Package 'FAVA'

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Title Quantify Compositional Variability Across Relative Abundance Vectors

Version 1.0.7

Description Implements the statistic FAVA, an Fst-based Assessment of Variability across vectors of relative Abundances, as well as a suite of helper functions which enable the visualization and statistical analysis of relative abundance data. The 'FAVA' R package accompanies the paper, "Quantifying compositional variability in microbial communities with FAVA" by Morrison, Xue, and Rosenberg (2024) <doi:10.1101/2024.07.03.601929>.

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https://maikemorrison.github.io/FAVA/articles/microbiome_tutorial.html

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Contents

bootstrap_fava	. 2
fava	. 4
fava_norm	. 6
gini_simpson	. 7
gini_simpson_mean	. 8
gini_simpson_pooled	. 9
plot_relabund	. 11
relab_phyloseq	. 14
time_weights	. 15
window_fava	. 16
window_list	. 17
window_plot	. 18
xue_microbiome_sample	. 19
xue_species_info	. 19
xue_species_similarity	. 20
xue_species_tree	. 20
	21

Index

bootstrap_fava

Statistically compare FAVA values between pairs of relative abundance matrices.

Description

bootstrap_fava uses bootstrapping to statistically compare FAVA values between pairs of relative abundance matrices. bootstrap_fava takes the same options as fava, so, as with fava, you can separately analyze multiple populations or groups of samples (specify group), and account for similarity among categories (specify S) or uneven weighting of rows (specify w or time). bootstrap_fava follows the bootstrapping procedure defined by Efron and Tibshirani (1993). Details on the bootstrapping procedure are available in the Methods section of the accompanying paper.

Usage

```
bootstrap_fava(
  relab_matrix,
  n_replicates = 1000,
  group,
  K = NULL,
  S = NULL,
  w = NULL,
  time = NULL,
  normalized = FALSE,
  seed = NULL,
  alternative = "two.sided"
)
```

bootstrap_fava

Arguments

relab_matrix	A matrix or data frame with rows containing non-negative entries that sum to 1. Each row represents a sample, each column represents a category, and each entry represents the abundance of that category in the sample. If relab_matrix contains any metadata, it must be on the left-hand side of the matrix, the right K entries of each row must sum to 1, and K must be specified. Otherwise, all entries of each row must sum to 1.
n_replicates	The number of bootstrap replicate matrices to generate. Default is n_replicates = 1000.
group	A string (or vector of strings) specifying the name(s) of the column(s) that de- scribes which group(s) each row (sample) belongs to. Use if relab_matrix is a single matrix containing multiple groups of samples you wish to compare.
К	Optional; an integer specifying the number of categories in the data. Default is K=ncol(relab_matrix).
S	Optional; a K x K similarity matrix with diagonal elements equal to 1 and off- diagonal elements between 0 and 1. Entry S[i,k] for i!=k is the similarity between category and i and category k, equaling 1 if the categories are to be treated as identical and equaling 0 if they are to be treated as totally dissimilar. The default value is S = diag(ncol(relab_matrix)).
w	Optional; a vector of length I with non-negative entries that sum to 1. Entry w[i] represents the weight placed on row i in the computation of the mean abundance of each category across rows. The default value is w = rep(1/nrow(relab_matrix), nrow(relab_matrix)).
time	Optional; a string specifying the name of the column that describes the sampling time for each row. Include if you wish to weight FAVA by the distance between samples.
normalized	Optional; should normalized FAVA be used? Default is normalized = FALSE; use normalized = TRUE to compute normalized FAVA. FAVA can only be nor- malized if it is not weighted.
seed	Optional; an integer to be used as a random seed for the simulations.
alternative	Optional; do you want to do a one- or two.sided test? Default is alternative = "two.sided". If you wish to do a one-sided test, specify either alternative = "lesser" or alternative = "greater".

