FALSE). If paired = TRUE be sure to provide the object properly ordered (by the grouping variable).
- **verbose**: an optional logical value. If TRUE, information about the steps of the algorithm is printed. Default verbose = TRUE.
**Value**

A list object containing the matrix of p-values 'pValMat', a matrix of summary statistics for each tag 'statInfo', and a suggested 'name' of the final object considering the parameters passed to the function.

**See Also**

`DA_Seurat` for a similar implementation of basic tests.

**Examples**

```r
set.seed(1)
# Create a very simple phyloseq object
counts <- matrix(rnbinom(n = 60, size = 3, prob = 0.5), nrow = 10, ncol = 6)
metadata <- data.frame("Sample" = c("S1", "S2", "S3", "S4", "S5", "S6"),
                      "group" = as.factor(c("A", "A", "A", "B", "B", "B")))
ps <- phyloseq::phyloseq(phyloseq::otu_table(counts, taxa_are_rows = TRUE),
                        phyloseq::sample_data(metadata))
# Differential abundance
DA_basic(object = ps, pseudo_count = FALSE, contrast = c("group", "B", "A"),
         test = "t", verbose = FALSE)
```

---

**Description**

Fast run for corncob differential abundance detection method.

**Usage**

```r
DA_corncob(
  object,
  assay_name = "counts",
  pseudo_count = FALSE,
  formula,
  phi.formula,
  formula_null,
  phi.formula_null,
  test,
  boot = FALSE,
  coefficient = NULL,
  verbose = TRUE
)
```
**DA_corncob**

**Arguments**

- **object**
  a phyloseq or TreeSummarizedExperiment object.

- **assay_name**
  the name of the assay to extract from the TreeSummarizedExperiment object (default `assayName = "counts"`). Not used if the input object is a phyloseq.

- **pseudo_count**
  add 1 to all counts if TRUE (default `pseudo_count = FALSE`).

- **formula**
  an object of class `formula` without the response: a symbolic description of the model to be fitted to the abundance.

- **phi.formula**
  an object of class `formula` without the response: a symbolic description of the model to be fitted to the dispersion.

- **formula_null**
  Formula for mean under null, without response

- **phi.formula_null**
  Formula for overdispersion under null, without response

- **test**
  Character. Hypothesis testing procedure to use. One of "Wald" or "LRT" (likelihood ratio test).

- **boot**
  Boolean. Defaults to FALSE. Indicator of whether or not to use parametric bootstrap algorithm. (See `pbWald` and `pbLRT`).

- **coefficient**
  The coefficient of interest as a single word formed by the variable name and the non reference level. (e.g.: 'ConditionDisease' if the reference level for the variable 'Condition' is 'control').

- **verbose**
  an optional logical value. If TRUE, information about the steps of the algorithm is printed. Default verbose = TRUE.

**Value**

A list object containing the matrix of p-values `pValMat`, the matrix of summary statistics for each tag `statInfo`, and a suggested `name` of the final object considering the parameters passed to the function.

**See Also**

`bbdml` and `differentialTest` for differential abundance and differential variance evaluation.

**Examples**

```R
set.seed(1)
# Create a very simple phyloseq object
counts <- matrix(rnbinom(n = 60, size = 3, prob = 0.5), nrow = 10, ncol = 6)
metadata <- data.frame("Sample" = c("S1", "S2", "S3", "S4", "S5", "S6"),
                      "group" = as.factor(c("A", "A", "A", "B", "B", "B")))
ps <- phyloseq::phyloseq(phyloseq::otu_table(counts, taxa_are_rows = TRUE),
                       phyloseq::sample_data(metadata))

# Differential abundance
DA_corncob(object = ps, formula = ~ group, phi.formula = ~ group,
           formula_null = ~ 1, phi.formula_null = ~ group, coefficient = "groupB",
           test = "Wald")
```
Description

Fast run for dearseq differential abundance detection method.

Usage

DA_dearseq(
  object,
  assay_name = "counts",
  pseudo_count = FALSE,
  covariates = NULL,
  variables2test = NULL,
  sample_group = NULL,
  test = c("permutation", "asymptotic"),
  preprocessed = FALSE,
  n_perm = 1000,
  verbose = TRUE
)

Arguments

object a phyloseq or TreeSummarizedExperiment object.
assay_name the name of the assay to extract from the TreeSummarizedExperiment object (default assayName = "counts"). Not used if the input object is a phyloseq.
pseudo_count add 1 to all counts if TRUE (default pseudo_count = FALSE).
covariates a character vector containing the colnames of the covariates to include in the model.
variables2test a character vector containing the colnames of the variable of interest.
sample_group a vector of length n indicating whether the samples should be grouped (e.g. paired samples or longitudinal data). Coerced to be a factor. Default is NULL in which case no grouping is performed.
test a character string indicating which method to use to approximate the variance component score test, either 'permutation' or 'asymptotic' (default test = "permutation").
preprocessed a logical flag indicating whether the expression data have already been preprocessed (e.g. log2 transformed). Default is FALSE, in which case log(y) is assumed to contain raw counts and is normalized into log(counts) per million.
n_perm the number of perturbations. Default is 1000
verbose an optional logical value. If TRUE, information about the steps of the algorithm is printed. Default verbose = TRUE.
Value

A list object containing the matrix of p-values ‘pValMat’, a matrix of summary statistics for each tag ‘statInfo’ which are still the p-values as this method does not produce other values, and a suggested ‘name’ of the final object considering the parameters passed to the function.

See Also

dear_seq for analysis of differential expression/abundance through a variance component test.

Examples

```r
set.seed(1)
# Create a very simple phyloseq object
counts <- matrix(rnbinom(n = 60, size = 3, prob = 0.5), nrow = 10, ncol = 6)
metadata <- data.frame("Sample" = c("S1", "S2", "S3", "S4", "S5", "S6"),
"group" = as.factor(c("A", "A", "A", "B", "B", "B")))
ps <- phyloseq::phyloseq(phyloseq::otu_table(counts, taxa_are_rows = TRUE),
phyloseq::sample_data(metadata))
# Differential abundance
DA_dearseq(object = ps, pseudo_count = FALSE, covariates = NULL,
variables2test = "group", sample_group = NULL, test = "asymptotic",
preprocessed = FALSE, verbose = TRUE)
```

Description

Fast run for DESeq2 differential abundance detection method.

Usage

```
DA_DESeq2(
  object,
  assay_name = "counts",
  pseudo_count = FALSE,
  design = NULL,
  contrast = NULL,
  alpha = 0.05,
  norm = c("ratio", "poscounts", "iterate"),
  weights,
  verbose = TRUE
)
```
Arguments

- **object**: a phyloseq or TreeSummarizedExperiment object.
- **assay_name**: the name of the assay to extract from the TreeSummarizedExperiment object (default `assayName = "counts"`). Not used if the input object is a phyloseq.
- **pseudo_count**: add 1 to all counts if TRUE (default `pseudo_count = FALSE`).
- **design**: character or formula to specify the model matrix.
- **contrast**: character vector with exactly three elements: the name of a factor in the design formula, the name of the numerator level for the fold change, and the name of the denominator level for the fold change.
- **alpha**: the significance cutoff used for optimizing the independent filtering (by default 0.05). If the adjusted p-value cutoff (FDR) will be a value other than 0.05, alpha should be set to that value.
- **norm**: name of the normalization method to use in the differential abundance analysis. Choose between the native DESeq2 normalization methods, such as `ratio`, `poscounts`, or `iterate`. Alternatively (only for advanced users), if `norm` is equal to "TMM", "TMMwsp", "RLE", "upperquartile", "posupperquartile", or "none" from `norm_edgeR`, "CSS" from `norm_CSS`, or "TSS" from `norm_TSS`, the normalization factors are automatically transformed into size factors. If custom factors are supplied, make sure they are compatible with DESeq2 size factors.
- **weights**: an optional numeric matrix giving observational weights.
- **verbose**: an optional logical value. If TRUE, information about the steps of the algorithm is printed. Default `verbose = TRUE`.

Value

A list object containing the matrix of p-values ‘pValMat’, the dispersion estimates ‘dispEsts’, the matrix of summary statistics for each tag ‘statInfo’, and a suggested ‘name’ of the final object considering the parameters passed to the function.

See Also

- `phyloseq_to_deseq2` for phyloseq to DESeq2 object conversion, DESeq and results for the differential abundance method.

Examples

```r
set.seed(1)
# Create a very simple phyloseq object
counts <- matrix(rnbinom(n = 60, size = 3, prob = 0.5), nrow = 10, ncol = 6)
metadata <- data.frame("Sample" = c("S1", "S2", "S3", "S4", "S5", "S6"),
                      "group" = as.factor(c("A", "A", "A", "B", "B", "B"))
ps <- phyloseq::phyloseq(phyloseq::otu_table(counts, taxa_are_rows = TRUE),
                      phyloseq::sample_data(metadata))
# Calculate the poscounts size factors
ps_NF <- norm_DESeq2(object = ps, method = "poscounts")
# The phyloseq object now contains the size factors:
sizeFacts <- phyloseq::sample_data(ps_NF)[, "NF.poscounts"]
```
head(sizeFacts)
# Differential abundance
DA_DESeq2(object = ps_NF, pseudo_count = FALSE, design = ~ group, contrast =
c("group", "B", "A"), norm = "poscounts")

---

**Description**

Fast run for edgeR differential abundance detection method.

**Usage**

```
DA_edgeR(
  object,
  assay_name = "counts",
  pseudo_count = FALSE,
  group_name = NULL,
  design = NULL,
  robust = FALSE,
  coef = 2,
  norm = c("TMM", "TMMwsp", "RLE", "upperquartile", "posupperquartile", "none"),
  weights,
  verbose = TRUE
)
```

**Arguments**

- `object` a phyloseq or TreeSummarizedExperiment object.
- `assay_name` the name of the assay to extract from the TreeSummarizedExperiment object (default assayName = "counts"). Not used if the input object is a phyloseq.
- `pseudo_count` add 1 to all counts if TRUE (default pseudo_count = FALSE).
- `group_name` character giving the name of the column containing information about experimental group/condition for each sample/library.
- `design` character or formula to specify the model matrix.
- `robust` logical, should the estimation of prior_df be robustified against outliers?
- `coef` integer or character index vector indicating which coefficients of the linear model are to be tested equal to zero.
- `norm` name of the normalization method to use in the differential abundance analysis. Choose between the native edgeR normalization methods, such as TMM, TMMwsp, RLE, upperquartile, posupperquartile, or none. Alternatively (only for advanced users), if `norm` is equal to "ratio", "poscounts", or "iterate" from `norm_DESeq2`, "CSS" from `norm_CSS`, or "TSS" from `norm_TSS`, the scaling factors are automatically transformed into normalization factors. If custom factors are supplied, make sure they are compatible with edgeR normalization factors.
weights an optional numeric matrix giving observational weights.
verbose an optional logical value. If TRUE, information about the steps of the algorithm is printed. Default verbose = TRUE.

Value
A list object containing the matrix of p-values pValMat, the dispersion estimates dispEsts, the matrix of summary statistics for each tag statInfo, and a suggested name of the final object considering the parameters passed to the function.

See Also
DGEList for the edgeR DEG object creation, estimateDisp and estimateGLMRobustDisp for dispersion estimation, and glmQLFit and glmQLFTest for the quasi-likelihood negative binomial model fit.

Examples
set.seed(1)
# Create a very simple phyloseq object
counts <- matrix(rnbinom(n = 60, size = 3, prob = 0.5), nrow = 10, ncol = 6)
metadata <- data.frame("Sample" = c("S1", "S2", "S3", "S4", "S5", "S6"),
  "group" = as.factor(c("A", "A", "A", "B", "B", "B")))
ps <- phyloseq::phyloseq(phyloseq::otu_table(counts, taxa_are_rows = TRUE),
  phyloseq::sample_data(metadata))

  # Calculate the TMM normalization factors
ps_NF <- norm_edgeR(object = ps, method = "TMM")
  # The phyloseq object now contains the normalization factors:
normFacts <- phyloseq::sample_data(ps_NF)[, "NF.TMM"]
head(normFacts)

  # Differential abundance
DA_edgeR(object = ps_NF, pseudo_count = FALSE, group_name = "group",
  design = ~ group, coef = 2, robust = FALSE, norm = "TMM")
DA_limma

```r
design = NULL,
coef = 2,
norm = c("TMM", "TMMwsp", "RLE", "upperquartile", "posupperquartile", "none"),
weights,
verbose = TRUE
)
```

Arguments

- **object**: a phyloseq or TreeSummarizedExperiment object.
- **assay_name**: the name of the assay to extract from the TreeSummarizedExperiment object (default `assayName = "counts"`). Not used if the input object is a phyloseq.
- **pseudo_count**: add 1 to all counts if TRUE (default `pseudo_count = FALSE`).
- **design**: character name of the metadata columns, formula, or design matrix with rows corresponding to samples and columns to coefficients to be estimated.
- **coef**: integer or character index vector indicating which coefficients of the linear model are to be tested equal to zero.
- **norm**: name of the normalization method to use in the differential abundance analysis. Choose between the native edgeR normalization methods, such as TMM, TMMwsp, RLE, upperquartile, posupperquartile, or none. Alternatively (only for advanced users), if `norm` is equal to "ratio", "poscounts", or "iterate" from `norm_DESeq2`, "CSS" from `norm_CSS`, or "TSS" from `norm_TSS`, the scaling factors are automatically transformed into normalization factors. If custom factors are supplied, make sure they are compatible with edgeR normalization factors.
- **weights**: an optional numeric matrix giving observational weights.
- **verbose**: an optional logical value. If `TRUE`, information about the steps of the algorithm is printed. Default `verbose = TRUE`.

Value

A list object containing the matrix of p-values `pValMat`, the matrix of summary statistics for each tag `statInfo`, and a suggested `name` of the final object considering the parameters passed to the function.

See Also

- `voom` for the mean-variance relationship estimation,
- `lmFit` for the linear model framework.

Examples

```r
set.seed(1)
# Create a very simple phyloseq object
counts <- matrix(rnbinom(n = 60, size = 3, prob = 0.5), nrow = 10, ncol = 6)
metadata <- data.frame("Sample" = c("S1", "S2", "S3", "S4", "S5", "S6"),
    "group" = as.factor(c("A", "A", "A", "B", "B", "B")))
ps <- phyloseq::phyloseq(phyloseq::otu_table(counts, taxa_are_rows = TRUE),
    phyloseq::sample_data(metadata))
# Calculate the TMM normalization factors
```
ps_NF <- norm_edgeR(object = ps, method = "TMM")
# The phyloseq object now contains the normalization factors:
normFacts <- phyloseq::sample_data(ps_NF)[, "NF.TMM"]
head(normFacts)
# Differential abundance
DA_limma(object = ps_NF, pseudo_count = FALSE, design = ~ group, coef = 2,
       norm = "TMM")

---

**Description**

Fast run for MAST differential abundance detection method.

**Usage**

```r
DA_MAST(
  object,
  assay_name = "counts",
  pseudo_count = FALSE,
  rescale = c("median", "default"),
  design,
  coefficient = NULL,
  verbose = TRUE
)
```

**Arguments**

- `object`: a phyloseq or TreeSummarizedExperiment object.
- `assay_name`: the name of the assay to extract from the TreeSummarizedExperiment object (default assayName = "counts"). Not used if the input object is a phyloseq.
- `pseudo_count`: add 1 to all counts if TRUE (default pseudo_count = FALSE).
- `rescale`: Rescale count data, per million if 'default', or per median library size if 'median' ('median' is suggested for metagenomics data).
- `design`: The model for the count distribution. Can be the variable name, or a character similar to "~ 1 + group", or a formula, or a 'model.matrix' object.
- `coefficient`: The coefficient of interest as a single word formed by the variable name and the non reference level. (e.g.: 'ConditionDisease' if the reference level for the variable 'Condition' is 'control').
- `verbose`: an optional logical value. If TRUE, information about the steps of the algorithm is printed. Default verbose = TRUE.

**Value**

A list object containing the matrix of p-values 'pValMat', the matrix of summary statistics for each tag 'statInfo', and a suggested 'name' of the final object considering the parameters passed to the function.
**DA_metagenomeSeq**

**See Also**

`zlm` for the Truncated Gaussian Hurdle model estimation.

