# Package 'MMUPHin'

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Type Package

**Title** Meta-analysis Methods with Uniform Pipeline for Heterogeneity in Microbiome Studies

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**Description** MMUPHin is an R package for meta-analysis tasks of microbiome cohorts. It has function interfaces for:

- a) covariate-controlled batch- and cohort effect adjustment,
- b) meta-analysis differential abundance testing,
- c) meta-analysis unsupervised discrete structure (clustering) discovery, and
- d) meta-analysis unsupervised continuous structure discovery.

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VignetteBuilder knitr

**Depends** R (>= 3.6)

**Imports** Maaslin2, metafor, fpc, igraph, ggplot2, dplyr, tidyr, cowplot, utils, stats, grDevices

**Suggests** testthat, BiocStyle, knitr, rmarkdown, magrittr, vegan, phyloseq, curatedMetagenomicData, genefilter

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# **R** topics documented:

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# Description

adjust\_batch takes as input a feature-by-sample matrix of microbial abundances, and performs batch effect adjustment given provided batch and optional covariate variables. It returns the batch-adjusted abundance matrix. Additional options and parameters can be passed through the control parameter as a list (see details).

# Usage

```
adjust_batch(feature_abd, batch, covariates = NULL, data, control)
```

## **Arguments**

feature_abd	feature-by-sample matrix of abundances (proportions or counts).
batch	name of the batch variable. This variable in data should be a factor variable and will be converted to so with a warning if otherwise.
covariates	name(s) of covariates to adjust for in the batch correction model.
data	data frame of metadata, columns must include batch and covariates (if specified).
control	a named list of additional control parameters. See details.

## **Details**

control should be provided as a named list of the following components (can be a subset).

**zero\_inflation** logical. Indicates whether or not a zero-inflated model should be run. Default to TRUE (zero-inflated model). If set to FALSE then the correction will be similar to ComBat as provided in the sva package.

pseudo\_count numeric. Pseudo count to add feature\_abd before the methods' log transformation.
Default to NULL, in which case adjust\_batch will set the pseudo count automatically to half of minimal non-zero values in feature\_abd.

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**diagnostic\_plot** character. Name for the generated diagnostic figure file. Default to "adjust\_batch\_diagnostic.pdf". Can be set to NULL in which case no output will be generated.

**conv** numeric. Convergence threshold for the method's iterative algorithm for shrinking batch effect parameters. Default to 1e-4.

maxit integer. Maximum number of iterations allowed for the method's iterative algorithm. Default to 1000.

**verbose** logical. Indicates whether or not verbose information will be printed.

#### Value

a list, with the following components:

**feature\_abd\_adj** feature-by-sample matrix of batch-adjusted abundances, normalized to the same per-sample total abundance as feature\_abd.

**control** list of additional control parameters used in the function call.

#### Author(s)

```
Siyuan Ma, <siyuanma@g.harvard.edu>
```

## **Examples**

continuous\_discover

Unsupervised meta-analytical discovery and validation of continuous structures in microbial abundance data

## **Description**

continuous\_discover takes as input a feature-by-sample matrix of microbial abundances. It first performs unsupervised continuous structure discovery (PCA) within each batch. Loadings of top PCs from each batch are then mapped against each other to identify "consensus" loadings that are reproducible across batches with a network community discovery approach with **igraph**. The identified consensus loadings/scores can be viewed as continuous structures in microbial profiles that are recurrent across batches and valid in a meta-analytical sense. continuous\_discover returns, among other output, the identified consensus scores for continuous structures in the provided microbial abundance profiles, as well as the consensus PC loadings which can be used to assign continuous scores to any sample with the same set of microbial features.

## Usage

```
continuous_discover(feature_abd, batch, data, control)
```

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## Arguments

feature\_abd feature-by-sample matrix of abundances (proportions or counts).

batch name of the batch variable. This variable in data should be a factor variable and

will be converted to so with a warning if otherwise.

data frame of metadata, columns must include batch.

control a named list of additional control parameters. See details.

#### **Details**

control should be provided as a named list of the following components (can be a subset).

