Package ‘SYNCSA’

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Type Package

Title Analysis of Functional and Phylogenetic Patterns in Metacommunities

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Imports vegan, FD, permute, RcppArmadillo, utils, stats, graphics, parallel

Description Analysis of metacommunities based on functional traits and phylogeny of the community components. The functions that are offered here implement for the R environment methods that have been available in the SYNCSA application written in C++ (by Valerio Pillar, available at <http://ecoqua.ecologia.ufrgs.br/SYNCSA.html>).

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Collate 'CollectNames.R' 'ProgressBAR.R' 'belonging.R' 'cent.norm.R'
 'cor.matrix.R' 'cor.mantel.R' 'cor.matrix.partial.R'
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 'var.dummy.R' 'var.type.R'

NeedsCompilation no

Repository CRAN

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Description

Artificial data for run SYNCSA.

Usage

data(ADRS)

Format

A list with 4 matrices.

community Community data, observations of six species in ten sites.
traits Matrix data of species described by two traits.
phylo Matrix containing phylogenetic distance between species.
envir Two environmental variables for each community.
**belonging**

**Examples**

```r
data(ADRS)
```

---

**belonging  Degree of belonging of species**

---

**Description**

Function to obtain a matrix containing the degrees of belonging of each and every species. The degree of belonging of each species is defined based on its ecological or phylogenetic resemblance to every other species in the community. For more details, see [matrix.p](#), [matrix.x](#) and [syncsa](#).

**Usage**

```r
belonging(dis, standardize = TRUE)
```

**Arguments**

- `dis` Matrix containing distance between species.
- `standardize` Logical argument (TRUE or FALSE) to specify if dis must be standardized in values ranging from 0 to 1 (Default standardize = TRUE).

**Value**

Standardized matrix containing the degree of belonging of species in relation to each other. Row totals (species) = 1.

**Author(s)**

Vanderlei Julio Debastiani <vanderleidebastiani@yahoo.com.br>

**References**


**See Also**

- [matrix.p](#), [matrix.x](#), [syncsa](#)

**Examples**

```r
data(ADRS)
belonging(ADRS$phylo)
```
**cent.norm**  
*Matrix centralization and standardization*

**Description**

Internal function for centralization and standardization in a matrix.

**Usage**

```r
cent.norm(x, na.rm = FALSE)
```

**Arguments**

- `x`: A matrix
- `na.rm`: Logical argument (TRUE or FALSE) to specify if missing observations are removed (Default `na.rm = FALSE`).

**Author(s)**

Vanderlei Julio Debastiani <vanderleidebastiani@yahoo.com.br>

**See Also**

`syncsa`

---

**CollectNames**  
*Collect names an entire list*

**Description**

Internal function to collect names an entire list.

**Usage**

```r
CollectNames(l, prefix = NULL)
```

**Arguments**

- `l`: A list.
- `prefix`: A prefix to names.

**Value**

The names.
Function to obtain the correlation between two matrices and partial matrix correlation between three matrices.

**Description**

The functions `cor.matrix` and `cor.matrix.partial` are similar the function `mantel` and `mantel.partial`, although the significance of the statistics is evaluated differently from Mantel. The functions `pro.matrix` and `pro.matrix.partial` use symmetric Procrustes as a measure of concordance between data sets. The function `cor.mantel` is similar to the function `mantel`, but allows the use of a set of predefined permutation. For more details, see `syncsa`.

**Usage**

```r
cor.matrix(
  mx1,
  mx2,
  x,
  my1 = NULL,
  my2 = NULL,
  y,
  permut.my2 = FALSE,
  method = "pearson",
  dist = "euclidean",
  permutations = 999,
  norm = FALSE,
  norm.y = FALSE,
  strata = NULL,
  na.rm = FALSE,
  seqpermutation = NULL,
  parallel = NULL,
  newClusters = TRUE,
  CL = NULL
)

cor.mantel(
  dist.x,
  dist.y,
  method = "pearson",
  permutations = 999,
  strata = NULL,
  na.rm = FALSE,
  seqpermutation = NULL,
  parallel = NULL,
  newClusters = TRUE,
  CL = NULL
)
```
cor.matrix.partial(
  mx1,
  mx2,
  x,
  my1 = NULL,
  my2 = NULL,
  y,
  mz1 = NULL,
  mz2 = NULL,
  z,
  method = "pearson",
  dist = "euclidean",
  permute.my2 = FALSE,
  permute.mz2 = FALSE,
  permutations = 999,
  norm = FALSE,
  norm.y = FALSE,
  norm.z = FALSE,
  strata = NULL,
  na.rm = FALSE,
  seqpermuation = NULL,
  parallel = NULL,
  newClusters = TRUE,
  CL = NULL
)

cor.matrix2(
  mx1,
  mx2,
  x,
  y,
  method = "pearson",
  dist = "euclidean",
  put.together = NULL,
  permutations = 999,
  strata = NULL,
  na.rm = FALSE,
  seqpermuation = NULL,
  parallel = NULL,
  newClusters = TRUE,
  CL = NULL
)

cor.procrustes(
  x,
  y,
  permutations = 999,
cor.matrix

strata = NULL,
na.rm = FALSE,
seqpermutation = NULL,
parallel = NULL,
newClusters = TRUE,
CL = NULL
)

pro.matrix(
    mx1,
    mx2,
    x,
    my1 = NULL,
    my2 = NULL,
    y,
    permute.my2 = FALSE,
    permutations = 999,
    norm = FALSE,
    norm.y = FALSE,
    strata = NULL,
    seqpermutation = NULL,
    parallel = NULL,
    newClusters = TRUE,
    CL = NULL
)

pro.matrix.partial(
    mx1,
    mx2,
    x,
    my1 = NULL,
    my2 = NULL,
    y,
    mz1 = NULL,
    mz2 = NULL,
    z,
    permute.my2 = FALSE,
    permute.mz2 = FALSE,
    permutations = 999,
    norm = FALSE,
    norm.y = FALSE,
    norm.z = FALSE,
    strata = NULL,
    seqpermutation = NULL,
    parallel = NULL,
    newClusters = TRUE,
    CL = NULL
)
pro.matrix2(
  mx1,
  mx2,
  x,
  y,
  put.together = NULL,
  permutations = 999,
  strata = NULL,
  seqpermutation = NULL,
  parallel = NULL,
  newClusters = TRUE,
  CL = NULL
)