**Examples**

```r
set.seed(1)
# Create a very simple phyloseq object
counts <- matrix(rnbinom(n = 60, size = 3, prob = 0.5), nrow = 10, ncol = 6)
metadata <- data.frame("Sample" = c("S1", "S2", "S3", "S4", "S5", "S6"),
                        "group" = as.factor(c("A", "A", "A", "B", "B", "B")))
ps <- phyloseq::phyloseq(phyloseq::otu_table(counts, taxa_are_rows = TRUE),
                        phyloseq::sample_data(metadata))

# Differential abundance
DA_MAST(object = ps, pseudo_count = FALSE, rescale = "median",
        design = ~ group, coefficient = "groupB")
```

---

**Description**

Fast run for the metagenomeSeq's differential abundance detection method.

**Usage**

```r
DA_metagenomeSeq(
  object,
  assay_name = "counts",
  pseudo_count = FALSE,
  design = NULL,
  coef = 2,
  norm = "CSS",
  model = "fitFeatureModel",
  verbose = TRUE
)
```

**Arguments**

- **object**
  - a phyloseq or TreeSummarizedExperiment object.
- **assay_name**
  - the name of the assay to extract from the TreeSummarizedExperiment object (default `assayName = "counts"`). Not used if the input object is a phyloseq.
- **pseudo_count**
  - add 1 to all counts if TRUE (default `pseudo_count = FALSE`).
- **design**
  - the model for the count distribution. Can be the variable name, or a character similar to "~ 1 + group", or a formula.
- **coef**
  - coefficient of interest to grab log fold-changes.
**norm**

name of the normalization method to use in the differential abundance analysis. Choose the native metagenomeSeq normalization method CSS. Alternatively (only for advanced users), if norm is equal to "TMM", "TMMwsp", "RLE", "upperquartile", "posupperquartile", or "none" from norm_edgeR, "ratio", "poscounts", or "iterate" from norm_DESeq2, or "TSS" from norm_TSS, the factors are automatically transformed into scaling factors. If custom factors are supplied, make sure they are compatible with metagenomeSeq normalization factors.

**model**

character equal to "fitFeatureModel" for differential abundance analysis using a zero-inflated log-normal model, "fitZig" for a complex mathematical optimization routine to estimate probabilities that a zero for a particular feature in a sample is a technical zero or not. The latter model relies heavily on the limma package (default model = "fitFeatureModel").

**verbose**

an optional logical value. If TRUE, information about the steps of the algorithm is printed. Default verbose = TRUE.

**Value**

A list object containing the matrix of p-values 'pValMat', the matrix of summary statistics for each tag 'statInfo', and a suggested 'name' of the final object considering the parameters passed to the function.

**See Also**

*fitZig* for the Zero-Inflated Gaussian regression model estimation and *MRfulltable* for results extraction.

**Examples**

```r
set.seed(1)
# Create a very simple phyloseq object
counts <- matrix(rnbinom(n = 60, size = 3, prob = 0.5), nrow = 10, ncol = 6)
metadata <- data.frame("Sample" = c("S1", "S2", "S3", "S4", "S5", "S6"),
                      "group" = as.factor(c("A", "A", "A", "B", "B", "B")))
ps <- phyloseq::phyloseq(phyloseq::otu_table(counts, taxa_are_rows = TRUE),
                      phyloseq::sample_data(metadata))
# Calculate the CSS normalization factors
ps_NF <- norm_CSS(object = ps, method = "CSS")
# The phyloseq object now contains the normalization factors:
normFacts <- phyloseq::sample_data(ps_NF)[, "NF.CSS"]
head(normFacts)
# Differential abundance
DA_metagenomeSeq(object = ps_NF, pseudo_count = FALSE, design = ~ group,
                 coef = 2, norm = "CSS")
```
**Description**

Fast run for NOISeqBIO differential abundance detection method. It computes differential expression between two experimental conditions.

**Usage**

```r
DA_NOISeq(
  object,
  assay_name = "counts",
  pseudo_count = FALSE,
  contrast = NULL,
  norm = c("rpkm", "uqua", "tmm", "n"),
  verbose = TRUE
)
```

**Arguments**

- `object`: a phyloseq or TreeSummarizedExperiment object.
- `assay_name`: the name of the assay to extract from the TreeSummarizedExperiment object (default `assayName = "counts"`). Not used if the input object is a phyloseq.
- `pseudo_count`: add 1 to all counts if TRUE (default `pseudo_count = FALSE`).
- `contrast`: character vector with exactly, three elements: a string indicating the name of factor whose levels are the conditions to be compared, the name of the level of interest, and the name of the other level.
- `norm`: name of the normalization method to use in the differential abundance analysis. Choose between the native edgeR normalization methods, such as TMM, TMMwsp, RLE, upperquartile, posupperquartile, or none. Alternatively (only for advanced users), if `norm` is equal to "ratio", "poscounts", or "iterate" from `norm_DESeq2`, "CSS" from `norm_CSS`, or "TSS" from `norm_TSS`, the scaling factors are automatically transformed into normalization factors. If custom factors are supplied, make sure they are compatible with edgeR normalization factors.
- `verbose`: an optional logical value. If TRUE, information about the steps of the algorithm is printed. Default `verbose = TRUE`.

**Value**

A list object containing the matrix of p-values `pValMat`, a matrix of summary statistics for each tag `statInfo`, and a suggested `name` of the final object considering the parameters passed to the function. NOISeq does not produce p-values but an estimated probability of differential expression for each feature. Note that these probabilities are not equivalent to p-values. The higher the probability, the more likely that the difference in abundance is due to the change in the experimental condition and not to chance... Hence, `pValMat` matrix is filled with 1 - prob values which can be interpreted as 1 - FDR. Where FDR can be considered as an adjusted p-value (see NOISeq vignette).
See Also

noiseqbio for analysis of differential expression/abundance between two experimental conditions from read count data.

Examples

```r
set.seed(1)
# Create a very simple phyloseq object
counts <- matrix(rnbinom(n = 60, size = 3, prob = 0.5), nrow = 10, ncol = 6)
metadata <- data.frame("Sample" = c("S1", "S2", "S3", "S4", "S5", "S6"),
  "group" = as.factor(c("A", "A", "A", "B", "B", "B")))
ps <- phyloseq::phyloseq(phyloseq::otu_table(counts, taxa_are_rows = TRUE),
  phyloseq::sample_data(metadata))
# Differential abundance
DA_NOISeq(object = ps, pseudo_count = FALSE, contrast = c("group", "B", "A"),
  norm = "tmm", verbose = FALSE)
```

Description

Fast run for Seurat differential abundance detection method.

Usage

```r
DA_Seurat(
  object,
  assay_name = "counts",
  pseudo_count = FALSE,
  norm = "LogNormalize",
  scale.factor = 10000,
  test = "wilcox",
  contrast,
  verbose = TRUE
)
```

Arguments

- **object**: a phyloseq or TreeSummarizedExperiment object.
- **assay_name**: the name of the assay to extract from the TreeSummarizedExperiment object (default assayName = "counts"). Not used if the input object is a phyloseq.
- **pseudo_count**: add 1 to all counts if TRUE (default pseudo_count = FALSE).
- **norm**: Method for normalization.
  - *LogNormalize*: Feature counts for each sample are divided by the total counts of that sample and multiplied by the scale.factor. This is then natural-log transformed using log1p;
- CLR Applies a centered log ratio transformation;
- RC Relative counts. Feature counts for each sample are divided by the total counts of that sample and multiplied by the scale.factor. No log-transformation is applied. For counts per million (CPM) set scale.factor = 1e6;
- none No normalization

scale.factor: Sets the scale factor for cell-level normalization

test: Denotes which test to use. Available options are:
  - "wilcox" Identifies differentially abundant features between two groups of samples using a Wilcoxon Rank Sum test (default).
  - "bimod" Likelihood-ratio test for the feature abundances, (McDavid et al., Bioinformatics, 2013).
  - "roc" Identifies 'markers' of feature abundance using ROC analysis. For each feature, evaluates (using AUC) a classifier built on that feature alone, to classify between two groups of cells. An AUC value of 1 means that abundance values for this feature alone can perfectly classify the two groupings (i.e. Each of the samples in group.1 exhibit a higher level than each of the samples in group.2). An AUC value of 0 also means there is perfect classification, but in the other direction. A value of 0.5 implies that the feature has no predictive power to classify the two groups. Returns a 'predictive power' (abs(AUC-0.5) * 2) ranked matrix of putative differentially expressed genes.
  - "t" Identify differentially abundant features between two groups of samples using the Student’s t-test.
  - "negbinom" Identifies differentially abundant features between two groups of samples using a negative binomial generalized linear model.
  - "poisson" Identifies differentially abundant features between two groups of samples using a poisson generalized linear model.
  - "LR" Uses a logistic regression framework to determine differentially abundant features. Constructs a logistic regression model predicting group membership based on each feature individually and compares this to a null model with a likelihood ratio test.
  - "MAST" Identifies differentially expressed genes between two groups of cells using a hurdle model tailored to scRNA-seq data. Utilizes the MAST package to run the DE testing.
  - "DESeq2" Identifies differentially abundant features between two groups of samples based on a model using DESeq2 which uses a negative binomial distribution (Love et al, Genome Biology, 2014).

contrast: character vector with exactly three elements: the name of a factor in the design formula, the name of the numerator level for the fold change, and the name of the denominator level for the fold change.

verbose: an optional logical value. If TRUE, information about the steps of the algorithm is printed. Default verbose = TRUE.
**Value**

A list object containing the matrix of p-values 'pValMat', the matrix of summary statistics for each tag 'statInfo', and a suggested ‘name’ of the final object considering the parameters passed to the function.

**See Also**

CreateSeuratObject to create the Seurat object, AddMetaData to add metadata information, NormalizeData to compute the normalization for the counts, FindVariableFeatures to estimate the mean-variance trend, ScaleData to scale and center features in the dataset, and FindMarkers to perform differential abundance analysis.

**Examples**

```r
set.seed(1)
# Create a very simple phyloseq object
counts <- matrix(rnbinom(n = 60, size = 3, prob = 0.5), nrow = 10, ncol = 6)
metadata <- data.frame("Sample" = c("S1", "S2", "S3", "S4", "S5", "S6"),
                      "group" = as.factor(c("A", "A", "A", "B", "B", "B")))
ps <- phyloseq::phyloseq(phyloseq::otu_table(counts, taxa_are_rows = TRUE),
                      phyloseq::sample_data(metadata))

# Differential abundance
DA_Seurat(object = ps, contrast = c("group","B","A"))

# Perform a simple Wilcoxon test using Seurat on raw data
DA_Seurat(object = ps, contrast = c("group","B","A"), norm = "none",
             test = "wilcox")
```

---

enrichmentTest

**Description**

Perform the Fisher exact test for all the possible 2x2 contingency tables, considering differential abundance direction and enrichment variable.

**Usage**

```r
enrichmentTest(method, enrichmentCol, alternative = "greater")
```

**Arguments**

- **method**: Output of differential abundance detection method in which DA information is extracted by the getDA function and the information related to enrichment is appropriately added through the addKnowledge.
- **enrichmentCol**: name of the column containing information for enrichment analysis.
- **alternative**: indicates the alternative hypothesis and must be one of "two.sided", "greater" or "less". You can specify just the initial letter. Only used in the 2 x 2 case.
Value

a list of objects:

• data a data.frame object with DA directions, statistics, and feature names;
• tables a list of 2x2 contingency tables;
• tests the list of Fisher exact tests’ p-values for each contingency table;
• summaries a list with the first element of each contingency table and its p-value (for graphical purposes);

See Also

extractDA, addKnowledge, and createEnrichment

Examples

data("ps_plaque_16S")
data("microbial_metabolism")

# Extract genera from the phyloseq tax_table slot
genera <- phyloseq::tax_table(ps_plaque_16S)[, "GENUS"]
# Genera as rownames of microbial_metabolism data.frame
rownames(microbial_metabolism) <- microbial_metabolism$Genus
# Match OTUs to their metabolism
priorInfo <- data.frame(genera,
   "Type" = microbial_metabolism[, "Type"])
# Unmatched genera becomes "Unknown"
unknown_metabolism <- is.na(priorInfo$Type)
priorInfo[unknown_metabolism, "Type"] <- "Unknown"
priorInfo$Type <- factor(priorInfo$Type)
# Add a more informative names column
priorInfo[, "newNames"] <- paste0(rownames(priorInfo), priorInfo[, "GENUS"])

# DA Analysis

# Make sure the subject ID variable is a factor
phyloseq::sample_data(ps_plaque_16S)[, "RSID"] <- as.factor(
   phyloseq::sample_data(ps_plaque_16S)[["RSID"]])

# Add scaling factors
ps_plaque_16S <- norm_edgeR(object = ps_plaque_16S, method = "TMM")

# DA analysis
da.limma <- DA_limma(
   object = ps_plaque_16S,
   design = ~ 1 + RSID + HMP_BODY_SUBSITE,
   coef = "HMP_BODY_SUBSITESupragingival Plaque",
   norm = "TMM"
)

DA <- getDA(method = da.limma, slot = "pValMat", colName = "adjP",
   type = "pvalue", direction = "logFC", threshold_pvalue = 0.05,
threshold_logfc = 1, top = NULL)

# Add a priori information
DA_info <- addKnowledge(method = DA, priorKnowledge = priorInfo,
                          enrichmentCol = "Type", namesCol = "newNames")

# Create contingency tables and compute F tests
DA_info_enriched <- enrichmentTest(method = DA_info, enrichmentCol = "Type",
                                    alternative = "greater")

---

**extractDA**

**extractDA**

---

**Description**

Inspect the list of p-values or/and log fold changes from the output of differential abundance detection methods.

**Usage**

```r
extractDA(
  object,
  slot = "pValMat",
  colName = "adjP",
  type = "pvalue",
  direction = NULL,
  threshold_pvalue = 1,
  threshold_logfc = 0,
  top = NULL,
  verbose = FALSE
)
```

**Arguments**

- **object**: Output of differential abundance detection methods. `pValMat`, `statInfo` matrices, and method's name must be present (See vignette for detailed information).
- **slot**: A character vector with 1 or number-of-methods-times repeats of the slot names where to extract values for each method (default slot = "pValMat").
- **colName**: A character vector with 1 or number-of-methods-times repeats of the column name of the slot where to extract values for each method (default colName = "rawP").
- **type**: A character vector with 1 or number-of-methods-times repeats of the value type of the column selected where to extract values for each method. Two values are possible: "pvalue" or "logfc" (default type = "pvalue").
- **direction**: A character vector with 1 or number-of-methods-times repeats of the `statInfo`'s column name containing information about the signs of differential abundance (usually log fold changes) for each method (default direction = NULL).
threshold_pvalue
A single or a numeric vector of thresholds for p-values. If present, features with p-values lower than threshold_pvalue are considered differentially abundant. Set threshold_pvalue = 1 to not filter by p-values.

threshold_logfc
A single or a numeric vector of thresholds for log fold changes. If present, features with log fold change absolute values higher than threshold_logfc are considered differentially abundant. Set threshold_logfc = 0 to not filter by log fold change values.

top
If not null, the top number of features, ordered by p-values or log fold change values, are considered as differentially abundant (default top = NULL).

verbose
Boolean to display the kind of extracted values (default verbose = FALSE).

Value
A data.frame with several columns for each method:

- stat which contains the p-values or the absolute log fold change values;
- direction which is present if direction was supplied, it contains the information about directionality of differential abundance (usually log fold changes);
- DA which can contain several values according to thresholds and inputs. "DA" or "non-DA" if direction = NULL, "UP Abundant", "DOWN Abundant", or "non-DA" otherwise.

See Also
getDA, extractStatistics

Examples

data("ps_plaque_16S")
# Add scaling factors
my_norm <- setNormalizations(fun = c("norm_edgeR", "norm_CSS"),
  method = c("TMM", "CSS"))
ps_plaque_16S <- runNormalizations(normalization_list = my_norm,
  object = ps_plaque_16S)
# Perform DA analysis
my_methods <- set_limma(design = ~ 1 + HMP_BODY_SUBSITE, coef = 2,
  norm = c("TMM", "CSS"))
Plaque_16S_DA <- runDA(method_list = my_methods, object = ps_plaque_16S)
# Top 10 features (ordered by 'direction') are DA
DA_1 <- extractDA(
  object = Plaque_16S_DA, slot = "pValMat", colName = "adjP",
  type = "pvalue", direction = "logFC", threshold_pvalue = 1,
  threshold_logfc = 0, top = 10
)
# Features with p-value < 0.05 and |logFC| > 1 are DA
DA_2 <- extractDA(
  object = Plaque_16S_DA, slot = "pValMat", colName = "adjP",
  type = "pvalue", direction = "logFC", threshold_pvalue = 0.05,
  threshold_logfc = 1, top = NULL
)
extractStatistics

Description

Extract the list of p-values or/and log fold changes from the outputs of the differential abundance detection methods.