Value

A named list containing the following entries:

- p_values: The probability of observing the observed difference in variability between each pair of groups if there were no difference between groups. Computed as the fraction of bootstrap differences greater than or equal to the observed difference. Depends on what alternative is specified ("greater", "lesser", or "two.sided").
- bootstrap_distribution_plot: The distribution of bootstrap replicate differences in each variability value. The observed differences are shown in red. The further the red points are from 0, the more significant the statistical difference between groups.
- observed_stats: The observed diversity statistics for the groups.
- bootstrap_stats: The bootstrap replicate diversity statistics for the groups.

Examples

Plots of the bootstrap distributions of differences in FAVA between each pair of matrices, # and how the true observed differences (red dots) compare to the distribution. boot_out\$bootstrap_distribution_plot

fava

Compute the Fst of a matrix of compositional vectors

Description

This function computes the population-genetic statistic Fst on any matrix with rows that sum to 1. Values of 0 are achieved when each row is a permutation of (1,0,...,0) and at least two categories have non-zero abundance across all rows. The value equals 1 when each row is identical.

Usage

```
fava(
   relab_matrix,
   K = NULL,
   S = NULL,
   w = NULL,
   time = NULL,
   group = NULL,
   normalized = FALSE
)
```

Arguments

```
relab_matrix A matrix or data frame with rows containing non-negative entries that sum to
1. Each row represents a sample, each column represents a category, and each
entry represents the abundance of that category in the sample. If relab_matrix
contains any metadata, it must be on the left-hand side of the matrix, the right
K entries of each row must sum to 1, and K must be specified. Otherwise, all
entries of each row must sum to 1.
```

fava	

К	Optional; an integer specifying the number of categories in the data. Default is K=ncol(relab_matrix).
S	Optional; a K x K similarity matrix with diagonal elements equal to 1 and off- diagonal elements between 0 and 1. Entry S[i,k] for i!=k is the similarity between category and i and category k, equaling 1 if the categories are to be treated as identical and equaling 0 if they are to be treated as totally dissimilar. The default value is S = diag(ncol(relab_matrix)).
W	Optional; a vector of length I with non-negative entries that sum to 1. Entry w[i] represents the weight placed on row i in the computation of the mean abundance of each category across rows. The default value is w = rep(1/nrow(relab_matrix), nrow(relab_matrix)).
time	Optional; a string specifying the name of the column that describes the sampling time for each row. Include if you wish to weight FAVA by the distance between samples.
group	Optional; a string (or vector of strings) specifying the name(s) of the column(s) that describes which group(s) each row (sample) belongs to. Use if relab_matrix is a single matrix containing multiple groups of samples you wish to compare.
normalized	Optional; should normalized FAVA be used? Default is normalized = FALSE; use normalized = TRUE to compute normalized FAVA. FAVA can only be normalized if it is not weighted.

Value

A numeric value between 0 and 1.

Examples

```
row_weights = c(0.5, 0, 0, 0.5)
fava(relative_abundances, w = row_weights)
```

```
# Compute fava assuming that
# categories 1 and 2 are identical:
similarity_matrix = diag(4)
similarity_matrix[1,2] = 1
```

```
similarity_matrix[2,1] = 1
fava(relative_abundances, S = similarity_matrix)
# Assume categories 1 and 2 are identical AND
# ignore rows 2 and 4:
row_weights = c(0.5, 0, 0.5, 0)
fava(relative_abundances, w = row_weights, S = similarity_matrix)
```

```
fava_norm
```

Compute the normalized Fst of a matrix of compositional vectors

Description

This function computes the normalized Fst given the number of rows and the mean abundance of the most abundant category. We employ the normalization employed in the FSTruct package by Morrison, Alcala, and Rosenberg (2020) doi: 10.1111/17550998.13647.

Usage

fava_norm(relab_matrix, K = ncol(relab_matrix))

Arguments

relab_matrix	A matrix or data frame with rows containing non-negative entries that sum to
	1. Each row represents a sample, each column represents a category, and each
	entry represents the abundance of that category in the sample. If relab_matrix
	contains any metadata, it must be on the left-hand side of the matrix, the right
	K entries of each row must sum to 1, and K must be specified. Otherwise, all entries of each row must sum to 1.
К	Optional; an integer specifying the number of categories in the data. Default is K=ncol(relab_matrix).