Arguments

mx1  Matrix that multiplied by mx2 results in the matrix x.
mx2  Matrix that when multiplied by mx1 results in the matrix x. See ‘details’ below.
x    Matrix that will be correlated with the matrix y.
my1  Matrix that multiplied by my2 results in the matrix y.
my2  Matrix that when multiplied by my1 results in the matrix y. See ‘details’ below.
y    Matrix that will be correlated with the matrix x.
permute.my2 Logical argument (TRUE or FALSE) to specify if realize parallel permutation in matrix my2.
method Correlation method, as accepted by cor: "pearson", "spearman" or "kendall".
dist  Dissimilarity index, as accepted by vegdist: "manhattan", "euclidean", "canberra", "bray", "kulczynski", "jaccard", "gower", "altGower", "morisita", "horn", "mountford", "raup", "binomial" or "chao".
permutations Number of permutations in assessing significance.
norm  Logical argument (TRUE or FALSE) to specify if x is standardized within variables (Default norm = FALSE).
norm.y Logical argument (TRUE or FALSE) to specify if y is standardized within variables (Default norm = FALSE).
strata Argument to specify restricting permutations within species groups (Default strata = NULL).
na.rm Logical argument (TRUE or FALSE) to specify if pairwise deletion of missing observations when computing dissimilarities (Default na.rm = FALSE).
seqpermutation A set of predefined permutation, with the same dimensions of permutations (Default seqpermutation = NULL).
parallel Number of parallel processes. Tip: use parallel::detectCores() (Default parallel = NULL).
newClusters Logical argument (TRUE or FALSE) to specify if make new parallel processes or use predefined socket cluster. Only if parallel is different of NULL (Default newClusters = TRUE).
cor.matrix

CL A predefined socket cluster done with parallel package.
dist.x Dissimilarity matrices of class dist.
dist.y Dissimilarity matrices of class dist.
mz1 Matrix that multiplied by mz2 results in the matrix z.
mz2 Matrix that when multiplied by mz1 results in the matrix z. See ‘details’ below.
z Matrix whose effect will be removed from the correlation between x and y.
permute.mz2 Logical argument (TRUE or FALSE) to specify if realize parallel permutation in matrix mz2.
norm.z Logical argument (TRUE or FALSE) to specify if z is standardized within variables (Default norm = FALSE).
put.together List to specify group of traits. Each group specify receive the same weight that one trait outside any group, in the way each group is considered as unique trait (Default put.together = NULL). This argument must be a list, see examples in syncsa.

Details

The null model is based on permutations in the matrix mx2, typically the matrices B, U and Q, except in the function cor.mantel when the permutations are done in one of distance matrix.

Null model described by Pillar et al. (2009) and Pillar & Duarte (2010). For more details on the matrices and the null model, see syncsa.

Value

Obs Correlation between matrices.
p Significance level based on permutations.

Author(s)

Vanderlei Julio Debastiani <vanderleidebastiani@yahoo.com.br>

References


See Also

syncsa, organize.syncsa, mantel, procrustes
flona Hypothetical data for SYNCSA

Description
Hypothetical data for running examples.

Usage
data(flona)

Format
A list with 4 matrices.

community Community data, observations of 59 species in 39 sites.
traits Matrix data of species described by five traits.
phylo Matrix containing phylogenetic distance between species.
environment Three environmental variables for each community.

Examples
data(flona)

matmult.syncsa Matrix multiplication

Description
Internal function to get the matrix product when missing data (NA) is found in matrix Y.

Usage
matmult.syncsa(X, Y)

Arguments
X A matrix, typically the standardized community matrix (W).
Y A matrix.
Details

The function ignore missing data when found in matrix Y. Before multiplication of matrices the missing data in Y are replaced by 0 and multiplication is performed, and then, an adjustment is performed. This adjustment is done by divide each cell of the product matrix by the sum of proportions of nonzero at X with complete data in Y. In SYNCSA context this adjustment is done by divide each cell of the product matrix by the sum of species proportion with trait data in Y. Important, the matrix X must be standardized, in other words, row totals must be equal to 1.

Value

The matrix product.

Author(s)

Vanderlei Julio Debastiani <vanderleidebastiani@yahoo.com.br>

See Also

syncsa, matmult,
Value

- **matrix.w**: Standardized community matrix, where rows are communities and columns species. Row totals (communities) = 1.
- **matrix.q**: Standardized matrix containing the degree of belonging of species in relation to each other. Row totals (species) = 1.

Note

**IMPORTANT**: Species sequence in the community data matrix MUST be the same as the one in the phylogenetic distance matrix. See `organize.syncsa`.

Author(s)

Vanderlei Julio Debastiani <vanderleidebastiani@yahoo.com.br>

References


See Also

`syncsa`, `organize.syncsa`, `belonging`, `matrix.t`, `matrix.x`

Examples

```r
data(ADRS)
matrix.p(ADRS$community, ADRS$phylo)
```

Description

Function to obtain a matrix containing trait averages at community level. For more details, see `syncsa`.

Usage

```r
matrix.t(comm, traits, scale = TRUE, ranks = TRUE, notification = TRUE)
```
Arguments

comm  Community data, with species as columns and sampling units as rows. This matrix can contain either presence/absence or abundance data.

traits  Matrix or data frame of species described by traits, with traits as columns and species as rows.

scale  Logical argument (TRUE or FALSE) to specify if the traits are measured on different scales (Default scale = TRUE). When scale = TRUE traits are measured on different scales and the matrix T is subjected to standardization within each trait. When scale = FALSE traits are measured on the same scale the matrix T is not subjected to standardization.

ranks  Logical argument (TRUE or FALSE) to specify if ordinal variables are convert to ranks (Default ranks = TRUE).

notification  Logical argument (TRUE or FALSE) to specify if notifications of missing observations are shown (Default notification = TRUE).