Usage

extractStatistics(
  object,
  slot = "pValMat",
  colName = "rawP",
  type = "pvalue",
  direction = NULL,
  verbose = FALSE
)

Arguments

  object
    Output of differential abundance detection methods. pValMat, statInfo matrices, and method’s name must be present (See vignette for detailed information).
  slot
    A character vector with 1 or number-of-methods-times repeats of the slot names where to extract values for each method (default slot = "pValMat").
  colName
    A character vector with 1 or number-of-methods-times repeats of the column name of the slot where to extract values for each method (default colName = "rawP").
  type
    A character vector with 1 or number-of-methods-times repeats of the value type of the column selected where to extract values for each method. Two values are possible: "pvalue" or "logfc" (default type = "pvalue").
  direction
    A character vector with 1 or number-of-methods-times repeats of the statInfo’s column name containing information about the signs of differential abundance (usually log fold changes) for each method (default direction = NULL).
  verbose
    Boolean to display the kind of extracted values (default verbose = FALSE).

Value

A vector or a data.frame for each method. If direction = NULL, the colname column values, transformed according to type (not tranformed if type = "pvalue", -abs(value) if type = "logfc"), of the slot are reported, otherwise the direction column of the statInfo matrix is added to the output.

See Also

generateStatistics
Examples

data("ps_plaque_16S")
# Add scaling factors
my_norm <- setNormalizations(fun = c("norm_edgeR", "norm_CSS"),
method = c("TMM", "CSS"))
ps_plaque_16S <- runNormalizations(normalization_list = my_norm,
object = ps_plaque_16S)
# Perform DA analysis
my_methods <- set_limma(design = ~ 1 + HMP_BODY_SUBSITE, coef = 2,
norm = c("TMM", "CSS"))
Plaque_16S_DA <- runDA(method_list = my_methods, object = ps_plaque_16S)
### Extract statistics for concordance analysis:
# Only p-values
extracted_pvalues <- extractStatistics(
  object = Plaque_16S_DA, slot =
  "pValMat", colName = "rawP", type = "pvalue"
)
# Only transformed log fold changes -\text{abs}(\text{logFC})
extracted_abslfc <- extractStatistics(
  object = Plaque_16S_DA, slot =
  "statInfo", colName = "logFC", type = "logfc"
)
# Only transformed log fold changes for a method and p-values for the other
extracted_abslfc_pvalues <- extractStatistics(
  object = Plaque_16S_DA,
  slot = c("statInfo", "pValMat"), colName = c("logFC", "rawP"), type =
  c("logfc", "pvalue")
)
### Extract statistics for enrichment analysis:
# p-values and log fold changes
extracted_pvalues_and_lfc <- extractStatistics(
  object = Plaque_16S_DA,
  slot = "pValMat", colName = "rawP", type = "pvalue", direction = "logFC"
)
# transformed log fold changes and untouched log fold changes
extracted_abslfc_and_lfc <- extractStatistics(
  object = Plaque_16S_DA,
  slot = "statInfo", colName = "logFC", type = "logfc", direction =
  "logFC"
)
# Only transformed log fold changes for a method and p-values for the other
extracted_mix <- extractStatistics(
  object = Plaque_16S_DA,
  slot = c("statInfo", "pValMat"), colName = c("logFC", "rawP"), type =
  c("logfc", "pvalue"), direction = "logFC"
)
Description

Fit a Dirichlet-Multinomial (DM) distribution for each taxon of the count data. The model estimation procedure is performed by MGLM MGLMreg function without assuming the presence of any group in the samples (design matrix equal to a column of ones.)

Usage

\[ \text{fitDM}\text{(object, assay\_name = "counts", verbose = TRUE)} \]

Arguments

- **object**: a phyloseq object, a TreeSummarizedExperiment object, or a matrix of counts.
- **assay\_name**: the name of the assay to extract from the TreeSummarizedExperiment object (default assayName = "counts"). Not used if the input object is a phyloseq.
- **verbose**: an optional logical value. If TRUE information on the steps of the algorithm is printed. Default verbose = TRUE.

Value

A data frame containing the continuity corrected logarithms of the average fitted values for each row of the matrix of counts in the \( Y \) column, and the estimated probability to observe a zero in the \( Y0 \) column.

Examples

```r
# Generate some random counts
counts = matrix(rnbinom(n = 60, size = 3, prob = 0.5), nrow = 10, ncol = 6)
# Fit model on the counts matrix
DM <- fitDM(counts)
head(DM)
```

---

Description

Fit a truncated gaussian hurdle model for each taxon of the count data. The hurdle model estimation procedure is performed by MAST zlm function without assuming the presence of any group in the samples (design matrix equal to a column of ones.)

Usage

\[ \text{fitHURDLE}\text{(object, assay\_name = "counts", scale = "default", verbose = TRUE)} \]
**Arguments**

- **object**: a phyloseq object, a TreeSummarizedExperiment object, or a matrix of counts.
- **assay_name**: the name of the assay to extract from the TreeSummarizedExperiment object (default assayName = "counts"). Not used if the input object is a phyloseq.
- **scale**: Character vector, either median or default to choose between the median of the library size or one million to scale raw counts.
- **verbose**: an optional logical value. If TRUE information on the steps of the algorithm is printed. Default verbose = TRUE.

**Value**

A data frame containing the continuity corrected logarithms of the average fitted values for each row of the matrix of counts in the Y column, and the estimated probability to observe a zero in the Y0 column.

**Examples**

```r
# Generate some random counts
counts = matrix(rnbinom(n = 60, size = 3, prob = 0.5), nrow = 10, ncol = 6)
# Fit model on the counts matrix
HURDLE <- fitHURDLE(counts, scale = "median")
head(HURDLE)
```

**Description**

A wrapper function that fits the specified models for each taxon of the count data and computes the mean difference (MD) and zero probability difference (ZPD) between estimated and observed values.

**Usage**

```r
fitModels(
  object, 
  assay_name = "counts", 
  models = c("NB", "ZINB", "DM", "ZIG", "HURDLE"), 
  scale_HURDLE = c("default", "median"), 
  verbose = TRUE 
)
```
Arguments

object a phyloseq object, a TreeSummarizedExperiment object, or a matrix of counts.
assay_name the name of the assay to extract from the TreeSummarizedExperiment object (default assayName = "counts"). Not used if the input object is a phyloseq.
models character vector which assumes the values NB, ZINB, DM, ZIG, and HURDLE.
scale_HURDLE character vector, either median or default to choose between the median of the library size or one million to scale raw counts for the truncated gaussian hurdle model.
verbose an optional logical value. If TRUE information on the steps of the algorithm is printed. Default verbose = TRUE.

Value

list of data.frame objects for each model. The first two columns contain the properly transformed observed values for mean and zero proportion, while the third and the fourth columns contain the estimated values for the mean and the zero rate respectively.

See Also

fitNB, fitZINB, fitDM, fitZIG, and fitHURDLE for the model estimations, prepareObserved for raw counts preparation, and meanDifferences for the Mean Difference (MD) and Zero Probability Difference (ZPD) computations.

Examples

# Generate some random counts
counts <- matrix(rnbinom(n = 60, size = 3, prob = 0.5), nrow = 10, ncol = 6)
# Estimate the counts assuming several distributions
GOF <- fitModels(
  object = counts, models = c("NB", "ZINB", "DM", "ZIG", "HURDLE"),
  scale_HURDLE = c("median", "default")
)
head(GOF)
Usage

fitNB(object, assay_name = "counts", verbose = TRUE)

Arguments

- object: a phyloseq object, a TreeSummarizedExperiment object, or a matrix of counts.
- assay_name: the name of the assay to extract from the TreeSummarizedExperiment object (default assayName = "counts"). Not used if the input object is a phyloseq.
- verbose: an optional logical value. If TRUE information on the steps of the algorithm is printed. Default verbose = TRUE.

Value

A data frame containing the continuity corrected logarithms of the average fitted values for each row of the ‘counts’ matrix in the ‘Y’ column, and the estimated probability to observe a zero in the ‘Y0’ column.

Examples

```r
# Generate some random counts
counts = matrix(rnbinom(n = 60, size = 3, prob = 0.5), nrow = 10, ncol = 6)
# Fit model on the matrix of counts
NB <- fitNB(counts)
head(NB)
```

Description

Fit a Zero-Inflated Gaussian (ZIG) distribution for each taxon of the count data. The model estimation procedure is performed by metagenomeSeq fitZig function without assuming the presence of any group in the samples (design matrix equal to a column of ones.)

Usage

fitZIG(object, assay_name = "counts", verbose = TRUE)

Arguments

- object: a phyloseq object, a TreeSummarizedExperiment object, or a matrix of counts.
- assay_name: the name of the assay to extract from the TreeSummarizedExperiment object (default assayName = "counts"). Not used if the input object is a phyloseq.
- verbose: an optional logical value. If TRUE information on the steps of the algorithm is printed. Default verbose = TRUE.
**fitZINB**

**Value**

A data frame containing the continuity corrected logarithms of the average fitted values for each row of the matrix of counts in the $Y$ column, and the estimated probability to observe a zero in the $Y0$ column.

**Examples**

```r
# Generate some random counts
counts = matrix(rnbinom(n = 60, size = 3, prob = 0.5), nrow = 10, ncol = 6)

# Fit model on the counts matrix
ZIG <- fitZIG(counts)
head(ZIG)
```

**Description**

Fit a Zero-Inflated Negative Binomial (ZINB) distribution for each taxon of the countdata. The ZINB estimation procedure is performed by zinbwave `zinbFit` function with `commondispersion = FALSE`, regularization parameter `epsilon = 1e10`, and not assuming the presence of any group in the samples (design matrix equal to a column of ones.)

**Usage**

```r
fitZINB(object, assay_name = "counts", verbose = TRUE)
```

**Arguments**

- `object`: a phyloseq object, a TreeSummarizedExperiment object, or a matrix of counts.
- `assay_name`: the name of the assay to extract from the TreeSummarizedExperiment object (default `assayName = "counts"`). Not used if the input object is a phyloseq.
- `verbose`: an optional logical value. If `TRUE` information on the steps of the algorithm is printed. Default `verbose = TRUE`.

**Value**

A data frame containing the continuity corrected logarithms of the average fitted values for each row of the matrix of counts in the $Y$ column, and the estimated probability to observe a zero in the $Y0$ column.
getDA

Examples

# Generate some random counts
counts = matrix(rnbinom(n = 60, size = 3, prob = 0.5), nrow = 10, ncol = 6)

# Fit model on the counts matrix
ZINB <- fitZINB(counts)
head(ZINB)

description

Inspect the list of p-values or/and log fold changes from the output of a differential abundance detection method.

Usage

getDA(
  method,
  slot = "pValMat",
  colName = "rawP",
  type = "pvalue",
  direction = NULL,
  threshold_pvalue = 1,
  threshold_logfc = 0,
  top = NULL,
  verbose = FALSE
)

Arguments

method Output of a differential abundance detection method. pValMat, statInfo matrices, and method’s name must be present (See vignette for detailed information).
slot The slot name where to extract values (default slot = "pValMat").
colName The column name of the slot where to extract values (default colName = "rawP").
type The value type of the column selected where to extract values. Two values are possible: "pvalue" or "logfc" (default type = "pvalue").
direction statInfo’s column name containing information about the signs of differential abundance (usually log fold changes) (default direction = NULL).
threshold_pvalue Threshold value for p-values. If present, features with p-values lower than threshold_pvalue are considered differentially abundant. Set threshold_pvalue = 1 to not filter by p-values.
threshold_logfc
Threshold value for log fold changes. If present, features with log fold change absolute values higher than threshold_logfc are considered differentially abundant. Set threshold_logfc = 0 to not filter by log fold change values.

top
If not null, the top number of features, ordered by p-values or log fold change values, are considered as differentially abundant (default top = NULL).

verbose
Boolean to display the kind of extracted values (default verbose = FALSE).

Value
A data.frame with several columns:
• stat which contains the p-values or the absolute log fold change values;
• direction which is present if method was a data.frame object, it contains the information about directionality of differential abundance (usually log fold changes);
• DA which can contain several values according to thresholds and inputs. "DA" or "non-DA" if method object was a vector, "UP Abundant", "DOWN Abundant", or "non-DA" if method was a data.frame.

See Also
getStatistics, extractDA

Examples

data("ps_plaque_16S")
# Add scaling factors
ps_plaque_16S <- norm_edgeR(object = ps_plaque_16S, method = "TMM")
# DA analysis
da_limma <- DA_limma(
  object = ps_plaque_16S,
  design = ~ 1 + HMP_BODY_SUBSITE,
  coef = 2,
  norm = "TMM"
)
# features with p-value < 0.1 as DA
getDA(
  method = da_limma, slot = "pValMat", colName = "rawP", type = "pvalue",
  direction = NULL, threshold_pvalue = 0.1, threshold_logfc = 0,
  top = NULL
)
# top 10 feature with highest logFC are DA
getDA(
  method = da_limma, slot = "pValMat", colName = "rawP", type = "pvalue",
  direction = "logFC", threshold_pvalue = 1, threshold_logfc = 0, top = 10
)
# features with p-value < 0.1 and |logFC| > 1 are DA
getDA(
  method = da_limma, slot = "pValMat", colName = "rawP", type = "pvalue",
  direction = "logFC", threshold_pvalue = 0.1, threshold_logfc = 1, top = NULL
)
getPositives

# top 10 features with |logFC| > 1 are DA
getDA(
    method = da.limma, slot = "pValMat", colName = "rawP", type = "pvalue",
    direction = "logFC", threshold_pvalue = 1, threshold_logfc = 1, top = 10
)

getPositives

Description

Inspect the list of p-values or/and log fold changes from the output of a differential abundance detection method and count the True Positives (TP) and the False Positives (FP).

Usage

getPositives(method, enrichmentCol, TP, FP)

Arguments

method Output of differential abundance detection method in which DA information is extracted by the getDA function, information related to enrichment is appropriately added through the addKnowledge function and the Fisher exact tests is performed for the contingency tables by the enrichmentTests function.

enrichmentCol name of the column containing information for enrichment analysis.

TP A list of length-2 vectors. The entries in the vector are the direction ("UP Abundant", "DOWN Abundant", or "non-DA") in the first position, and the level of the enrichment variable (enrichmentCol) which is expected in that direction, in the second position.

FP A list of length-2 vectors. The entries in the vector are the direction ("UP Abundant", "DOWN Abundant", or "non-DA") in the first position, and the level of the enrichment variable (enrichmentCol) which is not expected in that direction, in the second position.

Value

A named vector containing the number of TPs and FPs.

See Also

createPositives.
Examples

data("ps_plaque_16S")
data("microbial_metabolism")
# Extract genera from the phyloseq tax_table slot
genera <- phyloseq::tax_table(ps_plaque_16S)[, "GENUS"]
# Genera as rownames of microbial_metabolism data.frame
rownames(microbial_metabolism) <- microbial_metabolism$Genus
# Match OTUs to their metabolism
priorInfo <- data.frame(genera,
    "Type" = microbial_metabolism[genera, "Type"]
)
# Unmatched genera becomes "Unknown"
unknown_metabolism <- is.na(priorInfo$Type)
priorInfo[unknown_metabolism, "Type"] <- "Unknown"
priorInfo$Type <- factor(priorInfo$Type)
# Add a more informative names column
priorInfo[, "newNames"] <- paste0(rownames(priorInfo), priorInfo[, "GENUS"])

# DA Analysis
# Add scaling factors
ps_plaque_16S <- norm_edgeR(object = ps_plaque_16S, method = "TMM")
# DA analysis
da.limma <- DA_limma(
    object = ps_plaque_16S,
    design = ~ 1 + HMP_BODY_SUBSITE,
    coef = 2,
    norm = "TMM"
)

DA <- getDA(
    method = da.limma, slot = "pValMat", colName = "adjP",
    type = "pvalue", direction = "logFC", threshold_pvalue = 0.05,
    threshold_logfc = 1, top = NULL
)
# Add a priori information
DA_info <- addKnowledge(
    method = DA, priorKnowledge = priorInfo,
    enrichmentCol = "Type", namesCol = "newNames"
)
# Create contingency tables and compute F tests
DA_info_enriched <- enrichmentTest(
    method = DA_info, enrichmentCol = "Type",
    alternative = "greater"
)
# Count True and False Positives
DA_TP_FP <- getPositives(
    method = DA_info_enriched, enrichmentCol = "Type",
    TP = list(c("UP Abundant", "Aerobic"), c("DOWN Abundant", "Anaerobic")),
    FP = list(c("UP Abundant", "Anaerobic"), c("DOWN Abundant", "Aerobic"))
)
getStatistics

Description

Extract the list of p-values or/and log fold changes from the output of a differential abundance
detection method.