Value

A numeric value between 0 and 1.

Examples

fava_norm(relative_abundances)

gini_simpson

Description

This function computes the Gini-Simpson index, a statistical measure of variability known in population genetics as heterozygosity, of avector of non-negative entries which sum to 1. The function returns a number between 0 and 1 which quantifies the variability of the vector. Values of 0 are achieved when the vector is a permutation of (1,0,...,0). The value approaches 1 as the number of categories K increases when the vector is equal to (1/K, 1/K, ..., 1/K).

Usage

gini_simpson(q, K = length(q), S = diag(K))

Arguments

q	A vector with K=length(q) non-negative entries that sum to 1.
К	Optional; an integer specifying the number of categories in the data. Default is $K=length(q)$.
S	Optional; a K x K similarity matrix with diagonal elements equal to 1 and off- diagonal elements between 0 and 1. Entry S[i,k] for i!=k is the similarity between category and i and category k, equalling 1 if the categories are to be treated as identical and equaling 0 if they are to be treated as totally dissimilar. The default value is S = diag(ncol(q)).

Value

A numeric value between 0 and 1.

Examples

```
# Compute unweighted Gini-Simpson index:
gini_simpson(q = c(0.4, 0.3, 0.3))
```

```
# Compute Gini-Simpson index assuming that
# categories 1 and 2 are identical:
similarity_matrix = diag(3)
similarity_matrix[1,2] = 1
similarity_matrix[2,1] = 1
gini_simpson(q = c(0.4, 0.3, 0.3), S = similarity_matrix)
```

gini_simpson_mean

Description

This function computes the mean Gini-Simpson index, a statistical measure of variability known in population genetics as heterozygosity, of a set of vectors of non-negative entries which sum to 1. The function returns a number between 0 and 1 which quantifies the mean variability of the vectors. Values of 0 are achieved when each vector is a permutation of (1,0,...,0). The value approaches 1 as the number of categories K increases when the vectors are equal to (1/K, 1/K, ..., 1/K).

Usage

```
gini_simpson_mean(
  relab_matrix,
  K = NULL,
  S = NULL,
  w = NULL,
  time = NULL,
  group = NULL
)
```

Arguments

relab_matrix	A matrix or data frame with rows containing non-negative entries that sum to 1. Each row represents a sample, each column represents a category, and each entry represents the abundance of that category in the sample. If relab_matrix contains any metadata, it must be on the left-hand side of the matrix, the right K entries of each row must sum to 1, and K must be specified. Otherwise, all entries of each row must sum to 1.
К	Optional; an integer specifying the number of categories in the data. Default is K=ncol(relab_matrix).
S	Optional; a K x K similarity matrix with diagonal elements equal to 1 and off- diagonal elements between 0 and 1. Entry S[i,k] for i!=k is the similarity between category and i and category k, equalling 1 if the categories are to be treated as identical and equaling 0 if they are to be treated as totally dissimilar. The default value is S = diag(ncol(relab_matrix)).
W	Optional; a vector of length I with non-negative entries that sum to 1. Entry w[i] represents the weight placed on row i in the computation of the mean abundance of each category across rows. The default value is w = rep(1/nrow(relab_matrix), nrow(relab_matrix)).
time	Optional; a string specifying the name of the column that describes the sampling time for each row. Include if you wish to weight FAVA by the distance between samples.

group Optional; a string (or vector of strings) specifying the name(s) of the column(s) that describes which group(s) each row (sample) belongs to. Use if relab_matrix is a single matrix containing multiple groups of samples you wish to compare.

Value

A numeric value between 0 and 1.