**normalization** character. Similar to the normalization parameter in Maaslin2 but only "TSS" and "NONE" are allowed. Default to "TSS" (total sum scaling).

**transform** character. Similar to the transform parameter in Maaslin2 but only "AST" and "LOG" are allowed. Default to "AST" (arcsine square root transformation).

**pseudo\_count** numeric. Pseudo count to add feature\_abd before the transformation. Default to NULL, in which case pseudo count will be set automatically to 0 if transform="AST", and half of minimal non-zero values in feature\_abd if transform="LOG".

var\_perc\_cutoff numeric. A value between 0 and 1 that indicates the percentage variability explained to cut off at for selecting top PCs in each batch. Across batches, the top PCs that in total explain more than var\_perc\_cutoff of the total variability will be selected for meta-analytical continuous structure discovery. Default to 0.8 (PCs included need to explain at least 80 total variability).

cos\_cutoff numeric. A value between 0 and 1 that indicates cutoff for absolute cosine coefficients between PC loadings to construct the method's network with. Once the top PC loadings from each batch are selected, cosine coefficients between each loading pair are calculated which indicate their similarity. Loading pairs with absolute cosine coefficients surpassing cos\_cutoff are then considered as associated with each other, and represented as an edge between the pair in a PC loading network. Network community discovery can then be performed on this network to identified densely connected "clusters" of PC loadings, which represent meta-analytically recurrent continuous structures.

cluster\_function function. cluster\_function is used to perform community structure discovery in the constructed PC loading network. This can be any of the network cluster functions provided in igraph. Default to cluster\_optimal. Note that this option can be slow for larger datasets, in which case cluster\_fast\_greedy is recommended.

**network\_plot** character. Name for the generated network figure file. Default to "clustered\_network.pdf". Can be set to NULL in which case no output will be generated.

plot\_size\_cutoff integer. Clusters with sizes smaller than or equal to plot\_size\_cutoff will be excluded in the visualized network. Defaul to 2 - visualized clusters must have at least three nodes (PC loadings).

**diagnostic\_plot** character. Name for the generated diagnostic figure file. Default to "continuous\_diagnostic.pdf". Can be set to NULL in which case no output will be generated.

**verbose** logical. Indicates whether or not verbose information will be printed.

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#### Value

a list, with the following components:

**consensus\_scores** matrix of identified consensus continuous scores. Columns are the identified consensus scores and rows correspond to samples in feature\_abd.

**consensus\_loadings** matrix of identified consensus loadings. Columns are the identified consensus scores and rows correspond to features in feature\_abd.

mat\_vali matrix of validation cosine coefficients of the identified consensus loadings. Columns correspond to the identified consensus scores and rows correspond to batches.

network, communities, mat\_cos components for the constructed PC loading network and community discovery results. network is a igraph graph object for the constructed network of associated PC loadings. communities is a communities object for the identified consensus loading clusters in network (output from control\$cluster\_function). mat\_cos is the matrix of cosine coefficients between all selected top PCs from all batches.

**control** list of additional control parameters used in the function call.

#### Author(s)

```
Siyuan Ma, <siyuanma@g.harvard.edu>
```

## **Examples**

CRC\_abd

Species level feature abundance data of five public CRC studies

## **Description**

Species level relative abundance profiles of CRC and control patients in the five public studies used in Thomas et al. (2019). These were accessed through curatedMetagenomicData.

## Usage

```
data(CRC_abd)
```

## Format

A feature-by-sample matrix of species-level profiles

## Source

curated Metagenomic Data

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#### References

Thomas, Andrew Maltez, Paolo Manghi, Francesco Asnicar, Edoardo Pasolli, Federica Armanini, Moreno Zolfo, Francesco Beghini et al. "Metagenomic analysis of colorectal cancer datasets identifies cross-cohort microbial diagnostic signatures and a link with choline degradation." Nature medicine 25, no. 4 (2019): 667.