Usage

getStatistics(
  method,
  slot = "pValMat",
  colName = "rawP",
  type = "pvalue",
  direction = NULL,
  verbose = FALSE
)

Arguments

method  Output of a differential abundance detection method. pValMat, statInfo matrices, and method's name must be present (See vignette for detailed information).
slot    The slot name where to extract values (default slot = "pValMat").
colName The column name of the slot where to extract values (default colName = "rawP").
type    The value type of the column selected where to extract values. Two values are possible: "pvalue" or "logfc" (default type = "pvalue").
direction statInfo's column name containing information about the signs of differential abundance (usually log fold changes) (default direction = NULL).
verbose  Boolean to display the kind of extracted values (default verbose = FALSE).

Value

A vector or a data.frame. If direction = NULL, the colname column values, transformed according to type (not tranformed if type = "pvalue", -abs(value) if type = "logfc"), of the slot are reported, otherwise the direction column of the statInfo matrix is added to the output.

See Also

eextractStatistics
Examples

data("ps_plaque_16S")
# Add scaling factors
ps_plaque_16S <- norm_edgeR(object = ps_plaque_16S, method = "TMM")
# DA analysis
da.limma <- DA_limma(
    object = ps_plaque_16S,
    design = ~ 1 + HMP_BODY_SUBSITE,
    coef = 2,
    norm = "TMM"
)
# get p-values
getStatistics(
    method = da.limma, slot = "pValMat", colName = "rawP",
    type = "pvalue", direction = NULL
)
# get negative abs(logFC) values
getStatistics(
    method = da.limma, slot = "statInfo", colName = "logFC",
    type = "logfc", direction = NULL
)
# get p-values and logFC
getStatistics(
    method = da.limma, slot = "pValMat", colName = "rawP",
    type = "pvalue", direction = "logFC"
)

Description

Check whether the input object is a phyloseq or a TreeSummarizedExperiment, then extract the requested data slots.

Usage

get_counts_metadata(
    object, assay_name = "counts", min_counts = 0, min_samples = 0
)

Arguments

object a phyloseq or TreeSummarizedExperiment object.
assay_name the name of the assay to extract from the TreeSummarizedExperiment object (default assayName = "counts"). Not used if the input object is a phyloseq.
min_counts  Parameter to filter taxa. Set this number to keep features with more than min_counts counts in more than min_samples samples (default min_counts = 0).

min_samples  Parameter to filter taxa. Set this number to keep features with a min_counts counts in more than min_samples samples (default min_samples = 0).

Value

A list of results:

- counts: the otu_table slot or assayName assay of the phyloseq or TreeSummarizedExperiment object;
- metadata: the sample_data or colData slot of the phyloseq or TreeSummarizedExperiment object;
- is_phyloseq: a boolean equal to TRUE if the input is a phyloseq object.

Examples

```r
set.seed(1)
# Create a very simple phyloseq object
counts <- matrix(rnbinom(n = 60, size = 3, prob = 0.5), nrow = 10, ncol = 6)
metadata <- data.frame("Sample" = c("S1", "S2", "S3", "S4", "S5", "S6"),
                       "group" = as.factor(c("A", "A", "A", "B", "B", "B")))
ps <- phyloseq::phyloseq(phyloseq::otu_table(counts, taxa_are_rows = TRUE),
                        phyloseq::sample_data(metadata))
get_counts_metadata(ps)

# Or with a TreeSummarizedExperiment
tse <- TreeSummarizedExperiment::TreeSummarizedExperiment(
                                   assays = list("counts" = counts), colData = metadata)
get_counts_metadata(tse)
```

**iterative_ordering**

**iterativeOrdering**

**Description**

Turn the chosen columns (factor) of the input `data.frame` into ordered factors. For each factor, the order is given by the number of elements in each level of that factor.

**Usage**

`iterative_ordering(df, var_names, i = 1, decreasing = TRUE)`

**Arguments**

- **df**  A `data.frame` object.
- **var_names**  Character vector containing the names of one or more columns of `df`.
- **i**  Iteration index (default `i = 1`).
- **decreasing**  Logical value or a vector of them. Each value should be associated to a `var_name` vector’s element. Should the sort order be increasing or decreasing?
meanDifferences

Value

the input data.frame with the var_names variables as ordered factors.

See Also

plotMutualFindings

Examples

# From a dataset with some factor columns
mpg <- data.frame(ggplot2::mpg)
# Order the levels of the desired factors based on the cardinality of each
# level.
ordered_mpg <- iterative_ordering(df = mpg,
   var_names = c("manufacturer", "model"),
   decreasing = c(TRUE, TRUE))
# Now the levels of the factors are ordered in a decreasing way
levels(ordered_mpg$manufacturer)
levels(ordered_mpg$model)

meanDifferences

Description

Compute the differences between the estimated and the observed continuity corrected logarithms of the average count values (MD), and between the estimated average probability to observe a zero and the the observed zero rate (ZPD).

Usage

meanDifferences(estimated, observed)

Arguments

estimated a two column data.frame, output of fitNB, fitZINB, fitDM, fitZIG, or fitHURDLE functions. More in general, a data frame containing the continuity corrected logarithm for the average of the fitted values for each row of a matrix of counts in the Y column, and the estimated probability to observe a zero in the Y0 column.

observed a two column data.frame, output of prepareObserved function. More in general, a data frame containing the continuity corrected logarithm for the average of the observed values for each row of a matrix of counts in the Y column, and the estimated proportion of zeroes in the Y0 column.

Value

a data.frame containing the differences between the estimated and the observed continuity corrected logarithms of the average count values in the MD column, and between the estimated average probability to observe a zero and the the observed zero rate in the ZPD column.
See Also

prepareObserved.

Examples

# Randomly generate the observed and estimated data.frames
observed <- data.frame(Y = rpois(10, 5), Y0 = runif(10, 0, 1))
estimated <- data.frame(Y = rpois(10, 5), Y0 = runif(10, 0, 1))

# Compute the mean differences between estimated and observed data.frames
meanDifferences(estimated, observed)

---

**microbial_metabolism**  
*(Data)* Microbial metabolism

**Description**

Aerobic, Anaerobic, or Facultative Anaerobic microbes by genus *(NYC-HANES study)*.

**Usage**

data(microbial_metabolism)

**Format**

A data.frame object

**norm_CSS**  
norm_CSS

**Description**

Calculate normalization factors from a phyloseq or TreeSummarizedExperiment object. Inherited from metagenomeSeq *calcNormFactors* function which performs the Cumulative Sum Scaling normalization.

**Usage**

norm_CSS(object, assay_name = "counts", method = "CSS", verbose = TRUE)

**Arguments**

- **object**: a phyloseq or TreeSummarizedExperiment object.
- **assay_name**: the name of the assay to extract from the TreeSummarizedExperiment object (default assayName = "counts"). Not used if the input object is a phyloseq.
- **method**: normalization method to be used (only CSS).
- **verbose**: an optional logical value. If TRUE, information about the steps of the algorithm is printed. Default verbose = TRUE.
A new column containing the CSS normalization factors is added to the sample_data slot of the phyloseq object or the colData slot of the TreeSummarizedExperiment object.

See Also

calcNormFactors for details. setNormalizations and runNormalizations to fastly set and run normalizations.

Examples

```r
set.seed(1)
# Create a very simple phyloseq object
counts <- matrix(rnbinom(n = 60, size = 3, prob = 0.5), nrow = 10, ncol = 6)
ps <- phyloseq::phyloseq(phyloseq::otu_table(counts, taxa_are_rows = TRUE), phyloseq::sample_data(metadata))

# Calculate the normalization factors
ps_NF <- norm_CSS(object = ps, method = "CSS")
# The phyloseq object now contains the normalization factors:
CSSFacts <- phyloseq::sample_data(ps_NF)[, "NF.CSS"]
head(CSSFacts)

# VERY IMPORTANT: metagenomeSeq uses scaling factors to normalize counts
# (even though they are called normalization factors). These factors are used
# internally by a regression model. To make CSS size factors available for
# edgeR, we need to transform them into normalization factors. This is
# possible by dividing the factors for the library sizes and renormalizing.
normCSSFacts = CSSFacts / colSums(phyloseq::otu_table(ps_stool_16S))
# Renormalize: multiply to 1
normFacts = normCSSFacts/exp(colMeans(log(normCSSFacts)))
```

Description

Calculate size factors from a phyloseq or TreeSummarizedExperiment object. Inherited from DESeq2 estimateSizeFactors function.

Usage

```r
norm_DESeq2(
  object,
  assay_name = "counts",
  method = c("ratio", "poscounts", "iterate"),
  verbose = TRUE,
)```
norm_DESeq2

Arguments

- **object**: a phyloseq or TreeSummarizedExperiment object.
- **assay_name**: the name of the assay to extract from the TreeSummarizedExperiment object (default assayName = "counts"). Not used if the input object is a phyloseq.
- **method**: Method for estimation: either "ratio", "poscounts", or "iterate". "ratio" uses the standard median ratio method introduced in DESeq. The size factor is the median ratio of the sample over a "pseudosample": for each gene, the geometric mean of all samples. "poscounts" and "iterate" offer alternative estimators, which can be used even when all features contain a sample with a zero (a problem for the default method, as the geometric mean becomes zero, and the ratio undefined). The "poscounts" estimator deals with a feature with some zeros, by calculating a modified geometric mean by taking the n-th root of the product of the non-zero counts. This evolved out of use cases with Paul McMurdie’s phyloseq package for metagenomic samples. The "iterate" estimator iterates between estimating the dispersion with a design of ~1, and finding a size factor vector by numerically optimizing the likelihood of the ~1 model.
- **verbose**: an optional logical value. If TRUE, information about the steps of the algorithm is printed. Default verbose = TRUE.
- **...**: other parameters for DESeq2 estimateSizeFactors function.

Value

A new column containing the chosen DESeq2-based size factors is added to the sample_data slot of the phyloseq object or the colData slot of the TreeSummarizedExperiment object.

See Also

estimateSizeFactors for details. setNormalizations and runNormalizations to fastly set and run normalizations.

Examples

```r
set.seed(1)
# Create a very simple phyloseq object
counts <- matrix(rbinom(n = 60, size = 3, prob = 0.5), nrow = 10, ncol = 6)
metadata <- data.frame("Sample" = c("S1", "S2", "S3", "S4", "S5", "S6"),
  "group" = as.factor(c("A", "A", "A", "B", "B", "B")))
ps <- phyloseq::phyloseq(phyloseq::otu_table(counts, taxa_are_rows = TRUE),
  phyloseq::sample_data(metadata))

# Calculate the size factors
ps_NF <- norm_DESeq2(object = ps, method = "poscounts")
# The phyloseq object now contains the size factors:
sizeFacts <- phyloseq::sample_data(ps_NF)[, "NF.poscounts"]
head(sizeFacts)
```
# VERY IMPORTANT: DESeq2 uses size factors to normalize counts.
# These factors are used internally by a regression model. To make DEseq2
# size factors available for edgeR, we need to transform them into
# normalization factors. This is possible by dividing the factors by the
# library sizes and renormalizing.

normSizeFacts = sizeFacts / colSums(phyloseq::otu_table(ps_stool_16S))
# Renormalize: multiply to 1
normFacts = normSizeFacts/exp(colMeans(log(normSizeFacts)))

---

### Description

Calculate normalization factors from a phyloseq or TreeSummarizedExperiment object. Inherited from edgeR `calcNormFactors` function.

### Usage

```r
norm_edgeR(
  object,
  assay_name = "counts",
  method = c("TMM", "TMMwsp", "RLE", "upperquartile", "posupperquartile", "none"),
  refColumn = NULL,
  logratioTrim = 0.3,
  sumTrim = 0.05,
  doWeighting = TRUE,
  Acutoff = -1e+10,
  p = 0.75,
  verbose = TRUE,
  ...
)
```

### Arguments

- **object**: a phyloseq or TreeSummarizedExperiment object.
- **assay_name**: the name of the assay to extract from the TreeSummarizedExperiment object (default `assayName = "counts"`). Not used if the input object is a phyloseq.
- **method**: normalization method to be used. Choose between TMM, TMMwsp, RLE, upperquartile, posupperquartile or none.
- **refColumn**: column to use as reference for `method="TMM"`. Can be a column number or a numeric vector of length `nrow(object)`.
- **logratioTrim**: the fraction (0 to 0.5) of observations to be trimmed from each tail of the distribution of log-ratios (M-values) before computing the mean. Used by `method="TMM"` for each pair of samples.
sumTrim
the fraction (0 to 0.5) of observations to be trimmed from each tail of the distribution of A-values before computing the mean. Used by method="TMM" for each pair of samples.

doWeighting
logical, whether to use (asymptotic binomial precision) weights when computing the mean M-values. Used by method="TMM" for each pair of samples.

Acutoff
minimum cutoff applied to A-values. Count pairs with lower A-values are ignored. Used by method="TMM" for each pair of samples.

p
numeric value between 0 and 1 specifying which quantile of the counts should be used by method="upperquartile".

verbose
an optional logical value. If TRUE, information about the steps of the algorithm is printed. Default verbose = TRUE.

... other arguments are not currently used.

Value
A new column containing the chosen edgeR-based normalization factors is added to the sample_data slot of the phyloseq object or the colData slot of the TreeSummarizedExperiment object.

See Also
calcNormFactors for details.
setNormalizations and runNormalizations to fastly set and run normalizations.

Examples
set.seed(1)
# Create a very simple phyloseq object
counts <- matrix(rnbinom(n = 60, size = 3, prob = 0.5), nrow = 10, ncol = 6)
metadata <- data.frame("Sample" = c("S1", "S2", "S3", "S4", "S5", "S6"),
                        "group" = as.factor(c("A", "A", "A", "B", "B", "B")))
ps <- phyloseq::phyloseq(phyloseq::otu_table(counts, taxa_are_rows = TRUE),
                        phyloseq::sample_data(metadata))

# Calculate the normalization factors
ps_NF <- norm_edgeR(object = ps, method = "TMM")

# The phyloseq object now contains the normalization factors:
normFacts <- phyloseq::sample_data(ps_NF)[, "NF.TMM"]
head(normFacts)

# VERY IMPORTANT: edgeR uses normalization factors to normalize library sizes not counts. They are used internally by a regression model. To make edgeR normalization factors available for other methods, such as DESeq2 or other DA methods based on scaling or size factors, we need to transform them into size factors. This is possible by multiplying the factors for the library sizes and renormalizing.
normLibSize = normFacts * colSums(phyloseq::otu_table(ps_stool_16S))
# Renormalize: multiply to 1
sizeFacts = normLibSize/exp(colMeans(log(normLibSize)))
Description

Calculate the Total Sum Scaling (TSS) factors for a phyloseq or a TreeSummarizedExperiment object, i.e. the library sizes. If the counts are divided by the scaling factors, a relative abundance is obtained.

Usage

`norm_TSS(object, assay_name = "counts", method = "TSS", verbose = TRUE)`

Arguments

- `object`: a phyloseq or TreeSummarizedExperiment object.
- `assay_name`: the name of the assay to extract from the TreeSummarizedExperiment object (default `assayName = "counts"`). Not used if the input object is a phyloseq.
- `method`: normalization method to be used.
- `verbose`: an optional logical value. If `TRUE`, information about the steps of the algorithm is printed. Default `verbose = TRUE`.

Value

A new column containing the TSS scaling factors is added to the `sample_data` slot of the phyloseq object or the `colData` slot of the TreeSummarizedExperiment object.

See Also

`setNormalizations` and `runNormalizations` to fastly set and run normalizations.

Examples