Examples

```
# To compute the mean Gini-Simpson index of
# the following compositional vectors...
q1 = c(1, 0, 0,
                     0)
q2 = c(0.5, 0.5, 0,
                     0)
q3 = c(1/4, 1/4, 1/4, 1/4)
q4 = c(0, 0, 1, 0)
# we could compute the mean manually:
mean(sapply(list(q1, q2, q3, q4), gini_simpson))
# Or we could use gini_simpson_mean:
relative_abundances = matrix(c(q1, q2, q3, q4),
                  byrow = TRUE, nrow = 4)
gini_simpson_mean(relative_abundances)
# Incoporating weights:
# Compute mean Gini-Simpson index ignoring
# rows 2 and 3
row_weights = c(0.5, 0, 0, 0.5)
gini_simpson_mean(relative_abundances, w = row_weights)
# Compute mean Gini-Simpson index assuming that
# categories 1 and 2 are identical:
similarity_matrix = diag(4)
similarity_matrix[1,2] = 1
similarity_matrix[2,1] = 1
gini_simpson_mean(relative_abundances, S = similarity_matrix)
# Assume categories 1 and 2 are identical AND
# ignore rows 2 and 4:
row_weights = c(0.5, 0, 0.5, 0)
gini_simpson_mean(relative_abundances, w = row_weights, S = similarity_matrix)
```

gini_simpson_pooled Compute the pooled Gini-Simpson index of the rows in a matrix of compositional vectors

Description

This function computes the Gini-Simpson index of a "pooled" vector equal to colMeans(relab_matrix). Values of 0 are achieved when this pooled vector is a permutation of (1,0,..., 0). The value approaches 1 as the number of categories K increases when this pooled vector is equal to (1/K, 1/K, ..., 1/K).

Usage

```
gini_simpson_pooled(
  relab_matrix,
  K = NULL,
  S = NULL,
  w = NULL,
  time = NULL,
  group = NULL
)
```

Arguments

relab_matrix	A matrix or data frame with rows containing non-negative entries that sum to 1. Each row represents a sample, each column represents a category, and each entry represents the abundance of that category in the sample. If relab_matrix contains any metadata, it must be on the left-hand side of the matrix, the right K entries of each row must sum to 1, and K must be specified. Otherwise, all entries of each row must sum to 1.
К	Optional; an integer specifying the number of categories in the data. Default is K=ncol(relab_matrix).
S	Optional; a K x K similarity matrix with diagonal elements equal to 1 and off- diagonal elements between 0 and 1. Entry S[i,k] for i!=k is the similarity between category and i and category k, equalling 1 if the categories are to be treated as identical and equaling 0 if they are to be treated as totally dissimilar. The default value is S = diag(ncol(relab_matrix)).
W	Optional; a vector of length I with non-negative entries that sum to 1. Entry w[i] represents the weight placed on row i in the computation of the mean abundance of each category across rows. The default value is w = rep(1/nrow(relab_matrix), nrow(relab_matrix)).
time	Optional; a string specifying the name of the column that describes the sampling time for each row. Include if you wish to weight FAVA by the distance between samples.
group	Optional; a string (or vector of strings) specifying the name(s) of the column(s) that describes which group(s) each row (sample) belongs to. Use if relab_matrix is a single matrix containing multiple groups of samples you wish to compare.

Value

A numeric value between 0 and 1.

plot_relabund

Examples

```
# To compute the pooled Gini-Simpson index of
# the following compositional vectors...
q1 = c(1, 0, 0,
                      0)
q2 = c(0.5, 0.5, 0,
                      0)
q3 = c(1/4, 1/4, 1/4, 1/4)
q4 = c(0, 0, 1, 0)
# we could compute the mean manually:
qPooled = (q1 + q2 + q3 + q4)/4
gini_simpson(qPooled)
# Or we could use gini_simpson_pooled:
relative_abundances = matrix(c(q1, q2, q3, q4),
                  byrow = TRUE, nrow = 4)
gini_simpson_pooled(relative_abundances)
# Incoporating weights:
# Compute pooled Gini-Simpson index ignoring
# rows 2 and 3
row_weights = c(0.5, 0, 0, 0.5)
gini_simpson_pooled(relative_abundances, w = row_weights)
# Compute pooled Gini-Simpson index assuming that
# categories 1 and 2 are identical:
similarity_matrix = diag(4)
similarity_matrix[1,2] = 1
similarity_matrix[2,1] = 1
gini_simpson_pooled(relative_abundances, S = similarity_matrix)
# Assume categories 1 and 2 are identical AND
# ignore rows 2 and 4:
row_weights = c(0.5, 0, 0.5, 0)
gini_simpson_pooled(relative_abundances, w = row_weights, S = similarity_matrix)
```