Value

matriz.w  Standardized community matrix, where rows are communities and columns species. Row totals (communities) = 1.

matriz.b  Matrix of traits, exactly the same data input.

matriz.T  Matrix containing trait averages at community level. If Scale = TRUE the matrix T is standardized within the traits.

Note

IMPORTANT: The sequence species show up in community data matrix MUST be the same as they show up in traits matrix. See organize.syncsa.

Author(s)

Vanderlei Julio Debastiani <vanderleidebastiani@yahoo.com.br>

References


See Also

syncsa, organize.syncsa, matrix.p, matrix.x

Examples

data(ADRS)
matrix.t(ADRS$community, ADRS$traits)
Description

Function to obtain a matrix containing trait-weighted species composition. For more details, see `syncsa`.

Usage

```r
matrix.x(
  comm,
  traits,
  scale = TRUE,
  ranks = TRUE,
  ord,
  notification = TRUE,
  ...
)
```

Arguments

- `comm`: Community data, with species as columns and sampling units as rows. This matrix can contain either presence/absence or abundance data.
- `traits`: Matrix or data frame of species described by traits, with traits as columns and species as rows.
- `scale`: Logical argument (TRUE or FALSE) to specify if the traits are measured on different scales (Default scale = TRUE). When scale = TRUE traits are measured on different scales the matrix of traits is subjected to standardization within each trait, and Gower Index is used to calculate the degree of belonging to the species. When scale = FALSE traits are measured on the same scale the matrix of traits is not subjected to standardization, and Euclidean distance is calculated to determine the degree of belonging to the species.
- `ranks`: Logical argument (TRUE or FALSE) to specify if ordinal variables are convert to ranks (Default ranks = TRUE).
- `ord`: Method to be used for ordinal variables, see `gowdis`, if any method is forneced the rank parameter is not apply.
- `notification`: Logical argument (TRUE or FALSE) to specify if notifications of missing observations are shown (Default notification = TRUE).

Value

- `matriz.w`: Standardized community matrix, where rows are communities and columns species. Row totals (communities) = 1.
matriz.u  Standardized matrix containing the degree of belonging of each species in relation to each other species. Row totals (species) = 1.


Note

IMPORTANT: The sequence species show up in community data matrix MUST be the same as they show up in traits matrix. See organize.syncsa.

Author(s)

Vanderlei Julio Debastiani <vanderleidebastiani@yahoo.com.br>

References


See Also

syncsa, organize.syncsa, belonging, matrix.t, matrix.p, gowdis

Examples

data(ADRS)
matrix.x(ADRS$community, ADRS$traits)

---

optimal  Searching for optimal traits

Description

Maximize trait-convergence assembly patterns (TCAP = roTE), trait-divergence assembly patterns (TDAP = roXE.T), maximize both trait-divergence assembly patterns and trait-convergence assembly patterns (TCAP.TDAP = roXE) or alpha divergence (roR) For more details, see syncsa.

Usage

optimal(
  comm,
  traits = NULL,
  envir = NULL,
  checkdata = TRUE,
  subset.min = 1,
  subset.max = ncol(traits),
)
pattern = NULL,
ro.method = "mantel",
dist = "euclidean",
method = "pearson",
scale = TRUE,
scale.envir = TRUE,
ranks = TRUE,
ord = "metric",
put.together = NULL,
na.rm = FALSE,
notification = TRUE,
progressbar = FALSE
)

## S3 method for class 'optimal'
print(x, ...)

### Arguments

- **comm**: Community data, with species as columns and sampling units as rows. This matrix can contain either presence/absence or abundance data. Alternatively comm can be an object of class `metacommunity.data`, an alternative way to set all data.frames/matrices. When you use the class `metacommunity.data` the arguments traits, envir and put.together must be null. See details.

- **traits**: Matrix data of species described by traits, with traits as columns and species as rows (Default traits = NULL).

- **envir**: Environmental variables for each community, with variables as columns and sampling units as rows (Default envir = NULL).

- **checkdata**: Logical argument (TRUE or FALSE) to check if species sequence in the community data follows the same order as the one in the trait and if sampling units in the community data follows the same order as the one in the environmental matrices (Default checkdata = TRUE).

- **subset.min**: Minimum of traits in each subset (Default subset.min = 1).

- **subset.max**: Maximum of traits in each subset (Default subset.max = ncol(traits)).

- **pattern**: Patterns for maximize correlation, "tcap", "tdap", "tcap.tdap" or "rao" (Default pattern = NULL).

- **ro.method**: Method to obtain the correlation, "mantel" or "procrustes" (Default ro.method = "mantel").

- **dist**: Dissimilarity index, as accepted by vegdist: "manhattan", "euclidean", "canberra", "bray", "kulczynski", "jaccard", "gower", "altGower", "morisita", "horn", "mountford", "raup", "binomial" or "choa".

- **method**: Correlation method, as accepted by cor: "pearson", "spearman" or "kendall".

- **scale**: Logical argument (TRUE or FALSE) to specify if the traits are measured on different scales (Default Scale = TRUE). When scale = TRUE traits are measured on different scales and the matrix T is subjected to standardization within each trait. When scale = FALSE if traits are measured on the same scale and
the matrix $T$ is not subjected to standardization. Furthermore, if scale = TRUE the matrix of traits is subjected to standardization within each trait, and Gower Index is used to calculate the degree of belonging to the species, and if scale = FALSE the matrix of traits is not subjected to standardization, and Euclidean distance is calculated to determine the degree of belonging to the species.

**scale.envir** Logical argument (TRUE or FALSE) to specify if the environmental variables are measured on different scales (Default scale = TRUE). If the environmental variables are measured on different scales, the matrix is subjected to centralization and standardization within each variable.