```r
set.seed(1)
# Create a very simple phyloseq object
counts <- matrix(rnbinom(n = 60, size = 3, prob = 0.5), nrow = 10, ncol = 6)
metadata <- data.frame("Sample" = c("S1", "S2", "S3", "S4", "S5", "S6"),
                      "group" = as.factor(c("A", "A", "A", "B", "B", "B")))
ps <- phyloseq::phyloseq(phyloseq::otu_table(counts, taxa_are_rows = TRUE),
                        phyloseq::sample_data(metadata))

# Calculate the scaling factors
ps_NF <- norm_TSS(object = ps)
# The phyloseq object now contains the scaling factors:
scaleFacts <- phyloseq::sample_data(ps_NF)[, "NF.TSS"]
head(scaleFacts)
```
plotConcordance

Description

Produce a list of graphical outputs summarizing the between and within method concordance.

Usage

plotConcordance(concordance, threshold = NULL, cols = NULL)

Arguments

concordance A long format data.frame produced by createConcordance function.
threshold The threshold for rank (x-axis upper limit if all methods have a higher number of computed statistics).
cols A named vector containing the color hex codes.

Value

A 2 elements list of ggplot2 class objects:

- concordanceDendrogram which contains the vertically directioned dendrogram for the methods involved in the concordance analysis;
- concordanceHeatmap which contains the heatmap of between and within method concordances.

See Also

createConcordance

Examples

data(ps_plaque_16S)

# Balanced design
my_splits <- createSplits(
  object = ps_plaque_16S, varName = "HMP_BODY_SUBSITE", balanced = TRUE,
  paired = "RSID", N = 10 # N = 100 suggested
)

# Make sure the subject ID variable is a factor
phyloseq::sample_data(ps_plaque_16S)[, "RSID"] <- as.factor(
  phyloseq::sample_data(ps_plaque_16S)[["RSID"]]
)

# Initialize some limma based methods
my_limma <- set_limma(design = ~ RSID + HMP_BODY_SUBSITE,
  coef = "HMP_BODY_SUBSITESupragingival Plaque",
  etc.
norm = c("TMM", "CSS"))

# Set the normalization methods according to the DA methods
my_norm <- setNormalizations(fun = c("norm_edgeR", "norm_CSS"),
    method = c("TMM", "CSS"))

# Run methods on split datasets
results <- runSplits(split_list = my_splits, method_list = my_limma,
    normalization_list = my_norm, object = ps_plaque_16S)

# Concordance for p-values
concordance_pvalues <- createConcordance(
    object = results, slot = "pValMat", colName = "rawP", type = "pvalue"
)

# plot concordances from rank 1 to 50.
plotConcordance(
    concordance = concordance_pvalues,
    threshold = 50
)

---

**plotConcordanceDendrogram**

**plotConcordanceDendrogram**

**Description**

Plots the method’s dendrogram of concordances.

**Usage**

```
plotConcordanceDendrogram(hc, direction = "v", cols)
```

**Arguments**

- `hc`: Hierarchical clustering results produced in `plotConcordance` function.
- `direction`: vertical (default direction = "v") or horizontal (direction = "h").
- `cols`: A named vector containing the color hex codes.

**Value**

A ggplot2 object

**See Also**

`createConcordance` and `plotConcordance`
Description

Plots the heatmap of concordances.

Usage

plotConcordanceHeatmap(c_df, threshold, cols)

Arguments

c_df A simplified concordance data frame produced in plotConcordance function.
threshold The threshold for rank (x-axis upper limit if all methods have a higher number of computed statistics).
cols A named vector containing the color hex codes.

Value

a ggplot2 object

See Also

createConcordance and plotConcordance

Description

Plot of the contingency tables for the chosen method. The top-left cells are colored, according to Fisher exact tests’ p-values, if the number of features in those cells are enriched.

Usage

plotContingency(enrichment, method, levels_to_plot)

Arguments

enrichment enrichment object produced by createEnrichment function.
method name of the method to plot.
levels_to_plot A character vector containing the levels of the enrichment variable to plot.
Value

a ggplot2 object.

See Also

createEnrichment, plotEnrichment, and plotMutualFindings.

Examples

data("ps_plaque_16S")
data("microbial_metabolism")

# Extract genera from the phyloseq tax_table slot
genera <- phyloseq::tax_table(ps_plaque_16S)[, "GENUS"]
# Genera as rownames of microbial_metabolism data.frame
rownames(microbial_metabolism) <- microbial_metabolism$Genus
# Match OTUs to their metabolism
priorInfo <- data.frame(genera,
   "Type" = microbial_metabolism[genera, "Type"])
# Unmatched genera becomes "Unknown"
unknown_metabolism <- is.na(priorInfo$"Type")
priorInfo[unknown_metabolism, "Type"] <- "Unknown"
priorInfo$Type <- factor(priorInfo$"Type")
# Add a more informative names column
priorInfo[, "newNames"] <- paste0(rownames(priorInfo), priorInfo[, "GENUS"])

# Add some normalization/scaling factors to the phyloseq object
my_norm <- setNormalizations(fun = c("norm_edgeR", "norm_CSS"),
   method = c("TMM", "CSS"))
ps_plaque_16S <- runNormalizations(normalization_list = my_norm,
   object = ps_plaque_16S)

# Initialize some limma based methods
my_limma <- set_limma(design = ~ 1 + RSID + HMP_BODY_SUBSITE,
   coef = "HMP_BODY_SUBSITEsupragingival Plaque",
   norm = c("TMM", "CSS"))

# Make sure the subject ID variable is a factor
phyloseq::sample_data(ps_plaque_16S)[, "RSID"] <- as.factor(
   phyloseq::sample_data(ps_plaque_16S)[["RSID"]])

# Perform DA analysis
Plaque_16S_DA <- runDA(method_list = my_limma, object = ps_plaque_16S)

# Enrichment analysis
enrichment <- createEnrichment(object = Plaque_16S_DA,
   priorKnowledge = priorInfo, enrichmentCol = "Type", namesCol = "GENUS",
   slot = "pValMat", colName = "adjP", type = "pvalue", direction = "logFC",
   threshold_pvalue = 0.1, threshold_logfc = 1, top = 10, verbose = TRUE)

# Contingency tables
plotContingency(enrichment = enrichment, method = "limma.TMM")
Description

Summary plot for the number of differentially abundant (DA) features and their association with enrichment variable. If some features are UP-Abundant or DOWN-Abundant (or just DA), several bars will be represented in the corresponding direction. Significance thresholds are reported over/under each bar, according to the Fisher exact tests.

Usage

plotEnrichment(enrichment, enrichmentCol, levels_to_plot)

Arguments

enrichment enrichment object produced by createEnrichment function.
enrichmentCol name of the column containing information for enrichment analysis.
levels_to_plot A character vector containing the levels of the enrichment variable to plot.

Value

a ggplot2 object.

See Also

createEnrichment, plotContingency, and plotMutualFindings.

Examples

data("ps_plaque_16S")
data("microbial_metabolism")

# Extract genera from the phyloseq tax_table slot
genera <- phyloseq::tax_table(ps_plaque_16S)[, "GENUS"]
# Genera as rownames of microbial_metabolism data.frame
rownames(microbial_metabolism) <- microbial_metabolism$Genus
# Match OTUs to their metabolism
priorInfo <- data.frame(genera,
    "Type" = microbial_metabolism[genera, "Type"]
)  
# Unmatched genera becomes "Unknown"
unknown_metabolism <- is.na(priorInfo$Type)
priorInfo[unknown_metabolism, "Type"] <- "Unknown"
priorInfo$Type <- factor(priorInfo$Type)
# Add a more informative names column
priorInfo[, "newNames"] <- paste0(rownames(priorInfo), priorInfo[, "GENUS"])

# Add some normalization/scaling factors to the phyloseq object
my_norm <- setNormalizations(fun = c("norm_edgeR", "norm_CSS"),
method = c("TMM", "CSS"))
ps_plaque_16S <- runNormalizations(normalization_list = my_norm,
object = ps_plaque_16S)

# Initialize some limma based methods
my_limma <- set_limma(design = ~ 1 + RSID + HMP_BODY_SUBSITE,
coef = "HMP_BODY_SUBSITESupragingival Plaque",
norm = c("TMM", "CSS"))

# Make sure the subject ID variable is a factor
phyloseq::sample_data(ps_plaque_16S)[,"RSID"] <- as.factor(
  phyloseq::sample_data(ps_plaque_16S)[["RSID"]])

# Perform DA analysis
Plaque_16S_DA <- runDA(method_list = my_limma, object = ps_plaque_16S)

# Enrichment analysis
enrichment <- createEnrichment(object = Plaque_16S_DA,
priorKnowledge = priorInfo, enrichmentCol = "Type", namesCol = "GENUS",
slot = "pValMat", colName = "adjP", type = "pvalue", direction = "logFC",
threshold_pvalue = 0.1, threshold_logfc = 1, top = 10, verbose = TRUE)

# Contingency tables
plotContingency(enrichment = enrichment, method = "limma.TMM")
# Barplots
plotEnrichment(enrichment, enrichmentCol = "Type")
# Mutual findings
plotMutualFindings(
enrichment = enrichment, enrichmentCol = "Type",
n_methods = 1
)

plotFDR(df_FDR, cols = NULL)

Description

Draw the nominal false discovery rates for the 0.01, 0.05, and 0.1 levels.

Usage

plotFDR(df_FDR, cols = NULL)
Arguments

- `df_FDR`: a data.frame produced by the `createTIEC` function, containing the FDR values.
- `cols`: named vector of colors.

Value

A ggplot object.

Examples

# Load some data
data(ps_stool_16S)

# Generate the patterns for 10 mock comparison for an experiment
# (N = 1000 is suggested)
mocks <- createMocks(nsamples = phyloseq::nsamples(ps_stool_16S), N = 10)
head(mocks)

# Add some normalization/scaling factors to the phyloseq object
my_norm <- setNormalizations(fun = c("norm_edgeR", "norm_CSS"),
                              method = c("TMM", "CSS"))
ps_stool_16S <- runNormalizations(normalization_list = my_norm,
                                 object = ps_stool_16S)

# Initialize some limma based methods
my_limma <- set_limma(design = ~ group, coef = 2,
                      norm = c("TMM", "CSS"))

# Run methods on mock datasets
results <- runMocks(mocks = mocks, method_list = my_limma,
                     object = ps_stool_16S)

# Prepare results for Type I Error Control
TIEC_summary <- createTIEC(results)

# Plot the results
plotFPR(df_FPR = TIEC_summary$df_FPR)
plotFDR(df_FDR = TIEC_summary$df_FDR)
plotQQ(df_QQ = TIEC_summary$df_QQ, zoom = c(0, 0.1))
plotKS(df_KS = TIEC_summary$df_KS)
plotLogP(df_QQ = TIEC_summary$df_QQ)

Description

Draw the boxplots of the proportions of p-values lower than 0.01, 0.05, and 0.1 thresholds for each method.
Usage

plotFPR(df_FPR, cols = NULL)

Arguments

df_FPR  a data.frame produced by the createTIEC function, containing the FPR values.
cols    named vector of colors.

Value

A ggplot object.

Examples

# Load some data
data(ps_stool_16S)

# Generate the patterns for 10 mock comparison for an experiment
# (N = 1000 is suggested)
mocks <- createMocks(nsamples = phyloseq::nsamples(ps_stool_16S), N = 10)
head(mocks)

# Add some normalization/scaling factors to the phyloseq object
my_norm <- setNormalizations(fun = c("norm_edgeR", "norm_CSS"),
                              method = c("TMM", "CSS"))
ps_stool_16S <- runNormalizations(normalization_list = my_norm,
                                  object = ps_stool_16S)

# Initialize some limma based methods
my_limma <- set_limma(design = ~ group, coef = 2,
                       norm = c("TMM", "CSS"))

# Run methods on mock datasets
results <- runMocks(mocks = mocks, method_list = my_limma,
                     object = ps_stool_16S)

# Prepare results for Type I Error Control
TIEC_summary <- createTIEC(results)

# Plot the results
plotFPR(df_FPR = TIEC_summary$df_FPR)
plotFDR(df_FDR = TIEC_summary$df_FDR)
plotQQ(df_QQ = TIEC_summary$df_QQ, zoom = c(0, 0.1))
plotKS(df_KS = TIEC_summary$df_KS)
plotLogP(df_QQ = TIEC_summary$df_QQ)
Description

Draw the boxplots of the Kolmogorov-Smirnov test statistics for the p-value distributions across the mock comparisons.

Usage

plotKS(df_KS, cols = NULL)

Arguments

df_KS a data.frame produced by the createTIEC function containing the KS statistics and their p-values.
cols named vector of colors.

Value

A ggplot object.

Examples

# Load some data
data(ps_stool_16S)

# Generate the patterns for 10 mock comparison for an experiment
# (N = 1000 is suggested)
mocks <- createMocks(nsamples = phyloseq::nsamples(ps_stool_16S), N = 10)
head(mocks)

# Add some normalization/scaling factors to the phyloseq object
my_norm <- setNormalizations(fun = c("norm_edgeR", "norm_CSS"),
method = c("TMM", "CSS"))
ps_stool_16S <- runNormalizations(normalization_list = my_norm,
object = ps_stool_16S)

# Initialize some limma based methods
my_limma <- set_limma(design = ~ group, coef = 2,
norm = c("TMM", "CSS"))

# Run methods on mock datasets
results <- runMocks(mocks = mocks, method_list = my_limma,
object = ps_stool_16S)

# Prepare results for Type I Error Control
TIEC_summary <- createTIEC(results)
# Plot the results
plotFPR(df_FPR = TIEC_summary$df_FPR)
plotFDR(df_FDR = TIEC_summary$df_FDR)
plotQQ(df_QQ = TIEC_summary$df_QQ, zoom = c(0, 0.1))
plotKS(df_KS = TIEC_summary$df_KS)
plotLogP(df_QQ = TIEC_summary$df_QQ)

---

**plotLogP**

**Description**

Draw the p-values or the average p-values distribution across the mock comparisons in logarithmic scale.

**Usage**

```r
plotLogP(df_pval = NULL, df_QQ = NULL, cols = NULL)
```

**Arguments**

- `df_pval`: a data.frame produced by the `createTIEC` function, containing the p-values for each taxon, method, and mock comparison. It is used to draw the negative log10 p-values distribution. If `df_pval` is supplied, let `df_QQ = NULL`.
- `df_QQ`: a data.frame produced by the `createTIEC` function, containing the average p-values for each quantile and method. It is used to draw the negative log10 average p-values distribution. If `df_QQ` is supplied, let `df_pval = NULL`.
- `cols`: named vector of colors.

**Value**

A ggplot object.

**Examples**

```r
# Load some data
data(ps_stool_16S)

# Generate the patterns for 10 mock comparison for an experiment
# (N = 1000 is suggested)
mocks <- createMocks(nsamples = phyloseq::nsamples(ps_stool_16S), N = 10)
head(mocks)

# Add some normalization/scaling factors to the phylloseq object
my_norm <- setNormalizations(fun = c("norm_edgeR", "norm_CSS"),
method = c("TMM", "CSS"))
ps_stool_16S <- runNormalizations(normalization_list = my_norm,
object = ps_stool_16S)
```
# Initialize some limma based methods
my_limma <- set_limma(design = ~ group, coef = 2,
                   norm = c("TMM", "CSS"))

# Run methods on mock datasets
results <- runMocks(mocks = mocks, method_list = my_limma,
                   object = ps_stool_16S)

# Prepare results for Type I Error Control
TIEC_summary <- createTIEC(results)

# Plot the results
plotFPR(df_FPR = TIEC_summary$df_FPR)
plotFDR(df_FDR = TIEC_summary$df_FDR)
plotQQ(df_QQ = TIEC_summary$df_QQ, zoom = c(0, 0.1))
plotKS(df_KS = TIEC_summary$df_KS)
plotLogP(df_QQ = TIEC_summary$df_QQ)

---

plotMD

Description

A function to plot mean difference (MD) and zero probability difference (ZPD) values between estimated and observed values.

Usage

plotMD(data, difference = NULL, split = TRUE)

Arguments

data a list, output of the fitModels function. Each element of the list is a ‘data.frame’ object with Model, Y, Y0, MD, and ZPD columns containing the model name, the observed values for the mean and the zero proportion and the differences between observed and estimated values.

difference character vector, either MD or ZPD to plot the differences between estimated and observed mean counts or the differences between estimated zero probability and observed zero proportion.

split Display each model mean differences in different facets (default split = TRUE). If FALSE, points are not displayed for more clear representation.

Value

a ggplot object.
See Also

`fitModels` and `RMSE` for the model estimations and the RMSE computations respectively. `plotRMSE` for the graphical evaluation of the RMSE values.

Examples