plot_relabund Visualize a relative abundance matrix as a stacked bar
--

Description

This function enables graphical visualization of a matrix of compositional data. In the output plot, each vertical bar represents a single vector; the height of each color in the bar corresponds to the abundance of each category in that vector. Because this function produces a ggplot object, its output can be modified using standard ggplot2 syntax.

Usage

```
plot_relabund(
   relab_matrix,
   group = NULL,
   time = NULL,
   w = NULL,
   K = NULL,
   arrange = FALSE
)
```

Arguments

relab_matrix	A matrix or data frame with rows containing non-negative entries that sum to 1. Each row represents a sample, each column represents a category, and each entry represents the abundance of that category in the sample. If relab_matrix contains any metadata, it must be on the left-hand side of the matrix, the right K entries of each row must sum to 1, and K must be specified. Otherwise, all entries of each row must sum to 1.
group	Optional; a string specifying the name of the column that describes which group each row (sample) belongs to. Use if matrices is a single matrix containing multiple groups of samples you wish to compare.
time	Optional; a string specifying the name of the column that describes the sampling time for each row. Include if you wish to weight FAVA by the distance between samples.
W	Optional; a vector of length I with non-negative entries that sum to 1. Entry w[i] represents the weight placed on row i in the computation of the mean abundance of each category across rows. The default value is w = rep(1/nrow(relab_matrix), nrow(relab_matrix)).
K	Optional; an integer specifying the number of categories in the data. Default is K=ncol(relab_matrix).
arrange	Optional; controls horizontal ordering of samples and vertical ordering of cat- egories. If arrange = TRUE or arrange = "both", samples are ordered by the categories of greatest abundance and categories are ordered in decreasing abun- dance. If arrange = "vertical", sample order is unchanged but categories are ordered in decreasing abundance. If arrange = "horizontal", samples are ordered by the most abundant categories, but category order is unchanged. If arrange is missing or arrange = FALSE, neither order is changed.

Value

A ggplot object containing a bar plot visualization of the relative abundance matrix.

Examples

```
# Make an example matrix of compositional data
# Each row is an individual. Rows sum to 1.
population_A = matrix(c(
```

```
.5, .3, .2,
  .4, .2, .4,
  .5, .4, .1,
  .6, .1, .3,
  .2, 0, .8
),
nrow = 5,
byrow = TRUE
)
plot_relabund(relab_matrix = population_A,
            K = 3, # How many categories per vector?
            arrange = FALSE
            )
plot_relabund(relab_matrix = population_A,
            K = 3, # How many categories per vector?
            arrange = "horizontal"
            )
plot_relabund(relab_matrix = population_A,
            K = 3, # How many categories per vector?
            arrange = "vertical"
            )
 plot_relabund(relab_matrix = population_A,
            K = 3, # How many categories per vector?
            arrange = TRUE # could also be "both"
            )
```

```
# Plot a dataset which has 2 populations
```

```
population_B = matrix(c(
   .9, 0, .1,
   .6, .4, 0,
   .7, 0, .3,
   .3, .4, .3,
   .5, .3, .2
),
nrow = 5,
byrow = TRUE
)
```

relab_phyloseq

Generate a relative abundance matrix with sample metadata and OTU abundances from a phyloseq object.

Description

The R package phyloseq streamlines the storage and analysis of microbiome sequence data. This function takes a phyloseq object and extracts the OTU table and the sample metadata and combines them into one relative abundance matrix with rows corresponding to samples, metadata on the left-hand side, and OTU relative abundances on the right-hand side.