**ranks** Logical argument (TRUE or FALSE) to specify if ordinal variables are converted to ranks (Default ranks = TRUE).

**ord** Method to be used for ordinal variables, see `gowdis` (Default ord = "metric").

**put.together** List to specify group traits that are added or removed together (Default put.together = NULL). This argument must be a list, see examples.

**na.rm** Logical argument (TRUE or FALSE) to specify if pairwise deletion of missing observations when computing dissimilarities (Default na.rm = FALSE).

**notification** Logical argument (TRUE or FALSE) to specify if notifications of missing observations are shown (Default notification = TRUE).

**progressbar** Logical argument (TRUE or FALSE) to specify if display a progress bar on the R console (Default progressbar = FALSE).

**x** An object of class `optimal`.

**...** Other parameters for the respective functions.

**Details**

Package SYNCSA requires that the species and community sequence in the data.frame or matrix must be the same for all data.frame/matrices. The function `organize.syncsa` organizes the data for the functions of the package, placing the matrices of community, traits and environmental variables in the same order. The function use of function organize.syncsa is not required to run the functions, but is recommended. In this way the arguments comm, traits, envir, as well as the argument put.together, can be specified them as normal arguments or by passing them with the object returned by the function `organize.syncsa` using, in this case only the argument comm. Using the object returned by organize.syncsa, the comm argument is used as an alternative way of entering to set all data.frames/matrices, and therefore the other arguments (traits, envir, and put.together) must be null.

**Value**

- **Subset** Subset of traits that maximizes the correlation.
- **ro** Correlation for the subset of traits.

**Note**

**IMPORTANT**: The sequence species show up in community data matrix MUST be the same as they show up in traits matrix. See details and `organize.syncsa`.
organize.syncsa

Function for organize data for Package SYNCSA

organize.syncsa

Description

Package SYNCSA requires that the species and community sequence in the data.frame or matrix must be the same for all data.frame/matrices.

Usage

organize.syncsa(
    comm,
    traits = NULL,
    phylodist = NULL,
    envir = NULL,
    strata = NULL,
    check.comm = TRUE,
    convert.traits = FALSE,
    ranks = TRUE
)
Arguments

- **comm**: Community data, with species as columns and sampling units as rows.
- **traits**: Matrix or data.frame of species described by traits, with traits as columns and species as rows (Default traits = NULL).
- **phylodist**: Matrix containing phylogenetic distance between species. Must be a complete matrix (not a half diagonal matrix). This matrix can be larger than community data (more species) as long as it has at least all species that are in community data (Default phylodist = NULL).
- **envir**: Environmental variables for each community, with variables as columns and sampling units as rows (Default envir = NULL).
- **strata**: Strata named vector to specify restricting permutations within species groups (Default strata = NULL).
- **check.comm**: Logical argument (TRUE or FALSE) to remove sampling units and species with total sums equal or less than zero (Default check.comm = TRUE).
- **convert.traits**: Logical argument (TRUE or FALSE) to convert factor traits in dummy traits and/or convert ordinal variables in numeric (see ranks argument) (Default convert.traits = FALSE).
- **ranks**: Logical argument (TRUE or FALSE) to specify if ordinal variables are convert to ranks. If ranks = TRUE all ordered variable are replaced by their ranks and if ranks = FALSE all ordinal variables are simply treated as continuous variables (Default ranks = TRUE).

Details

The function organizes the data for the functions of the package SYNCSA, placing the matrices of community, traits, phylogenetic distance, environmental variables and strata vector in the same order. The function use as reference the community data for organize all data.frame or matrices in the same order that the sampling units names and species names found in community data set. For this all data sets entered must be correctly named, with rows and columns named. The matrices phylodist, traits, envir can be larger than community data (more species and/or more sampling units) as long as it has at least all species and/or sampling units that are in community data. The function organizes the data despite the absence of one of the data.frames or matrices, provided that the community data had been entered. Unspecified data will appear as NULL.

When trait is a data.frame with differents types of variables correctly identified and the argument convert.traits is TRUE factor traits are expanded in dummy traits and ordinal variables are converted in numeric according to ranks argument.

The strata must be a named vector. The strata vector is a vector for restrict permutation within species groups, insofar as the SYNCSA package the null models are based in permutation of species rather than permutation of sample units.

Value

A object of class metacommunity.data (also of the class list) with the data.frames or matrices:
- **call**: The arguments used.
- **community**: Community data.
part.cor

Description

First-order partial correlation coefficient

Usage

part.cor(rxy, rxz, ryz)

Arguments

rxy Correlation between xy
rxz Correlation between xz
ryz Correlation between yz
Value

The first-order partial correlation coefficient.

Author(s)

Vanderlei Julio Debastiani <vanderleidebastiani@yahoo.com.br>

See Also

syncsa, cor.matrix,

Principal Components Analysis (PCA) with NA (missing data)

Description

The function use the option "pairwise.complete.obs" (in function cor) for calculate the correlation. The correlation between each pair of variables is computed using all complete pairs of observations on those variables.

Usage

pca(data)

## S3 method for class 'pcasyncsa'
plot(
  x,
  show = c("variables", "individuals"),
  axis = c(1, 2),
  xlab = axis[1],
  ylab = axis[2],
  arrows = TRUE,
  text = TRUE,
  points = FALSE,
  ...
)

Arguments

data A data frame or matrix with individuals in rows and variables in columns.

x A object of class pcasyncsa.

show Draw "variables" or "individuals".

axis Axis for draw, must have length equal to two (Default axis = c(1, 2)).

xlab Text for x label (Default xlab = axis[1]).

ylab Text for y label (Default ylab = axis[2]).
Logical argument (TRUE or FALSE) to specify if arrows are showed for variables (Default arrows = TRUE).

text

Logical argument (TRUE or FALSE) to specify if text are showed for individuals (Default text = TRUE).

points

Logical argument (TRUE or FALSE) to specify if points are showed for individuals (Default points = FALSE).

... Parameters for plot function.