```r
# Generate some random counts
counts = matrix(rnbinom(n = 60, size = 3, prob = 0.5), nrow = 10, ncol = 6)

# Estimate the counts assuming several distributions
GOF <- fitModels(
  object = counts, models = c("NB", "ZINB", "DM", "ZIG", "HURDLE"),
  scale_HURDLE = c("median", "default")
)

# Plot the results
plotMD(data = GOF, difference = "MD", split = TRUE)
plotMD(data = GOF, difference = "ZPD", split = TRUE)
```

Description

Plot and filter the features which are considered differentially abundant, simultaneously, by a specified number of methods.

Usage

```r
plotMutualFindings(enrichment, enrichmentCol, levels_to_plot, n_methods = 1)
```

Arguments

- `enrichment`: enrichment object produced by `createEnrichment` function.
- `enrichmentCol`: name of the column containing information for enrichment analysis.
- `levels_to_plot`: A character vector containing the levels of the enrichment variable to plot.
- `n_methods`: minimum number of method that mutually find the features.

Value

A `ggplot2` object.

See Also

`createEnrichment`, `plotEnrichment`, and `plotContingency`.
Examples

data("ps_plaque_16S")
data("microbial_metabolism")

# Extract genera from the phyloseq tax_table slot
genera <- phyloseq::tax_table(ps_plaque_16S)[, "GENUS"]

# Genera as rownames of microbial_metabolism data.frame
rownames(microbial_metabolism) <- microbial_metabolism$Genus

# Match OTUs to their metabolism
priorInfo <- data.frame(genera, 
    "Type" = microbial_metabolism[genera, "Type"])

# Unmatched genera becomes "Unknown"
unknown_metabolism <- is.na(priorInfo$Type)
priorInfo[unknown_metabolism, "Type"] <- "Unknown"
priorInfo$Type <- factor(priorInfo$Type)

# Add a more informative names column
priorInfo[, "newNames"] <- paste0(rownames(priorInfo), priorInfo[, "GENUS"])

# Add some normalization/scaling factors to the phyloseq object
my_norm <- setNormalizations(fun = c("norm_edgeR", "norm_CSS"), 
    method = c("TMM", "CSS"))
ps_plaque_16S <- runNormalizations(normalization_list = my_norm, 
    object = ps_plaque_16S)

# Initialize some limma based methods
my_limma <- set_limma(design = ~ 1 + RSID + HMP_BODY_SUBSITE, 
    coef = "HMP_BODY_SUBSITESupragingival Plaque", 
    norm = c("TMM", "CSS"))

# Make sure the subject ID variable is a factor
phyloseq::sample_data(ps_plaque_16S)[, "RSID"] <- as.factor( 
    phyloseq::sample_data(ps_plaque_16S)[["RSID"]])

# Perform DA analysis
Plaque_16S_DA <- runDA(method_list = my_limma, object = ps_plaque_16S)

# Enrichment analysis
enrichment <- createEnrichment(object = Plaque_16S_DA, 
    priorKnowledge = priorInfo, enrichmentCol = "Type", namesCol = "GENUS", 
    slot = "pValMat", colName = "adjP", type = "pvalue", direction = "logFC", 
    threshold_pvalue = 0.1, threshold_logfc = 1, top = 10, verbose = TRUE)

# Contingency tables
plotContingency(enrichment = enrichment, method = "limma.TMM")

# Barplots
plotEnrichment(enrichment, enrichmentCol = "Type")

# Mutual findings
plotMutualFindings( 
    enrichment = enrichment, enrichmentCol = "Type", 
    n_methods = 1)
)
Description

Plot the difference between the number of true positives (TP) and false positives (FP) for each method and for each 'top' threshold provided by the createPositives() function.

Usage

plotPositives(positives, cols = NULL)

Arguments

positives data.frame object produced by createPositives() function.

cols named vector of cols (default cols = NULL).

Value

a ggplot2 object.

See Also

getPositives, createPositives.

Examples

data("ps_plaque_16S")
data("microbial_metabolism")

# Extract genera from the phyloseq tax_table slot
genera <- phylloseq::tax_table(ps_plaque_16S)[, "GENUS"]
# Genera as rownames of microbial_metabolism data.frame
rownames(microbial_metabolism) <- microbial_metabolism$Genus
# Match OTUs to their metabolism
priorInfo <- data.frame(genera, "Type" = microbial_metabolism[genera, "Type"])
# Unmatched genera becomes "Unknown"
unknown_metabolism <- is.na(priorInfo$Type)
priorInfo[unknown_metabolism, "Type"] <- "Unknown"
priorInfo$Type <- factor(priorInfo$Type)
# Add a more informative names column
priorInfo[, "newNames"] <- paste0(rownames(priorInfo), priorInfo[, "GENUS"])

# Add some normalization/scaling factors to the phyloseq object
my_norm <- setNormalizations(fun = c("norm_edgeR", "norm_CSS"),
                            method = c("TMM", "CSS"))
ps_plaque_16S <- runNormalizations(normalization_list = my_norm,
                                    object = ps_plaque_16S)
# Initialize some limma based methods
my_limma <- set_limma(design = ~ 1 + RSID + HMP_BODY_SUBSITE,
  coef = "HMP_BODY_SUBSITE_supragingival Plaque",
  norm = c("TMM", "CSS"))

# Make sure the subject ID variable is a factor
phyloseq::sample_data(ps_plaque_16S)[, "RSID"] <- as.factor(phyloseq::sample_data(ps_plaque_16S)[["RSID"]])

# Perform DA analysis
Plaque_16S_DA <- runDA(method_list = my_limma, object = ps_plaque_16S)

# Count TPs and FPs, from the top 1 to the top 20 features.
# As direction is supplied, features are ordered by "logFC" absolute values.
positives <- createPositives(object = Plaque_16S_DA,
  priorKnowledge = priorInfo, enrichmentCol = "Type",
  namesCol = "newNames", slot = "pValMat", colName = "rawP",
  type = "pvalue", direction = "logFC", threshold_pvalue = 1,
  threshold_logfc = 0, top = 1:20, alternative = "greater",
  verbose = FALSE,
  TP = list(c("DOWN Abundant", "Anaerobic"), c("UP Abundant", "Aerobic")),
  FP = list(c("DOWN Abundant", "Aerobic"), c("UP Abundant", "Anaerobic")))

# Plot the TP-FP differences for each threshold
plotPositives(positives = positives)

---

**plotQQ**

**plotQQ**

---

**Description**

Draw the average QQ-plots across the mock comparisons.

**Usage**

```r
plotQQ(df_QQ, cols = NULL, zoom = c(0, 0.1), split = FALSE)
```

**Arguments**

- `df_QQ` Coordinates to draw the QQ-plot to compare the mean observed p-value distribution across comparisons, with the theoretical uniform distribution.
- `cols` named vector of colors.
- `zoom` 2-dimesional vector containing the starting and the final coordinates (default: `c(0, 0.1)`)
- `split` boolean value. If TRUE, the qq-plots are reported separately for each method (default `split = FALSE`). Setting it to TRUE is hardly suggested when the number of methods is high or when their colors are similar.
plotRMSE

Value

A ggplot object.

Examples

```r
# Load some data
data(ps_stool_16S)

# Generate the patterns for 10 mock comparison for an experiment
# (N = 1000 is suggested)
mocks <- createMocks(nsamples = phyloseq::nsamples(ps_stool_16S), N = 10)
head(mocks)

# Add some normalization/scaling factors to the phyloseq object
my_norm <- setNormalizations(fun = c("norm_edgeR", "norm_CSS"),
   method = c("TMM", "CSS"))
ps_stool_16S <- runNormalizations(normalization_list = my_norm,
   object = ps_stool_16S)

# Initialize some limma based methods
my_limma <- set_limma(design = ~ group, coef = 2,
   norm = c("TMM", "CSS"))

# Run methods on mock datasets
results <- runMocks(mocks = mocks, method_list = my_limma,
   object = ps_stool_16S)

# Prepare results for Type I Error Control
TIEC_summary <- createTIEC(results)

# Plot the results
plotFPR(df_FPR = TIEC_summary$df_FPR)
plotFDR(df_FDR = TIEC_summary$df_FDR)
plotQQ(df_QQ = TIEC_summary$df_QQ, zoom = c(0, 0.1))
plotKS(df_KS = TIEC_summary$df_KS)
plotLogP(df_QQ = TIEC_summary$df_QQ)
```

Description

A function to plot RMSE values computed for mean difference (MD) and zero probability difference (ZPD) values between estimated and observed values.

Usage

```r
plotRMSE(data, difference = NULL, plotIt = TRUE)
```
prepareObserved

Arguments

- **data**: a list, output of the `fitModels` function. Each element of the list is a ‘data.frame’ object with Model, Y, Y0, MD, and ZPD columns containing the model name, the observed values for the mean and the zero proportion and the differences between observed and estimated values.

- **difference**: character vector, either MD or ZPD to plot the differences between estimated and observed mean counts or the differences between estimated zero probability and observed zero proportion.

- **plotIt**: logical. Should plotting be done? (default `plotIt = TRUE`)

Value

- a ggplot object.

See Also

- `fitModels` and `RMSE` for the model estimations and the RMSE computations respectively. `plotMD` for the graphical evaluation.

Examples

```r
# Generate some random counts
counts = matrix(rnbinom(n = 60, size = 3, prob = 0.5), nrow = 10, ncol = 6)

# Estimate the counts assuming several distributions
GOF <- fitModels(
  object = counts, models = c(
    "NB", "ZINB",
    "DM", "ZIG", "HURDLE"
  ), scale_HURDLE = c("median", "default")
)

# Plot the RMSE results
plotRMSE(data = GOF, difference = "MD")
plotRMSE(data = GOF, difference = "ZPD")
```

Description

Continuity corrected logarithms of the average counts and fraction of zeroes by feature.

Usage

```r
prepareObserved(object, assay_name = "counts", scale = NULL)
```
Arguments

object a phyloseq object, a TreeSummarizedExperiment object, or a matrix of counts.
assay_name the name of the assay to extract from the TreeSummarizedExperiment object (default assayName = "counts"). Not used if the input object is a phyloseq.
scale If specified it refers to the character vector used in fitHURDLE function. Either median or default to choose between the median library size or one million as scaling factors for raw counts.

Value

A data frame containing the continuity corrected logarithm for the raw count mean values for each taxon of the matrix of counts in the Y column and the observed zero rate in the Y0 column. If scale is specified the continuity corrected logarithm for the mean CPM (scale = "default") or the mean counts per median library size (scale = "median") is computed instead.

See Also

meanDifferences

Examples

# Generate some random counts
counts <- matrix(rnbinom(n = 60, size = 3, prob = 0.5), nrow = 10, ncol = 6)

observed1 <- prepareObserved(counts)
# For the comparison with HURDLE model
observed2 <- prepareObserved(counts, scale = "median")

Description

A demonstrative purpose dataset containing microbial abundances for a total of 88 OTUs. The 60 Gingival Plaque paired samples belong to the Human Microbiome Project. This particular subset contains 30 Supragingival and 30 Subgingival Plaque samples from the SEX = "Male", RUN_CENTER = "WUCG", and VISITNO = "1" samples. It is possible to obtain the same dataset after basic filters (remove taxa with zero counts) and collapsing the counts to the genus level; HMP16SData Bioconductor package was used to download the data.

Usage

data(ps_plaque_16S)

Format

An object of class phyloseq
ps_stool_16S

(Data) 33 Stool samples of 16S rRNA (HMP 2012)

Description

A demonstrative purpose dataset containing microbial abundances for a total of 71 OTUs. The 32 Stool samples belong to the Human Microbiome Project. This particular subset contains the SEX = "Male", RUN_CENTER = "BI", and VISITNO = "1" samples. It is possible to obtain the same dataset after basic filters (remove taxa with zero counts) and collapsing the counts to the genus level; HMP16Data Bioconductor package was used to download the data.

Usage

data(ps_stool_16S)

Format

An object of class phyloseq

---

RMSE

RMSE

Description

Computes the Root Mean Square Error (RMSE) from a vector of differences.

Usage

RMSE(differences)

Arguments

differences a vector of differences.

Value

RMSE value

See Also

prepareObserved and meanDifferences.
# Examples

```r
# Generate the data.frame of Mean Differences and Zero Probability Difference
MD_df <- data.frame(MD = rpois(10, 5), ZPD = runif(10, -1, 1))

# Calculate RMSE for MD and ZPD values
RMSE(MD_df[, "MD"])
RMSE(MD_df[, "ZPD"])
```

---

### Description

Run the differential abundance detection methods.

### Usage

```r
runDA(method_list, object, weights = NULL, verbose = TRUE)
```

### Arguments

- `method_list`: a list object containing the methods and their parameters.
- `object`: a phyloseq object.
- `weights`: an optional numeric matrix giving observational weights.
- `verbose`: an optional logical value. If `TRUE`, information about the steps of the algorithm is printed. Default `verbose = TRUE`.

### Value

A named list containing the results for each method.

### Examples

```r
set.seed(1)
# Create a very simple phyloseq object
counts <- matrix(rnbinom(n = 60, size = 3, prob = 0.5), nrow = 10, ncol = 6)
metadata <- data.frame("Sample" = c("S1", "S2", "S3", "S4", "S5", "S6"),
"group" = as.factor(c("A", "A", "A", "B", "B", "B")))
ps <- phyloseq::phyloseq(phyloseq::otu_table(counts, taxa_are_rows = TRUE),
phyloseq::sample_data(metadata))

# Set some simple normalizations
my_norm <- setNormalizations()

# Add them to the phyloseq object
ps <- runNormalizations(normalization_list = my_norm, object = ps)

# Set some limma instances
```
my_methods <- set_limma(design = ~ group, coef = 2,
    norm = c("TMM", "poscounts", "CSS"))

# Run the methods
results <- runDA(method_list = my_methods, object = ps)

---

### runMocks

**Description**

Run the differential abundance detection methods on mock datasets.

**Usage**

```r
runMocks(
    mocks,
    method_list,
    object,
    weights = NULL,
    verbose = TRUE,
    BPPARAM = BiocParallel::SerialParam()
)
```

**Arguments**

- `mocks`: a data.frame containing N rows and nsamples columns (if even). Each cell of the data frame contains the "grp1" or "grp2" characters which represent the mock groups pattern. Produced by the `createMocks` function.
- `method_list`: a list object containing the methods and their parameters.
- `object`: a phyloseq or TreeSummarizedExperiment object.
- `weights`: an optional numeric matrix giving observational weights.
- `verbose`: an optional logical value. If TRUE, information about the steps of the algorithm is printed. Default verbose = TRUE.
- `BPPARAM`: An optional `BiocParallelParam` instance defining the parallel back-end to be used during evaluation.

**Value**

A named list containing the results for each method.
Examples

```r
# Load some data
data(ps_stool_16S)

# Generate the pattern for 10 mock comparisons
# (N = 1000 is suggested)
mocks <- createMocks(nsamples = phyloseq::nsamples(ps_stool_16S), N = 10)
head(mocks)

# Add some normalization/scaling factors to the phyloseq object
my_norm <- setNormalizations(fun = c("norm_edgeR", "norm_CSS"),
    method = c("TMM", "CSS"))
ps_stool_16S <- runNormalizations(normalization_list = my_norm,
    object = ps_stool_16S)

# Initialize some limma based methods
my_limma <- set_limma(design = ~ group, coef = 2, norm = c("TMM", "CSS"))

# Run methods on mock datasets
results <- runMocks(mocks = mocks, method_list = my_limma,
    object = ps_stool_16S)
```

Description

Add normalization/scaling factors to a phyloseq object

Usage

```r
runNormalizations(
    normalization_list,
    object,
    assay_name = "counts",
    verbose = TRUE
)
```

Arguments

- **normalization_list**: a list object containing the normalization methods and their parameters.
- **object**: a phyloseq or TreeSummarizedExperiment object.
- **assay_name**: the name of the assay to extract from the TreeSummarizedExperiment object (default `assayName = "counts"`). Not used if the input object is a phyloseq.
- **verbose**: an optional logical value. If `TRUE`, information about the steps of the algorithm is printed. Default `verbose = TRUE`. 

Value

A phyloseq object containing the normalization/scaling factors.

See Also

setNormalizations

Examples