## **Examples**

```
data(CRC_abd)
# features included
rownames(CRC_abd)
# These are relative abundances
apply(CRC_abd, 2, sum)
# The following were used to generate the object
# library(curatedMetagenomicData)
# library(phyloseq)
# library(genefilter)
# datasets <- curatedMetagenomicData(</pre>
   c("FengQ_2015.metaphlan_bugs_list.stool"
      "HanniganGD_2017.metaphlan_bugs_list.stool",
      "VogtmannE_2016.metaphlan_bugs_list.stool",
      "YuJ_2015.metaphlan_bugs_list.stool",
      "ZellerG_2014.metaphlan_bugs_list.stool"),
    dryrun = FALSE)
# Construct phyloseq object from the five datasets
# physeq <-
    # Aggregate the five studies into ExpressionSet
   mergeData(datasets) %>%
   # Convert to phyloseq object
   ExpressionSet2phyloseq() %>%
    # Subset samples to only CRC and controls
   subset_samples(study_condition %in% c("CRC", "control")) %>%
   # Subset features to species
   subset_taxa(!is.na(Species) & is.na(Strain)) %>%
   # Normalize abundances to relative abundance scale
   transform_sample_counts(function(x) x / sum(x)) %>%
    # Filter features to be of at least 1e-5 relative abundance in five
    # samples
   filter_taxa(kOverA(5, 1e-5), prune = TRUE)
# CRC_abd <- otu_table(physeq)@.Data
```

CRC\_meta

Sample metadata of five public CRC studies

## **Description**

Metadata information of CRC and control patients in the five public studies used in Thomas et al. (2019). These were accessed through curatedMetagenomicData.

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#### Usage

```
data(CRC_meta)
```

#### **Format**

A data. frame of per-sample metadata information

#### Source

curatedMetagenomicData

#### References

Thomas, Andrew Maltez, Paolo Manghi, Francesco Asnicar, Edoardo Pasolli, Federica Armanini, Moreno Zolfo, Francesco Beghini et al. "Metagenomic analysis of colorectal cancer datasets identifies cross-cohort microbial diagnostic signatures and a link with choline degradation." Nature medicine 25, no. 4 (2019): 667.

## **Examples**

```
data(CRC_meta)
# has CRC and control samples across five studies
table(CRC_meta$studyID, CRC_meta$study_condition)
# The following were used to generate the object
# library(curatedMetagenomicData)
# library(phyloseq)
# library(genefilter)
# datasets <- curatedMetagenomicData(</pre>
   c("FengQ_2015.metaphlan_bugs_list.stool"
      "HanniganGD_2017.metaphlan_bugs_list.stool",
      "VogtmannE_2016.metaphlan_bugs_list.stool",
      "YuJ_2015.metaphlan_bugs_list.stool",
      "ZellerG_2014.metaphlan_bugs_list.stool"),
   dryrun = FALSE)
# Construct phyloseq object from the five datasets
# physeq <-
    # Aggregate the five studies into ExpressionSet
   mergeData(datasets) %>%
   # Convert to phyloseg object
  ExpressionSet2phyloseq() %>%
   # Subset samples to only CRC and controls
   subset_samples(study_condition %in% c("CRC", "control")) %>%
    # Subset features to species
   subset_taxa(!is.na(Species) & is.na(Strain)) %>%
    # Normalize abundances to relative abundance scale
   transform_sample_counts(function(x) x / sum(x)) %>%
   # Filter features to be of at least 1e-5 relative abundance in five
    # samples
   filter_taxa(kOverA(5, 1e-5), prune = TRUE)
# CRC_meta <- data.frame(sample_data(physeq))</pre>
# CRC_meta$studyID <- factor(CRC_meta$studyID)</pre>
```

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discrete_discover	Unsupervised meta-analytical discovery and validation of discrete clustering structures in microbial abundance data

### **Description**

discrete\_discover takes as input sample-by-sample dissimilarity measurements (generated from microbial abundance profiles), and performs unsupervised clustering within each batch across a range of cluster numbers. It then evaluates the support for each cluster number with both internal (i.e., samples within the batch) and external (i.e., samples in other batches) data. Internal evaluation is realized with prediction.strength and external evaluation is based on a generalized version of the same method. discrete\_discover generates as output the evaluation statistics for each cluster number. A cluster number with good support from both internal and external evaluations provides meta-analytical evidence for discrete structures in the microbial abundance profiles.