Value

decomposition list with the results of decomposition of correlation matrix.
eigenvalues Data frame containing all the eigenvalues, the percentage of inertia and the cumulative percentage of inertia.
individuals Coordinates for the individuals.
variables Correlation between original variables and axes.

Author(s)

Vanderlei Julio Debastiani <vanderleidebastiani@yahoo.com.br>

See Also

syncsa syncsa

Examples

data(ADRS)
traits<-ADRS$traits
# Some NA
traits[c(1,5),1]<-NA
traits[3,2]<-NA
traits
res<-pca(traits)
res
plot(res, show = "variables", arrows = TRUE)
plot(res, show = "individuals", axis = c(1, 2), text = TRUE)
plot(res, show = "individuals", text = FALSE, points = TRUE)

Usage

permut.row.matrix(data, strata = NULL, seqpermutation = NULL)
**permut.vector**

Arguments

- **data**: A matrix.
- **strata**: Strata vector to specify restricting permutations in rows, must be the same length of number of rows in data matrix (Default strata = NULL).
- **seqpermutation**: A set of predefined permutation vector (Default seqpermutation = NULL).

Value

- **permut.matrix**: The matrix permuted.
- **samp**: The sequence of permutation.

Author(s)

Vanderlei Julio Debastiani <vanderleidebastiani@yahoo.com.br>

See Also

- `syncsa`, `permut.vector`

---

**permut.vector**: *Permutate a vector*

Description

Internal function to permutate a vector of size n using the function `shuffleSet`.

Usage

```r
permut.vector(n, strata = NULL, nset = 999)
```

Arguments

- **n**: The length of vector.
- **strata**: A vector to specify restricting permutations.
- **nset**: The number of permutations to generate for the set (Default strata = NULL).

Value

A matrix of permutations, where each row is a separate permutation.

Author(s)

Vanderlei Julio Debastiani <vanderleidebastiani@yahoo.com.br>

See Also

- `syncsa`, `permut.row.matrix`
procrustes.syncsa

Procrustes and Partial Procrustes correlations.

Description

Function to obtain the Procrustes correlation between two matrices and partial Procrustes correlation between three matrices.

Usage

procrustes.syncsa(x, y)

procrustes.partial(x, y, z)

Arguments

x, y Matrix that will be correlated.

z Matrix whose effect will be removed from the correlation between x and y.

Details

The function procrustes.syncsa is a small change in the function procrustes. The function uses a correlation-like statistic derived from the symmetric Procrustes sum of squares. Partial Procrustes correlations is obtained by Procrustes correlation of residuals between each variable in z being used as linear predictor of all variables in x and y. For more details, see syncsa.

Value

Procrustes correlation between matrices.

Author(s)

Vanderlei Julio Debastiani <vanderleidebastiani@yahoo.com.br>

References


See Also

syncsa, cor.matrix
**ProgressBAR**

**Text Progress Bar**

**Description**

Text progress bar in the R console. See `txtProgressBar`.

**Usage**

```r
ProgressBAR(n, N, ...) 
```

**Arguments**

- `n` Number of the current progress.
- `N` Total number of cases.
- `...` Other parameters for the `txtProgressBar` function.

**Author(s)**

Vanderlei Julio Debastiani <vanderleidebastiani@yahoo.com.br>

---

**rao.diversity**

Rao's quadratic entropy

**Description**

Calculates Rao's quadratic entropy, functional and phylogenetic redundancy.

**Usage**

```r
rao.diversity( 
  comm, 
  traits = NULL, 
  phylodist = NULL, 
  checkdata = TRUE, 
  ord = "metric", 
  put.together = NULL, 
  standardize = TRUE, 
  ... 
)
```
Arguments

comm Community data, with species as columns and sampling units as rows. This matrix can contain either presence/absence or abundance data. Alternatively, comm can be an object of class metacommunity.data, an alternative way to set all data.frames/matrices. When you use the class metacommunity.data the arguments traits, phylodist and put.together must be null. See details.

traits Data frame or matrix data of species described by traits, with traits as columns and species as rows (Default traits = NULL).

phylodist Matrix containing phylogenetic distance between species (Default phylodist = NULL).

checkdata Logical argument (TRUE or FALSE) to check if species sequence in the community data follows the same order as the one in the trait and in the phylodist matrices (Default checkdata = TRUE).

ord Method to be used for ordinal variables, see gowdis (Default ord = "metric").

put.together List to specify group of traits. Each group specify receive the same weight that one trait outside any group, in the way each group is considered as unique trait (Default put.together = NULL). This argument must be a list, see examples in syncsa.

standardize Logical argument (TRUE or FALSE) to specify if standardize phylogenetic distance to range into range 0 to 1. (Default standardize = TRUE).

... Parameters for gowdis function.

Details

Rao’s quadratic entropy is a measure of diversity of ecological communities defined by Rao (1982) and is based on the proportion of the abundance of species present in a community and some measure of dissimilarity among them. The dissimilarity range from 0 to 1 and is based on a set of specified functional traits or in the phylogenetic dissimilarity.

For the trait data, the function calculates the square root of the one-complement of Gower’s similarity index, in order to have a dissimilarity matrix with Euclidean metric properties. Gower’s index ranges from 0 to 1 and can handle traits measured indifferent scales. When the species are completely different in terms of their traits, Rao quadratic entropy is equivalent to the Gini-Simpson index. Traits data can be numeric, factor or ordered factor. For this be considered traits data must be of data.frame class and containing each variable type determined. For additional details and requirements of function please see gowdis.

Functional redundancy is defined purely as the difference between species diversity and Rao’s quadratic entropy based on their functional dissimilarity (de Bello et al. 2007). The same definition is used for phylogenetic redundancy.

Package SYNCSA requires that the species and community sequence in the data.frame or matrix must be the same for all dataframe/matrices. The function organize.syncsa organizes the data for the functions of the package, placing the matrices of community, traits, phylogenetic distance in the same order. The function use of function organize.syncsa is not required for run the functions, but is recommended. In this way the arguments comm, traits, phylodist, as well as the argument put.together, can be specified them as normal arguments or by passing them with the object returned by the function organize.syncsa using, in this case only the argument comm. Using the object
rao.diversity

returned by organize.syncsa, the comm argument is used as an alternative way of entering to set all
data.frames/matrices, and therefore the other arguments (traits, phylodist and put.together) must be
null.