```r
set.seed(1)
# Create a very simple phyloseq object
counts <- matrix(rnbinom(n = 60, size = 3, prob = 0.5), nrow = 10, ncol = 6)
metadata <- data.frame("Sample" = c("S1", "S2", "S3", "S4", "S5", "S6"),
                        "group" = as.factor(c("A", "A", "A", "B", "B", "B")))
ps <- phyloseq::phyloseq(phyloseq::otu_table(counts, taxa_are_rows = TRUE),
                        phyloseq::sample_data(metadata))

# Set some simple normalizations
my_normalizations <- setNormalizations()

# Add them to the phyloseq object
ps <- runNormalizations(normalization_list = my_normalizations, object = ps)
```

Description

Run the differential abundance detection methods on split datasets.

Usage

```r
runSplits(
  split_list,
  method_list,
  normalization_list,
  object,
  assay_name = "counts",
  min_counts = 0,
  min_samples = 0,
  verbose = TRUE,
  BPPARAM = BiocParallel::SerialParam()
)
```
Arguments

- **split_list**: A list of 2 data.frame objects: Subset1 and Subset2 produced by the `createSplits` function.
- **method_list**: a list object containing the methods and their parameters.
- **normalization_list**: a list object containing the normalization method names and their parameters produced by `setNormalizations`.
- **object**: a phyloseq object.
- **assay_name**: the name of the assay to extract from the TreeSummarizedExperiment object (default `assayName = "counts"`). Not used if the input object is a phyloseq.
- **min_counts**: Parameter to filter taxa. Set this number to keep features with more than `min_counts` counts in more than `min_samples` samples (default `min_counts = 0`).
- **min_samples**: Parameter to filter taxa. Set this number to keep features with a `min_counts` counts in more than `min_samples` samples (default `min_samples = 0`).
- **verbose**: an optional logical value. If TRUE, information about the steps of the algorithm is printed. Default `verbose = TRUE`.
- **BPPARAM**: An optional `BiocParallelParam` instance defining the parallel back-end to be used during evaluation.

Value

A named list containing the results for each method.

Examples

data(ps_plaque_16S)

# Balanced design
my_splits <- createSplits(
  object = ps_plaque_16S, varName = "HMP_BODY_SUBSITE", balanced = TRUE,
  paired = "RSID", N = 10 # N = 100 suggested
)

# Make sure the subject ID variable is a factor
phyloseq::sample_data(ps_plaque_16S)[, "RSID"] <- as.factor(
  phyloseq::sample_data(ps_plaque_16S)[["RSID"]]
)

# Initialize some limma based methods
my_limma <- set_limma(design = ~ RSID + HMP_BODY_SUBSITE,
  coef = "HMP_BODY_SUBSITESupragingival Plaque",
  norm = c("TMM", "CSS"))

# Set the normalization methods according to the DA methods
my_norm <- setNormalizations(fun = c("norm_edgeR", "norm_CSS"),
  method = c("TMM", "CSS"))

# Run methods on split datasets
results <- runSplits(split_list = my_splits, method_list = my_limma,
  normalization_list = my_norm, object = ps_plaque_16S)
setNormalizations

Description

Set the methods and parameters to compute normalization/scaling factors.

Usage

setNormalizations(
  fun = c("norm_edgeR", "norm_DESeq2", "norm_CSS"),
  method = c("TMM", "poscounts", "CSS")
)

Arguments

fun        a character with the name of normalization function (e.g. "norm_edgeR", "norm_DESeq2", "norm_CSS"...).
method     a character with the normalization method (e.g. "TMM", "upperquartile"... if the fun is "norm_edgeR").

Value

a list object containing the normalization methods and their parameters.

See Also

runNormalizations, norm_edgeR, norm_DESeq2, norm_CSS, norm_TSS

Examples

# Set a TMM normalization
my_TMM_normalization <- setNormalizations(fun = "norm_edgeR", method = "TMM")

# Set some simple normalizations
my_normalizations <- setNormalizations()

# Add a custom normalization
my_normalizations <- c(my_normalizations,
                        myNormMethod1 = list("myNormMethod", "parameter1", "parameter2"))
Description

Set the parameters for ALDEx2 differential abundance detection method.

Usage

```r
set_ALDEx2(
  assay_name = "counts",
  pseudo_count = FALSE,
  design = NULL,
  mc.samples = 128,
  test = "t",
  paired.test = FALSE,
  denom = "all",
  contrast = NULL,
  expand = TRUE
)
```

Arguments

- **assay_name**: the name of the assay to extract from the TreeSummarizedExperiment object (default `assayName = "counts"`). Not used if the input object is a phyloseq.
- **pseudo_count**: add 1 to all counts if TRUE (default `pseudo_count = FALSE`).
- **design**: a character with the name of a variable to group samples and compare them or a formula to compute a model.matrix (when `test = "glm"`).
- **mc.samples**: an integer. The number of Monte Carlo samples to use when estimating the underlying distributions. Since we are estimating central tendencies, 128 is usually sufficient.
- **test**: a character string. Indicates which tests to perform. "t" runs Welch’s t test while "wilcox" runs Wilcoxon test. "kw" runs Kruskal-Wallace test while "kw_glml" runs glm ANOVA-like test. "glm" runs a generalized linear model.
- **paired.test**: A boolean. Toggles whether to do paired-sample tests. Applies to `effect = TRUE` and `test = "t"`.
- **denom**: An any variable (all, iqlr, zero, lvha, median, user) indicating features to use as the denominator for the Geometric Mean calculation. The default "all" uses the geometric mean abundance of all features. Using "median" returns the median abundance of all features. Using "iqlr" uses the features that are between the first and third quartile of the variance of the clr values across all samples. Using "zero" uses the non-zero features in each group as the denominator. This approach is an extreme case where there are many nonzero features in one condition but many zeros in another. Using "lvha" uses features that have low variance (bottom quartile) and high relative abundance (top quartile in every sample). It is
also possible to supply a vector of row indices to use as the denominator. Here, the experimentalist is determining a-priori which rows are thought to be invariant. In the case of RNA-seq, this could include ribosomal protein genes and other house-keeping genes. This should be used with caution because the offsets may be different in the original data and in the data used by the function because features that are 0 in all samples are removed by `aldex.clr`.

**contrast** character vector with exactly three elements: the name of a variable used in "design", the name of the level of interest, and the name of the reference level. If "kw" or "kw_glm" as test, contrast vector is not used.

**expand** logical, if TRUE create all combinations of input parameters (default expand = TRUE)

**Value**
A named list containing the set of parameters for `DA_ALDEx2` method.

**See Also**
`DA_ALDEx2`

**Examples**

# Set some basic combinations of parameters for ALDEx2
default_ALDEx2 <- set_ALDEx2(design = "group",
                           contrast = c("group", "grp2", "grp1"))

# Set a specific set of normalization for ALDEx2 (even of other
# packages!)
defaultNorm_ALDEx2 <- set_ALDEx2(design = "group",
                               contrast = c("group", "grp2", "grp1"))

# Set many possible combinations of parameters for ALDEx2
defaultMany_ALDEx2 <- set_ALDEx2(design = "group",
                                denom = c("iqlr", "zero"),
                                test = c("t", "wilcox"), contrast = c("group", "grp2", "grp1"))

---

**set_ANCOM**

Set the parameters for ANCOM differential abundance detection method.

**Usage**

```r
set_ANCOM(
  assay_name = "counts",
  pseudo_count = FALSE,
  fix_formula = NULL,
  adj_formula = NULL,
  rand_formula = NULL,
)```

**Description**
Set the parameters for ANCOM differential abundance detection method.
lme_control = lme4::lmerControl()
contrast = NULL,
p_adj_method = "BH",
struc_zero = FALSE,
BC = TRUE,
n_cl = 1,
expand = TRUE
)

Arguments

assay_name
the name of the assay to extract from the TreeSummarizedExperiment object (default assayName = "counts"). Not used if the input object is a phyloseq.
pseudo_count
add 1 to all counts if TRUE (default pseudo_count = FALSE).
fix_formula
Used when BC = TRUE (ANCOM-BC2). The character string expresses how the microbial absolute abundances for each taxon depend on the fixed effects in metadata.
adj_formula
Used when BC = FALSE (ANCOM). The character string represents the formula for covariate adjustment. Default is NULL.
rand_formula
Optionally used when BC = TRUE or BC = FALSE. The character string expresses how the microbial absolute abundances for each taxon depend on the random effects in metadata. ANCOM and ANCOM-BC2 follows the lmerTest package in formulating the random effects. See ?lmerTest::lmer for more details. Default is rand_formula = NULL.
lme_control
a list of control parameters for mixed model fitting. See ?lme4::lmerControl for details.
contrast
character vector with exactly, three elements: a string indicating the name of factor whose levels are the conditions to be compared, the name of the level of interest, and the name of the other level.
p_adj_method
character. method to adjust p-values. Default is "holm". Options include "holm", "hochberg", "hommel", "bonferroni", "BH", "BY", "fdr", "none". See ?stats::p.adjust for more details.
struc_zero
logical. Whether to detect structural zeros based on group. Default is FALSE. See Details for a more comprehensive discussion on structural zeros.
BC
boolean for ANCOM method to use. If TRUE the bias correction (ANCOM-BC2) is computed (default BC = TRUE). When BC = FALSE computational time may increase and p-values are not computed.
n_cl
numeric. The number of nodes to be forked. For details, see ?parallel::makeCluster. Default is 1 (no parallel computing).
expand
logical, if TRUE create all combinations of input parameters (default expand = TRUE).

Value

A named list containing the set of parameters for DA_ANCOM method.
set_basic

See Also

DA_ANCOM

Examples

# Set some basic combinations of parameters for ANCOM with bias correction
base_ANCOMBC <- set_ANCOM(pseudo_count = FALSE, fix_formula = "group",
contrast = c("group", "B", "A"), BC = TRUE, expand = FALSE)
many_ANCOMs <- set_ANCOM(pseudo_count = c(TRUE, FALSE),
fix_formula = "group", contrast = c("group", "B", "A"),
struc_zero = c(TRUE, FALSE), BC = c(TRUE, FALSE))

Description

Set the parameters for basic differential abundance detection methods such as t and wilcox.

Usage

set_basic(
    assay_name = "counts",
    pseudo_count = FALSE,
    contrast = NULL,
    test = c("t", "wilcox"),
    paired = FALSE,
    expand = TRUE
)

Arguments

<table>
<thead>
<tr>
<th>Argument</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>assay_name</td>
<td>the name of the assay to extract from the TreeSummarizedExperiment object</td>
</tr>
<tr>
<td></td>
<td>(default assayName = &quot;counts&quot;). Not used if the input object is a phyloseq.</td>
</tr>
<tr>
<td>pseudo_count</td>
<td>add 1 to all counts if TRUE (default pseudo_count = FALSE).</td>
</tr>
<tr>
<td>contrast</td>
<td>character vector with exactly, three elements: a string indicating the name of</td>
</tr>
<tr>
<td></td>
<td>factor whose levels are the conditions to be compared, the name of the level</td>
</tr>
<tr>
<td></td>
<td>of interest, and the name of the other level.</td>
</tr>
<tr>
<td>test</td>
<td>name of the test to perform. Choose between &quot;t&quot; or &quot;wilcox&quot;.</td>
</tr>
<tr>
<td>paired</td>
<td>boolean. Choose whether the test is paired or not (default paired = FALSE).</td>
</tr>
<tr>
<td></td>
<td>If paired = TRUE be sure to provide the object properly ordered (by the grouping</td>
</tr>
<tr>
<td></td>
<td>variable).</td>
</tr>
<tr>
<td>expand</td>
<td>logical, if TRUE create all combinations of input parameters (default expand =</td>
</tr>
<tr>
<td></td>
<td>TRUE).</td>
</tr>
</tbody>
</table>
Value

A named list containing the set of parameters for DA_basic method.

See Also

DA_basic

Examples

# Set some basic methods
basic_methods <- set_basic(pseudo_count = FALSE, test = c("t", "wilcox"),
                            contrast = c("group", "B", "A"), expand = TRUE)

Description

Set the parameters for corncob differential abundance detection method.

Usage

set_corncob(
    assay_name = "counts",
    pseudo_count = FALSE,
    formula = NULL,
    phi.formula = NULL,
    formula_null = NULL,
    phi.formula_null = NULL,
    test = c("Wald", "LRT"),
    boot = FALSE,
    coefficient = NULL,
    expand = TRUE
)

Arguments

assay_name the name of the assay to extract from the TreeSummarizedExperiment object (default assayName = "counts"). Not used if the input object is a phyloseq.

pseudo_count add 1 to all counts if TRUE (default pseudo_count = FALSE).

formula an object of class formula without the response: a symbolic description of the model to be fitted to the abundance.

phi.formula an object of class formula without the response: a symbolic description of the model to be fitted to the dispersion.

formula_null Formula for mean under null, without response
phi.formula_null
- Formula for overdispersion under null, without response

test
- Character. Hypothesis testing procedure to use. One of "Wald" or "LRT" (likelihood ratio test).

boot
- Boolean. Defaults to FALSE. Indicator of whether or not to use parametric bootstrap algorithm. (See pbWald and pbLRT).

coefficient
- The coefficient of interest as a single word formed by the variable name and the non reference level. (e.g.: 'ConditionDisease' if the reference level for the variable 'Condition' is 'control').

expand
- logical, if TRUE create all combinations of input parameters (default expand = TRUE)

Value
- A named list containing the set of parameters for DA_corncob method.

See Also
- DA_corncob

Examples

```r
# Set some basic combinations of parameters for corncob
base_corncob <- set_corncob(formula = ~ group, phi.formula = ~ group,
                           formula_null = ~ 1, phi.formula_null = ~ group, coefficient = "groupB")

# Set many possible combinations of parameters for corncob
all_corncob <- set_corncob(pseudo_count = c(TRUE, FALSE), formula = ~ group,
                           phi.formula = ~ group, formula_null = ~ 1, phi.formula_null = ~ group,
                           coefficient = "groupB", boot = c(TRUE, FALSE))
```

Description
- Set the parameters for dearseq differential abundance detection method.