## Usage

```
discrete_discover(D, batch, data, control)
```

#### **Arguments**

D	sample-by-sample dissimilarity measurements. Should be provided as a dist object.
batch	name of the batch variable. This variable in data should be a factor variable and will be converted to so with a warning if otherwise.
data	data frame of metadata, columns must include batch.
control	a named list of additional control parameters. See details.

#### **Details**

control should be provided as a named list of the following components (can be a subset).

- **k\_max** integer. Maximum number of clusters to evaluate. discrete\_discover will evaluate clustering structures corresponding to cluster numbers ranging from 2 to k\_max. Default to 10.
- cluster\_function an interface function. This function will be used for unsupervised clustering for discrete structure evaluation. This corresponds to the clustermethod parameter in prediction.strength, and similarly, should also follow the specifications as detailed in clusterboot. Default to claraCBI
- classify\_method character. Classification method used to assign observations in the method's internal and external evaluation stage. Corresponds to the classification parameter in prediction.strength, and can only be either "centroid" or "knn". Default to "centroid".
- **M** integer. Number of random iterations to partition the batch during method's internal evaluation. Corresponds to the M parameter in prediction. strength. Default to 30.
- **nnk** integer. Numbber of nearest neighbors if classify\_method="knn". Corresponds to the nnk parameter in prediction.strength. Default to 1.

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**diagnostic\_plot** character. Name for the generated diagnostic figure file. Default to "discrete\_diagnostic.pdf". Can be set to NULL in which case no output will be generated.

**verbose** logical. Indicates whether or not verbose information will be printed.

## Value

a list, with the following components:

**internal\_mean, internal\_se** matrices of internal clustering structure evaluation measurements (prediction strengths). Columns and rows corresponds to different batches and different numbers of clusters, respectively. internal\_mean and internal\_se, as the names suggest, are the mean and standard error of prediction strengths for each batch/cluster number.

**external\_mean, external\_se** same structure as internal\_mean and internal\_se, but records external clustering structure evaluation measurements (generalized prediction strength).

**control** list of additional control parameters used in the function call.

#### Author(s)

Siyuan Ma, <siyuanma@g.harvard.edu>

## **Examples**

lm\_meta

Covariate adjusted meta-analytical differential abundance testing

## **Description**

Im\_meta runs differential abundance models on microbial profiles within individual studies/batches, and aggregates per-batch effect sizes with a meta-analysis fixed/random effects model. It takes as input a feature-by-sample microbial abundance table and the accompanying meta data data frame which should includes the batch indicator variable, the main exposure variable for differential abundance testing, and optional covariates and random covariates. The function first runs Maaslin2 models on the exposure with optional covariates/random covariates in each batch. The per-batch effect sizes are then aggregated with rma.uni and reported as output. Additional parameters, including those for both Maaslin2 and rma.uni can be provided through control (see details).

#### Usage

```
lm_meta(feature_abd, batch, exposure, covariates = NULL,
  covariates_random = NULL, data, control)
```

lm\_meta

## **Arguments**

feature\_abd feature-by-sample matrix of abundances (proportions or counts).

batch name of the batch variable. This variable in data should be a factor variable and

will be converted to so with a warning if otherwise.

exposure name of the exposure variable for differential abundance testing.

covariates names of covariates to adjust for in Maaslin2 differential abundance testing mod-

els.

covariates\_random

names of random effects grouping covariates to adjust for in Maaslin2 differen-

tial abundance testing models.

data frame of metadata, columns must include exposure, batch, and covariates

and covariates\_random (if specified).

control a named list of additional control parameters. See details.

#### **Details**

control should be provided as a named list of the following components (can be a subset).

**normalization** character. normalization parameter for Maaslin2. See Maaslin2 for details and allowed values. Default to "TSS" (total sum scaling).

**transform** character. transform parameter for Maaslin2. See Maaslin2 for details and allowed values. Default to "LOG" (log transformation).

**analysis\_method** character. analysis\_method parameter for Maaslin2. See Maaslin2 for details and allowed values. Default to "LM" (linear modeling).