Usage

```
relab_phyloseq(phyloseq_object)
```

Arguments

phyloseq_object

A phyloseq object containing both an OTU table (otu_table) and sample metadata (sample_data).

Value

A data frame with rows representing samples and columns representing sample data categories or OTU relative abundances. OTU abundances are automatically normalized so that they sum to 1 for each sample, though a warning will be provided if a renormalization was necessary.

Examples

```
if (requireNamespace("phyloseq", quietly = TRUE)) {
   data(GlobalPatterns, package = "phyloseq")
# Make a small phyloseq object for demonstration
```

phyloseq_subset = phyloseq::subset_taxa(phyloseq::subset_samples(GlobalPatterns,

```
X.SampleID %in%
c("CL3", "CC1")),
Order == "Cenarchaeales")
otu_table = relab_phyloseq(phyloseq_subset)
otu_table[, 1:10]
}
```

time_weights

Compute a normalized weighting vector based on a vector of sampling times.

Description

This function takes a vector of sampling times, $t = (t_1, t_2, ..., t_I)$ and computes a normalized vector which can be used to weight each sample based on the time between the subsequent and the preceding samples. The weighting vector w is defined such that each entry, $w_i = d_i/2T$, where $T = t_I - t_1$ and $d_i = t_{i+1} - t_{i-1}$ for i not equal to 1 or I. $d_1 = t_2 - t_1$ and $d_I = t_I - t_{I-1}$.

Usage

time_weights(times, group = NULL)

Arguments

times	A numeric vector of sampling times. Each entry must be greater than the previous entry.
group	Optional; a character vector specifying the group identity of each sampling time. Use if there are samples from multiple replicates or subjects in one dataset.

Value

A numeric vector. Each entry provides a weight for each entry in the provided times vector. If group is not specified, the vector sums to 1. If group is specified, the vector sums to the number of distinct groups.

Examples

```
time_vector = c(1, 8, 22, 23, 24, 25, 26, 27, 28, 29, 30, 31,
32, 33, 34, 35, 36, 37, 38, 39, 44, 50, 57, 64)
```

```
time_weights(times = time_vector)
```

window_fava

Description

This function computes FAVA in sliding window slices of a dataset.

Usage

```
window_fava(
   relab_matrix,
   window_size,
   window_step = 1,
   group = NULL,
   index = NULL,
   time = NULL,
   w = NULL,
   S = NULL,
   K = NULL,
   normalized = FALSE,
   alpha = 0.5
)
```

Arguments

relab_matrix	A matrix or data frame with rows containing non-negative entries that sum to 1. Each row represents a sample, each column represents a category, and each entry represents the abundance of that category in the sample. If relab_matrix contains any metadata, it must be on the left-hand side of the matrix, the right K entries of each row must sum to 1, and K must be specified. Otherwise, all entries of each row must sum to 1.
window_size	An integer number specifying the number of samples per window.
window_step	Optional; an integer specifying the distance between the first entry of adjacent windows. Default is window_step=1.
group	Optional; a string specifying the name of the column that describes which group each row (sample) belongs to. Use if relab_matrix is a single matrix containing multiple groups of samples you wish to compare.
index	Optional; a string specifying the name of the column in relab_matrix con- taining an index for each sample. For example, if relab_matrix contains time series data, index would be the column containing the time of each sample. If index is not specified but time is, time is by default used as the index.
time	Optional; a string specifying the name of the column that describes the sampling time for each row. Include if you wish to weight FAVA by the distance between samples.

W	Optional; a vector of length I with non-negative entries that sum to 1. Entry w[i] represents the weight placed on row i in the computation of the mean abundance of each category across rows. The default value is w = rep(1/nrow(relab_matrix) nrow(relab_matrix)).
S	Optional; a K x K similarity matrix with diagonal elements equal to 1 and off- diagonal elements between 0 and 1. Entry S[i,k] for i!=k is the similarity between category and i and category k, equaling 1 if the categories are to be treated as identical and equaling 0 if they are to be treated as totally dissimilar. The default value is S = diag(ncol(relab_matrix)).
К	Optional; an integer specifying the number of categories in the data. Default is K=ncol(relab_matrix).
normalized	Optional; should normalized FAVA be used? Default is normalized = FALSE; use normalized = TRUE to compute normalized FAVA. FAVA can only be normalized if it is not weighted.
alpha	Optional; number between 0 and 1 specifying the opacity of the horizontal lines plotted. Default is $alpha = 0.5$.