Value

Simpson  Gini-Simpson index within each community (equivalent to Rao quadratic en-
tropy with null, crisp, similarities).
FunRao    Rao quadratic entropy within each community, considering trait distance.
FunRedundancy Functional redundancy in each community.
PhyRao    Rao quadratic entropy within each community, considering phylogenetic dis-
tance.
PhyRedundancy Phylogenetic redundancy in each community.

Note

IMPORTANT: The sequence species show up in community data matrix MUST be the same as
they show up in traits and phylodist matrices. See details and organize.syncsa.

Author(s)

Vanderlei Julio Debastiani <vanderleidebastiani@yahoo.com.br>

References

de Bello, F.; Leps, J.; Lavorel, S. & Moretti, M. (2007). Importance of species abundance for as-
essment of trait composition: an example based on pollinator communities. Community Ecology,
8, 163:170.
Biology, 21, 24:43.

See Also

organize.syncsa, gowdis, syncsa

Examples

data(ADRS)
rao.diversity(ADRS$community)
rao.diversity(ADRS$community, traits = ADRS$traits)
**Description**

This function integrates several steps for the analysis of phylogenetic assembly patterns and their links to traits and ecological processes in a metacommunity (Pillar et al. 2009, Pillar & Duarte 2010, Debastiani & Pillar 2012). The function implement methods that have been available in the SYNCSA application written in C++ (by Valerio Pillar, available at http://ecoqua.ecologia.ufrgs.br/SYNCSA.html). See details.

**Usage**

```r
syncsa(
  comm,
  traits = NULL,
  phylodist = NULL,
  envir = NULL,
  checkdata = TRUE,
  ro.method = "mantel",
  method = "pearson",
  dist = "euclidean",
  scale = TRUE,
  scale.envir = TRUE,
  ranks = TRUE,
  ord,
  put.together = NULL,
  na.rm = FALSE,
  strata = NULL,
  permutations = 999,
  parallel = NULL,
  notification = TRUE
)
```

```r
## S3 method for class 'syncsa'
print(x, ...)
```

**Arguments**

- **comm**: Community data, with species as columns and sampling units as rows. This matrix can contain either presence/absence or abundance data. Alternatively comm can be an object of class metacommunity.data, an alternative way to set all data.frames/matrices. When you use the class metacommunity.data the arguments traits, phylodist, envir and put.together must be null. See details.

- **traits**: Data frame or matrix data of species described by traits, with traits as columns and species as rows (Default traits = NULL).
<table>
<thead>
<tr>
<th>Argument</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>phylodist</td>
<td>Matrix containing phylogenetic distance between species. Must be a complete matrix, not a half diagonal matrix (Default phylodist = NULL).</td>
</tr>
<tr>
<td>envir</td>
<td>Environmental variables for each community, with variables as columns and sampling units as rows (Default envir = NULL).</td>
</tr>
<tr>
<td>checkdata</td>
<td>Logical argument (TRUE or FALSE) to check if species sequence in the community data follows the same order as the one in the trait and in the phylodist matrices and if sampling units in the community data follows the same order as the one in the environmental data (Default checkdata = TRUE).</td>
</tr>
<tr>
<td>ro.method</td>
<td>Method to obtain the correlation, &quot;mantel&quot; or &quot;procrustes&quot; (Default ro.method = &quot;mantel&quot;).</td>
</tr>
<tr>
<td>method</td>
<td>Mantel correlation method, as accepted by cor: &quot;pearson&quot;, &quot;spearman&quot; or &quot;kendall&quot; (Default method = &quot;pearson&quot;).</td>
</tr>
<tr>
<td>dist</td>
<td>Dissimilarity index used for Mantel correlation, as accepted by vegdist: &quot;manhattan&quot;, &quot;euclidean&quot;, &quot;canberra&quot;, &quot;bray&quot;, &quot;kulczynski&quot;, &quot;jaccard&quot;, &quot;gower&quot;, &quot;alt-Gower&quot;, &quot;morisita&quot;, &quot;horn&quot;, &quot;mountford&quot;, &quot;raup&quot;, &quot;binomial&quot; or &quot;chao&quot; (Default dist = &quot;euclidean&quot;).</td>
</tr>
<tr>
<td>scale</td>
<td>Logical argument (TRUE or FALSE) to specify if the traits are measured on different scales (Default Scale = TRUE). When scale = TRUE traits are measured on different scales the the matrix T is subjected to standardization within each trait. When scale = FALSE traits are measured on the same scale and the matrix T is not subjected to standardization. Furthermore, if scale = TRUE the matrix of traits is subjected to standardization within each trait, and Gower Index is used to calculate the degree of belonging to the species, and if scale = FALSE the matrix of traits is not subjected to standardization, and Euclidean distance is calculated to determine the degree of belonging to the species.</td>
</tr>
<tr>
<td>scale.envir</td>
<td>Logical argument (TRUE or FALSE) to specify if the environmental variables are measured on different scales. If the enviromental variables are measured on different scales, the scale.envir = TRUE the matrix with enviromental variables is subjected to centralization and standardization within each variable. (Default scale.envir = TRUE).</td>
</tr>
<tr>
<td>ranks</td>
<td>Logical argument (TRUE or FALSE) to specify if ordinal variables are convert to ranks (Default ranks = TRUE).</td>
</tr>
<tr>
<td>ord</td>
<td>Method to be used for ordinal variables, see gowdis.</td>
</tr>
<tr>
<td>put.together</td>
<td>List to specify group of traits. Each group specify receive the same weight that one trait outside any group, in the way each group is considered as unique trait (Default put.together = NULL). This argument must be a list, see examples.</td>
</tr>
<tr>
<td>na.rm</td>
<td>Logical argument (TRUE or FALSE) to specify if pairwise distances should be deleted in cases of missing observations (Default na.rm = FALSE).</td>
</tr>
<tr>
<td>strata</td>
<td>Strata named vector to specify restricting permutations within species groups (Default strata = NULL).</td>
</tr>
<tr>
<td>permutations</td>
<td>Number of permutations in assessing significance.</td>
</tr>
<tr>
<td>parallel</td>
<td>Number of parallel processes. Tip: use parallel::detectCores() (Default parallel = NULL).</td>
</tr>
<tr>
<td>notification</td>
<td>Logical argument (TRUE or FALSE) to specify if notification of missing observations should to be shown (Default notification = TRUE).</td>
</tr>
</tbody>
</table>
x An object of class syncsa.