Usage

```r
set_dearseq(
    assay_name = "counts",
    pseudo_count = FALSE,
    covariates = NULL,
    variables2test = NULL,
    sample_group = NULL,
    test = c("permutation", "asymptotic"),
    preprocessed = FALSE,
)```
set_dearseq

```r
n_perm = 1000,
expand = TRUE
```

**Arguments**

- `assay_name` the name of the assay to extract from the TreeSummarizedExperiment object (default `assayName = "counts"`). Not used if the input object is a phyloseq.
- `pseudo_count` add 1 to all counts if TRUE (default `pseudo_count = FALSE`).
- `covariates` a character vector containing the colnames of the covariates to include in the model.
- `variables2test` a character vector containing the colnames of the variable of interest.
- `sample_group` a vector of length n indicating whether the samples should be grouped (e.g. paired samples or longitudinal data). Coerced to be a factor. Default is NULL in which case no grouping is performed.
- `test` a character string indicating which method to use to approximate the variance component score test, either 'permutation' or 'asymptotic' (default `test = "permutation"`).
- `preprocessed` a logical flag indicating whether the expression data have already been preprocessed (e.g. log2 transformed). Default is FALSE, in which case y is assumed to contain raw counts and is normalized into log(counts) per million.
- `n_perm` the number of perturbations. Default is 1000
- `expand` logical, if TRUE create all combinations of input parameters (default `expand = TRUE`).

**Value**

A named list containing the set of parameters for DA_dearseq method.

**See Also**

DA_dearseq

**Examples**

```r
# Set some basic combinations of parameters for dearseq
base_dearseq <- set_dearseq(pseudo_count = FALSE, variables2test = "group",
                           test = c("permutation", "asymptotic"), expand = TRUE)
```
**set_DESeq2**

**Description**

Set the parameters for DESeq2 differential abundance detection method.

**Usage**

```r
set_DESeq2(
  assay_name = "counts",
  pseudo_count = FALSE,
  design = NULL,
  contrast = NULL,
  alpha = 0.05,
  norm = c("ratio", "poscounts", "iterate"),
  weights_logical = FALSE,
  expand = TRUE
)
```

**Arguments**

- `assay_name` the name of the assay to extract from the TreeSummarizedExperiment object (default `assayName = "counts"`). Not used if the input object is a phyloseq.
- `pseudo_count` add 1 to all counts if TRUE (default `pseudo_count = FALSE`).
- `design` character or formula to specify the model matrix.
- `contrast` character vector with exactly three elements: the name of a factor in the design formula, the name of the numerator level for the fold change, and the name of the denominator level for the fold change.
- `alpha` the significance cutoff used for optimizing the independent filtering (by default 0.05). If the adjusted p-value cutoff (FDR) will be a value other than 0.05, alpha should be set to that value.
- `norm` name of the normalization method to use in the differential abundance analysis. Choose between the native DESeq2 normalization methods, such as `ratio`, `poscounts`, or `iterate`. Alternatively (only for advanced users), if `norm` is equal to "TMM", "TMMwsp", "RLE", "upperquartile", "posupperquartile", or "none" from `norm_edgeR`, "CSS" from `norm_CSS`, or "TSS" from `norm_TSS`, the normalization factors are automatically transformed into size factors. If custom factors are supplied, make sure they are compatible with DESeq2 size factors.
- `weights_logical` logical vector, if TRUE a matrix of observational weights will be used for differential abundance analysis (default `weights_logical = FALSE`).
- `expand` logical, if TRUE create all combinations of input parameters (default `expand = TRUE`).
set_edgeR

Value
A named list containing the set of parameters for DA_DESeq2 method.

See Also
DA_DESeq2

Examples
# Set some basic combinations of parameters for DESeq2
base_DESeq2 <- set_DESeq2(design = ~ group, contrast = c("group", "B", "A"))
# Set a specific set of normalization for DESeq2
setNorm_DESeq2 <- set_DESeq2(design = ~ group, contrast =
   c("group", "B", "A"), norm = c("ratio", "poscounts"))
# Set many possible combinations of parameters for DESeq2
all_DESeq2 <- set_DESeq2(pseudo_count = c(TRUE, FALSE), design = ~ group,
   contrast = c("group", "B", "A"), weights_logical = c(TRUE,FALSE))

set_edgeR

Description
Set the parameters for edgeR differential abundance detection method.

Usage
set_edgeR(
  assay_name = "counts",
  pseudo_count = FALSE,
  group_name = NULL,
  design = NULL,
  robust = FALSE,
  coef = 2,
  norm = c("TMM", "TMMwsp", "RLE", "upperquartile", "posupperquartile", "none"),
  weights_logical = FALSE,
  expand = TRUE
)

Arguments
assay_name the name of the assay to extract from the TreeSummarizedExperiment object
  (default assayName = "counts"). Not used if the input object is a phyloseq.
pseudo_count add 1 to all counts if TRUE (default pseudo_count = FALSE).
group_name character giving the name of the column containing information about experimental
group/condition for each sample/library.
design character or formula to specify the model matrix.
set_limma

robust  logical, should the estimation of prior.df be robustified against outliers?

coeff  integer or character index vector indicating which coefficients of the linear model are to be tested equal to zero.

norm  name of the normalization method to use in the differential abundance analysis. Choose between the native edgeR normalization methods, such as TMM, TMMwsp, RLE, upperquartile, posupperquartile, or none. Alternatively (only for advanced users), if norm is equal to "ratio", "poscounts", or "iterate" from norm_DESeq2, "CSS" from norm_CSS, or "TSS" from norm_TSS, the scaling factors are automatically transformed into normalization factors. If custom factors are supplied, make sure they are compatible with edgeR normalization factors.

weights_logical  logical vector, if true a matrix of observation weights must be supplied (default weights_logical = FALSE).

expand  logical, if TRUE create all combinations of input parameters (default expand = TRUE).

Value

A named list containing the set of parameters for DA_edgeR method.

See Also

DA_edgeR

Examples

# Set some basic combinations of parameters for edgeR
base_edgeR <- set_edgeR(group_name = "group", design = ~ group, coef = 2)

# Set a specific set of normalization for edgeR
setNorm_edgeR <- set_edgeR(group_name = "group", design = ~ group, coef = 2,
                          norm = c("TMM", "RLE"))

# Set many possible combinations of parameters for edgeR
all_edgeR <- set_edgeR(pseudo_count = c(TRUE, FALSE), group_name = "group",
                         design = ~ group, robust = c(TRUE, FALSE), coef = 2,
                         weights_logical = c(TRUE, FALSE))

Description

Set the parameters for limma differential abundance detection method.
Usage

set_limma(
  assay_name = "counts",
  pseudo_count = FALSE,
  design = NULL,
  coef = 2,
  norm = c("TMM", "TMMwsp", "RLE", "upperquartile", "posupperquartile", "none"),
  weights_logical = FALSE,
  expand = TRUE
)

Arguments

- **assay_name**: the name of the assay to extract from the TreeSummarizedExperiment object (default assayName = "counts"). Not used if the input object is a phyloseq.
- **pseudo_count**: add 1 to all counts if TRUE (default pseudo_count = FALSE).
- **design**: character name of the metadata columns, formula, or design matrix with rows corresponding to samples and columns to coefficients to be estimated.
- **coef**: integer or character index vector indicating which coefficients of the linear model are to be tested equal to zero.
- **norm**: name of the normalization method to use in the differential abundance analysis. Choose between the native edgeR normalization methods, such as TMM, TMMwsp, RLE, upperquartile, posupperquartile, or none. Alternatively (only for advanced users), if norm is equal to "ratio", "poscounts", or "iterate" from norm_DESeq2, "CSS" from norm_CSS, or "TSS" from norm_TSS, the scaling factors are automatically transformed into normalization factors. If custom factors are supplied, make sure they are compatible with edgeR normalization factors.
- **weights_logical**: logical vector, if TRUE a matrix of observational weights will be used for differential abundance analysis (default weights_logical = FALSE).
- **expand**: logical, if TRUE create all combinations of input parameters (default expand = TRUE).

Value

A named list containing the set of parameters for DA_limma method.

See Also

- DA_limma

Examples

# Set some basic combinations of parameters for limma
base_limma <- set_limma(design = ~ group, coef = 2)
# Set a specific set of normalization for limma (even of other packages!)
setNorm_limma <- set_limma(design = ~ group, coef = 2,
set_MAST

norm = c("TMM", "upperquartile")

# Set many possible combinations of parameters for limma
all_limma <- set_limma(pseudo_count = c(TRUE, FALSE), design = ~ group,
coef = 2, weights_logical = c(TRUE, FALSE))

Description

Set the parameters for MAST differential abundance detection method.

Usage

set_MAST(
  assay_name = "counts",
  pseudo_count = FALSE,
  rescale = c("median", "default"),
  design = NULL,
  coefficient = NULL,
  expand = TRUE
)

Arguments

  assay_name  the name of the assay to extract from the TreeSummarizedExperiment object
               (default assayName = "counts"). Not used if the input object is a phyloseq.
  pseudo_count   add 1 to all counts if TRUE (default pseudo_count = FALSE).
  rescale  Rescale count data, per million if 'default', or per median library size if 'median'
           ('median' is suggested for metagenomics data).
  design  The model for the count distribution. Can be the variable name, or a character
          similar to "~ 1 + group", or a formula, or a 'model.matrix' object.
  coefficient The coefficient of interest as a single word formed by the variable name and
          the non reference level. (e.g.: 'ConditionDisease' if the reference level for the
          variable 'Condition' is 'control').
  expand  logical, if TRUE create all combinations of input parameters (default expand =
          TRUE)

Value

A named list containing the set of parameters for DA_MAST method.

See Also

DA_MAST
Examples

# Set some basic combinations of parameters for MAST
base_MAST <- set_MAST(design = ~ group, coefficient = "groupB")

# Set many possible combinations of parameters for MAST
all_MAST <- set_MAST(pseudo_count = c(TRUE, FALSE), rescale = c("median", "default"), design = ~ group, coefficient = "groupB")

Description

Set the parameters for metagenomeSeq differential abundance detection method.

Usage

set_metagenomeSeq(
    assay_name = "counts",
    pseudo_count = FALSE,
    design = NULL,
    coef = 2,
    norm = "CSS",
    model = "fitFeatureModel",
    expand = TRUE
)

Arguments

assay_name the name of the assay to extract from the TreeSummarizedExperiment object (default assayName = "counts"). Not used if the input object is a phyloseq.
pseudo_count add 1 to all counts if TRUE (default pseudo_count = FALSE).
design the model for the count distribution. Can be the variable name, or a character similar to "~ 1 + group", or a formula.
coef coefficient of interest to grab log fold-changes.
norm name of the normalization method to use in the differential abundance analysis. Choose the native metagenomeSeq normalization method CSS. Alternatively (only for advanced users), if norm is equal to "TMM", "TMMwsp", "RLE", "upperquartile", "posupperquartile", or "none" from norm_edgeR, "ratio", "poscounts", or "iterate" from norm_DESeq2, or "TSS" from norm_TSS, the factors are automatically transformed into scaling factors. If custom factors are supplied, make sure they are compatible with metagenomeSeq normalization factors.
model character equal to "fitFeatureModel" for differential abundance analysis using a zero-inflated log-normal model, "fitZig" for a complex mathematical optimization routine to estimate probabilities that a zero for a particular feature in a sample is a technical zero or not. The latter model relies heavily on the limma package (default model = "fitFeatureModel").
set_NOISeq

set_NOISeq

expand logical, if TRUE create all combinations of input parameters (default expand = TRUE)

Value

A named list containing the set of parameters for DA_metagenomeSeq method.

See Also

DA_metagenomeSeq

Examples

# Set a basic combination of parameters for metagenomeSeq
base_mgs <- set_metagenomeSeq(design = ~ group, coef = 2)
# Set a specific model for metagenomeSeq
setModel_mgs <- set_metagenomeSeq(design = ~ group, coef = 2, model = "fitZig")
# Set many possible combinations of parameters for metagenomeSeq
all_mgs <- set_metagenomeSeq(pseudo_count = c(TRUE, FALSE), design = ~ group, coef = 2, model = c("fitFeatureModel", "fitZig"), norm = "CSS")
name of the normalization method to use in the differential abundance analysis. Choose between the native edgeR normalization methods, such as TMM, TMMwsp, RLE, upperquartile, posupperquartile, or none. Alternatively (only for advanced users), if `norm` is equal to "ratio", "poscounts", or "iterate" from `norm_DESeq2`, "CSS" from `norm_CSS`, or "TSS" from `norm_TSS`, the scaling factors are automatically transformed into normalization factors. If custom factors are supplied, make sure they are compatible with edgeR normalization factors.

`expand` logical, if TRUE create all combinations of input parameters (default `expand = TRUE`).

**Value**

A named list containing the set of parameters for DA_NOISeq method.

**See Also**

`DA_NOISeq`

**Examples**

```r
# Set a basic combination of parameters for NOISeq with 'tmm' normalization
base_NOISeq <- set_NOISeq(pseudo_count = FALSE, norm = "tmm",
 contrast = c("group", "B", "A"), expand = FALSE)
# try many normalizations
many_NOISeq <- set_NOISeq(pseudo_count = FALSE,
 norm = c("tmm", "uqua", "rpkm", "n"), contrast = c("group", "B", "A"))
```

**Description**

Set the parameters for Seurat differential abundance detection method.

**Usage**

```r
set_Seurat(
   assay_name = "counts",
   pseudo_count = FALSE,
   test = "wilcox",
   contrast = NULL,
   norm = "LogNormalize",
   scale.factor = 10000,
   expand = TRUE
)
```
Arguments

- **assay_name**: the name of the assay to extract from the TreeSummarizedExperiment object (default `assayName = "counts"`). Not used if the input object is a phyloseq.
- **pseudo_count**: add 1 to all counts if TRUE (default `pseudo_count = FALSE`).
- **test**: Denotes which test to use. Available options are:
  - "wilcox": Identifies differentially abundant features between two groups of samples using a Wilcoxon Rank Sum test (default).
  - "bimod": Likelihood-ratio test for the feature abundances, (McDavid et al., Bioinformatics, 2013).
  - "roc": Identifies 'markers' of feature abundance using ROC analysis. For each feature, evaluates (using AUC) a classifier built on that feature alone, to classify between two groups of cells. An AUC value of 1 means that abundance values for this feature alone can perfectly classify the two groupings (i.e. Each of the samples in group.1 exhibit a higher level than each of the samples in group.2). An AUC value of 0 also means there is perfect classification, but in the other direction. A value of 0.5 implies that the feature has no predictive power to classify the two groups. Returns a 'predictive power' (abs(AUC-0.5) * 2) ranked matrix of putative differentially expressed genes.
  - "t": Identify differentially abundant features between two groups of samples using the Student’s t-test.
  - "negbinom": Identifies differentially abundant features between two groups of samples using a negative binomial generalized linear model.
  - "poisson": Identifies differentially abundant features between two groups of samples using a poisson generalized linear model.
  - "LR": Uses a logistic regression framework to determine differentially abundant features. Constructs a logistic regression model predicting group membership based on each feature individually and compares this to a null model with a likelihood ratio test.
  - "MAST": Identifies differentially expressed genes between two groups of cells using a hurdle model tailored to scRNA-seq data. Utilizes the MAST package to run the DE testing.
  - "DESeq2": Identifies differentially abundant features between two groups of samples based on a model using DESeq2 which uses a negative binomial distribution (Love et al, Genome Biology, 2014).
- **contrast**: character vector with exactly three elements: the name of a factor in the design formula, the name of the numerator level for the fold change, and the name of the denominator level for the fold change.
- **norm**: Method for normalization.
  - LogNormalize: Feature counts for each sample are divided by the total counts of that sample and multiplied by the scale.factor. This is then natural-log transformed using log1p;
  - CLR: Applies a centered log ratio transformation;
weights_ZINB

- RC Relative counts. Feature counts for each sample are divided by the total counts of that sample and multiplied by the scale.factor. No log-transformation is applied. For counts per million (CPM) set scale.factor = 1e6;
- none No normalization

scale.factor Sets the scale factor for cell-level normalization
expand logical, if TRUE create all combinations of input parameters (default expand = TRUE)

Value
A named list containing the set of parameters for DA_Seurat method.

See Also
DA_Seurat

Examples
# Set some basic combinations of parameters for Seurat
base_Seurat <- set_Seurat(contrast = c("group", "B", "A"))
# Set many possible combinations of parameters for Seurat
all_Seurat <- set_Seurat(test = c("wilcox", "t", "negbinom", "poisson"),
                        norm = c("LogNormalize", "CLR", "RC", "none"),
                        scale.factor = c(1000, 10000), contrast = c("group", "B", "A"))

weights_ZINB

Description
Computes the observational weights of the counts under a zero-inflated negative binomial (ZINB) model. For each count, the ZINB distribution is parametrized by three parameters: the mean value and the dispersion of the negative binomial distribution, and the probability of the zero component.

Usage
weights_ZINB(
  object,
  assay_name = "counts",
  design,
  K = 0,
  commondispersion = TRUE,
  zeroinflation = TRUE,
  verbose = FALSE,
  ...
)
weights_ZINB

Arguments

object a phyloseq or TreeSummarizedExperiment object.
assay_name the name of the assay to extract from the TreeSummarizedExperiment object (default assayName = "counts"). Not used if the input object is a phyloseq.
design character name of the metadata columns, formula, or design matrix with rows corresponding to samples and columns to coefficients to be estimated (the user needs to explicitly include the intercept in the design).
K integer. Number of latent factors.
commondispersion Whether or not a single dispersion for all features is estimated (default TRUE).
zeroinflation Whether or not a ZINB model should be fitted. If FALSE, a negative binomial model is fitted instead.
verbose Print helpful messages.
... Additional parameters to describe the model, see zinbModel.

Value

A matrix of weights.

See Also

zinbFit for zero-inflated negative binomial parameters’ estimation and computeObservationalWeights for weights extraction.

Examples

set.seed(1)
# Create a very simple phyloseq object
counts <- matrix(rnbinom(n = 60, size = 3, prob = 0.5), nrow = 10, ncol = 6)
metadata <- data.frame("Sample" = c("S1", "S2", "S3", "S4", "S5", "S6"))
ps <- phyloseq::phyloseq(phyloseq::otu_table(counts, taxa_are_rows = TRUE),
phyloseq::sample_data(metadata))
# Calculate the ZINB weights
zinbweights <- weights_ZINB(object = ps, K = 0, design = "~ 1")
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