Value

A list of values of FAVA for each window.

Examples

window_list	Generate sliding windows of specified length given the maximum num-
	ber of samples

Description

This function generates a list of of sliding windows conditional on two parameters: the length of each window (number of samples) and the total number of samples present in the data.

Usage

```
window_list(window_size, length, window_step = 1)
```

Arguments

window_size	An integer number specifying the number of samples per window.
length	An integer number specifying the total number of samples.
window_step	Optional; an integer number specifying the distance between the first entry of adjacent windows. Default is window_step=1.

Value

A list of samples of sample indices. Each list entry represents one window.

Examples

```
window_list(window_size = 6, length = 40)
window_list(window_size = 6, length = 40, window_step = 2)
```

window_plot

Generate a plot of FAVA in sliding windows.

Description

This function generates a plot of normalized or unnormalized, weighted or unweighted FAVA computed in sliding windows across samples for one or many groups of samples.

Usage

window_plot(window_fava, alpha = 0.5)

Arguments

window_fava	The output of window_fava.
alpha	Optional; number between 0 and 1 specifying the opacity of the horizontal lines plotted. Default is $alpha = 0.5$.

Value

A ggplot2 object.

Examples

xue_microbiome_sample Temporal microbiome composition data

Description

A subset of the data generated by Xue et al. (2024) detailing longitudinal composition of the human gut microbiome for three subjects who experience an antiobitic perturbation between days 29 and 34. We include only the subjects XAA, XBA, and XCA. Data from doi: 10.1101/2023.09.26.559480.

Usage

xue_microbiome_sample

Format

xue_microbiome_sample:A data frame with 75 rows and 1,348 columns:subject Subject ID: XBA, XDA, or XMAtimepoint Time (days) of sample collection... Species names ...

xue_species_info Table of species information

Description

A data frame providing taxonomic information for the species included in xue_species_tree.

Usage

xue_species_info

Format

xue_species_info:

A data frame with 1346 rows and 9 columns:

species_id The species_id given in xue_microbiome_sample

kingdom, phylum, class, order, family, genus, species The corresponding taxonomic category for each species

species_id_number The numeric code associated with each species, as used in xue_species_tree

xue_species_similarity

Species similarity matrix for the species included in xue_microbiome_sample

Description

A similarity matrix, with entry (i,j) corresponding to the pairwise similarity between species i and species j. This similarity matrix was derived from a phylogenetic distance matrix, inferred from the tree xue_species_tree, using the expression s(i,j) = exp(-d(i,j)), where d(i,j) is the phylogenetic distance between species i and j.

Usage

```
xue_species_similarity
```

Format

```
xue_species_similarity:
A data frame with 524 rows and 524 columns, each corresponding to one species.
```

xue_species_tree Phylogenetic tree for the species included in xue_microbiome_sample

Description

A phylogenetic tree in the Newick format.

Usage

xue_species_tree

Format

xue_species_tree:
A Newick tree.

Index

* datasets xue_microbiome_sample, 19 xue_species_info, 19 xue_species_similarity, 20 xue_species_tree, 20 bootstrap_fava, 2 fava, 4 fava_norm, 6 gini_simpson, 7 gini_simpson_mean, 8 gini_simpson_pooled, 9 $\texttt{plot_relabund}, \texttt{11}$ relab_phyloseq, 14 time_weights, 15 window_fava, $16\,$ window_list, 17 window_plot, 18 xue_microbiome_sample, 19 xue_species_info, 19 xue_species_similarity, 20 xue_species_tree, 20