... Other parameters for the respective functions.

Details

Package SYNCSA requires that the species and community sequence in the data.frame or matrix must be the same for all dataframe/matrices. The function organize.syncsa organizes the data for the functions of the package, placing the matrices of community, traits, phylogenetic distance, environmental variables and strata vector in the same order. The function use of function organize.syncsa is not required for run the functions, but is recommended. It requires data organized into the following matrices: (1) the presences or abundances of species in a set of communities (W); (2) the phylogenetic pairwise dissimilarities of these species (DF); (3) a set of functional traits describing the species (B), which may be a mixture of binary and quantitative traits (continual and ordinal), but not nominal ones (these should be expanded into binary traits); and (4) the ecological gradient of interest, which may be one or more factors to which the communities respond or ecosystem effects of the communities (E). In this way the arguments comm, traits, phylodist, envir, as well as the arguments put.together and strata, can be specified as normal arguments or by passing them with the object returned by the function organize.syncsa using, in this case only the argument comm. Using the object returned by organize.syncsa, the comm argument is used as an alternative way of entering to set all data.frames/matrices, and therefore the other arguments (traits, phylodist, envir, put.together and strata) must be null.

Correlations

The function computes several correlations (Mantel or Procrustes) that express trait-convergence assembly patterns (TCAP), trait-divergence assembly patterns (TDAP), and phylogenetic signal in functional traits at the species pool level and at the metacomunity level. This function also generates P-values by permutation testing based on null models (Pillar et al. 2009, Pillar & Duarte 2010).

ro(TE)

This correlation refers to trait-convergence assembly patterns related to the ecological gradient (TCAP, Pillar et al. 2009). For evaluating TCAP, by matrix multiplication we define $T = WB$, which with previous standardization of $W$ to unit column totals will contain the trait averages in each community. The elements in $T$ are community weighted mean values or community functional parameters (Violle et al. 2007). Standardization of the traits (rows) in $T$ is needed if the trait set contains traits measured with different scales. By using matrix correlation, we evaluate how the trait patterns in $T$ are associated to ecological gradients in $E$. For relating $T$ to $E$, using Mantel correlation we define a distance matrix of the communities ($DT$) using $T$, and another distance matrix of the community sites ($DE$) using $E$. The matrix correlation $ro(TE) = ro(DT;DE)$ measures the level of congruence between TCAP and $E$. A strong correlation $ro(TE)$ indicates the factors directly or indirectly represented in $E$ are involved in ecological filtering of species that, at least for the traits considered in the analysis, consistently produce trait-convergence assembly patterns along the gradient comprising the metacommunity.

ro(XE) and ro(XE,T)

These matrix correlations refer to trait-divergence assembly patterns related to the ecological gradient (TDAP, Pillar et al. 2009). For the identification of TDAP, in a first step the species pairwise similarities (in the range 0 to 1) in matrix $SB$ based on traits in $B$ are used to define matrix $U$ with degrees of belonging of species to fuzzy sets. By matrix multiplication $X = WU$ will contain the species composition of the communities after fuzzy-weighting by their trait similarities (each row
in X will refer to a species). Matrix X expresses both TCAP and TDAP (Pillar et al. 2009). By using matrix correlation, we evaluate how the trait patterns in X (TCAP and TDAP) are associated to ecological gradients in E. For relating X to E, we define a distance matrix of the communities (DX) using X, and another distance matrix of the community sites (DE) using E. The matrix correlation ro(XE) = ro(DX,DE) between X and E is defined. We then remove the trait-convergence component ro(TE) from ro(XE) by computing the partial matrix correlation ro(XE,T), which measures the level of congruence between TDAP and E. Trait-divergence assembly patterns (TDAP, Pillar et al. 2009) may result from community assembly processes related to biotic interactions (Stubbs & Wilson 2004; Wilson 2007).

ro(PE)
This matrix correlation refers to the phylogenetic structure related to the ecological gradient comprising the metacommunity. The phylogenetic pairwise dissimilarities in DF are transformed into similarities and used to define degrees of belonging qij to fuzzy sets. This is analogous to the definition of functional fuzzy sets (Pillar & Orloci 1991; Pillar et al. 2009). Based on the phylogenetic similarities, every species i among s species in the metacommunity specifies a fuzzy set to which every species j (j = 1 to s species, including species i) belongs with a certain degree of belonging in the interval [0, 1]. In our definition, each row in matrix Q with the degrees of belonging must add to unit, i.e., the degrees of belonging of a given species across the fuzzy sets are standardized to unit total. By matrix multiplication P = WQ will contain the composition of the communities after fuzzy-weighting of species presences or abundances by the species’ phylogenetic similarities. Each column in matrix P holds the phylogenetic structure of a community. The standardization of Q is essential for the community totals in each column in W remaining the same in P. Further, matrix W is adjusted to unit column totals prior to multiplication, so that the total richness or abundance within each community in W will be standardized. Matrix correlation ro(PE) = ro(DP,DE) measures the strength of the association between community distances based on their phylogenetic structure in DP and distances based on their ecological conditions (DE). Further, P can be explored for phylogenetic patterns at the metacommunity level by using, e.g., ordination techniques.