**rma\_method** character. method parameter for rma.uni. See rma.uni for details and allowed values. Default to "REML" (estricted maximum-likelihood estimator).

**output** character. Output directory for intermediate Maaslin2 output and the optional forest plots. Default to "MMUPHin\_lm\_meta".

**forest\_plot** character. Suffix in the name for the generated forest plots visualizing significant metaanalytical differential abundance effects. Default to "forest.pdf". Can be set to NULL in which case no output will be generated.

**rma\_conv** numeric. Convergence threshold for rma.uni (corresponds to control\$threshold. See rma.uni for details. Default to 1e-4.

**rma\_maxit** integer. Maximum number of iterations allowed for rma.uni (corresponds to control\$maxiter. See rma.uni for details. Default to 1000.

verbose logical. Indicates whether or not verbose information will be printed.

#### Value

a list, with the following components:

**meta\_fits** data frame of per-feature meta-analytical differential abundance results, including columns for effect sizes, p-values and q-values, heterogeneity statistics such as  $\tau^2$  and  $I^2$ , as well as weights for individual batches. Many of these statistics are explained in detail in rma.uni.

**maaslin\_fits** list of data frames, each one corresponding to the fitted results of Maaslin2 in a individual batch. See Maaslin2 on details of these output.

**control** list of additional control parameters used in the function call.

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## Author(s)

```
Siyuan Ma, <siyuanma@g.harvard.edu>
```

## **Examples**

vaginal\_abd

Species level feature abundance data of two public vaginal studies

## **Description**

Species level relative abundance profiles of vaginal samples in the two public studies provided in curatedMetagenomicData.

## Usage

```
data(vaginal_abd)
```

#### **Format**

A feature-by-sample matrix of species-level profiles

# Source

curated Metagenomic Data

#### References

Pasolli, Edoardo, Lucas Schiffer, Paolo Manghi, Audrey Renson, Valerie Obenchain, Duy Tin Truong, Francesco Beghini et al. "Accessible, curated metagenomic data through ExperimentHub." Nature methods 14, no. 11 (2017): 1023.

## **Examples**

```
data(vaginal_abd)
# features included
rownames(vaginal_abd)
# These are relative abundances
apply(vaginal_abd, 2, sum)
# The following were used to generate the object
# library(curatedMetagenomicData)
# library(phyloseq)
# datasets <- curatedMetagenomicData(</pre>
```

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vaginal\_meta

Sample metadata of two public vaginal studies

## Description

Metadata information of vaginal samples in the two public studies provided in curatedMetagenomicData.

## Usage

```
data(vaginal_meta)
```

#### **Format**

A data. frame of per-sample metadata information

#### **Source**

curatedMetagenomicData

#### References

Pasolli, Edoardo, Lucas Schiffer, Paolo Manghi, Audrey Renson, Valerie Obenchain, Duy Tin Truong, Francesco Beghini et al. "Accessible, curated metagenomic data through ExperimentHub." Nature methods 14, no. 11 (2017): 1023.

## **Examples**

```
data(vaginal_meta)
# has vaginal samples across two studies
table(vaginal_meta$studyID, vaginal_meta$body_site)
# The following were used to generate the object
# library(curatedMetagenomicData)
# library(phyloseq)
# datasets <- curatedMetagenomicData(</pre>
```

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```
"*metaphlan_bugs_list.vagina*",
  dryrun = FALSE)
# Construct phyloseq object from the five datasets
# physeq <-
 # Aggregate the five studies into ExpressionSet
# mergeData(datasets) %>%
 # Convert to phyloseq object
# ExpressionSet2phyloseq() %>%
 # Subset features to species
# subset_taxa(!is.na(Species) & is.na(Strain)) %>%
 # Normalize abundances to relative abundance scale
# transform_sample_counts(function(x) x / sum(x)) %>%
 # Filter features to be of at least 1e-5 relative abundance in two samples
# filter_taxa(kOverA(2, 1e-5), prune = TRUE)
# vaginal_meta <- data.frame(sample_data(physeq))</pre>
# vaginal_meta$studyID <- factor(vaginal_meta$studyID)</pre>
```

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