ro(PT) and ro(PX.T)
These matrix correlations measure phylogenetic signal at the metacommunity level related to TCAP and to TDAP. We define phylogenetic signal at the metacommunity level related to TCAP (PSMT) as the correlation between the phylogenetic structure described in matrix P and the trait-convergence structure described in matrix T. For this, a proper distance matrix (e.g. Euclidean distances) of communities (DP) is computed using P and another distance matrix of the same communities (DT) is computed using T. Then matrix correlation ro(PT) = ro(DP,DT) will measure the level of congruence between variation in P and T, which is a measure of PSMT. A strong phylogenetic signal at the metacommunity level is expected when communities that are more similar in terms of phylogenetic structure are also similar regarding their average trait values. We also define phylogenetic signal at the metacommunity level related to TDAP (PSMX.T) as the partial matrix correlation ro(PX.T) = ro(DP,DX,DT) between community distances DP computed on phylogenetic structure and community distances DX computed on species composition after fuzzy-weighting by the species, or trait similarities, removing the effect of TCAP (DT). This is analogous to TDAP.

ro(BF)
This matrix correlation measures phylogenetic signal at the species pool level (PSS, Pillar & Duarte 2010). We define PSS as the matrix correlation ro(FB) = ro(DF,DB) between species phylogenetic dissimilarities (already defined as matrix DF) and species trait dissimilarities (derived from already defined matrix SB) computed on any number of traits from matrix B. The species pool refers to the species present in the metacommunity.
Additional matrix correlations
The matrix correlations $\rho(TE, P)$ and $\rho(XE, P)$ are also computed, which may be useful for evaluating causal models in path analysis.

Mantel correlations
The Mantel and Partial Mantel statistics are calculated simply as the correlation entries the dissimilarity matrices, using standard Mantel test (see \texttt{mantel} and \texttt{cor.mantel}). Partial Mantel statistic use paired correlation between the three matrices and obtains the partial correlation using the formula of first-order partial correlation coefficient. The significances are obtained using a different procedure than standard Mantel test, see section Testing against null models below.

Procrustes correlations
The Procrustes correlation uses symmetric Procrustes as a measure of concordance between the data matrices (see \texttt{procrustes} and \texttt{cor.procrustes}). Procrustes procedure use rotation, translation, and rescaling for minimizing sum of squared differences between two data sets. The correlation of Procrustes is calculated as the statistic derived from the symmetric Procrustes sum of squares, representing the optimal fit between the two data matrices. Partial Procrustes correlation is obtained by Procrustes correlation between residuals matrices. Firstly one Principal Components Analysis (PCA, see \texttt{prcomp}) is performed in the matrix $Z$ for dimensionality reduction. The max number of axis kept in the analysis is the number of sampling units divided by 2, this axes of PCA represent the total variation in the $Z$ matrix. After the kept axes are used as predictor in one linear model for each variable of the matrices $X$ and $Y$. For this a linear model is build using as response one variable of $X$ (same via for $Y$ matrix) and as predictor all remaining axes of PCA, after model fitted and the residual are extracted with the aim of form the residual matrix. The linear model is repeated in the other variables, only with the changed the response variable. The same procedure is performed in the matrix $Y$. Both residual matrices are submitted to Procrustes analysis and the statistic is returned as a partial correlation, the Partial Procrustes statistic. The significances are obtained using the same procedure than Mantel test, see section Testing against null models below.

Testing against null models
All the matrix correlations are tested against null models. The null model is defined according to the correlation being tested. Usually in the SYNCSA package the null models are based in permutation of species rather than permutation of sample units. For $\rho(TE)$, each permutation generates a random matrix $T$ calculated after the permutation among the species vectors in matrix $B$. For $\rho(XE)$ and $\rho(XE, T)$, each permutation generates a random matrix $X$ after the permutation among species fuzzy sets (rows) in matrix $U$. For $\rho(PE)$, $\rho(PT)$, and $\rho(PX, T)$, each permutation generates a random matrix $P$ after the permutation among species fuzzy sets (rows) in matrix $Q$. For $\rho(BF)$, a conventional Mantel test is performed with dissimilarity matrices $DF$ and $DB$. Analogous null models are used for testing the additional matrix correlations; that is, the same null model for $\rho(TE)$ is used for $\rho(TE, P)$, the same model for $\rho(XE)$ is used for $\rho(XE, P)$. The permutation can be restrict within species groups specifying the strata argument.

Traits types
Traits data can be numeric, factor or ordered factor. For this be considered in the analyses traits data must be of data.frame class and containing each variable type determined. Gower index is used to calculate the similarity between species, using the function \texttt{gowdis} of package FD. For additional details and requirements of function please see \texttt{gowdis}.

Missing data
The functions ignore missing data when specified. In the case of direct multiplication of matrices the missing data are replaced by 0, ignoring the cell with missing value. For the matrix $T = WB$ an
adjustment is done by divide each cell of the product matrix (T) by the sum of species proportion with trait data in B. Result matrices are shown without missing values. Where the matrices are calculated using a dissimilarity index (matrix U and correlations between matrices) the missing data are ignored as in `vegdist` function. In some cases the dissimilarity matrices obtained by the function `vegdist` still contain some missing values. In these cases the rest of the procedure will be affected. In these cases you can find solutions in impute the missing values.

**Error messenger and options**

The data pass by several check points that can produce error messenger. The matrices or data frames must be contain only numeric, binary or ordinal variables, in the way that nominal variable should be expanded into binary (see `var.dummy`). For enhance the code speed some functions use by default matrix algebra, this option can produce error under certain circumstances. This global option can be changed using options("SYNCSA.speed" = FALSE). If SYNCSA.speed = TRUE for use matrix algebra and if SYNCSA.speed = FALSE use not another function of same procedure.

**Value**

<table>
<thead>
<tr>
<th>call</th>
<th>The arguments used.</th>
</tr>
</thead>
<tbody>
<tr>
<td>notes</td>
<td>Some notes about the statistics.</td>
</tr>
<tr>
<td>statistics</td>
<td>Correlations roTE, roXE, roPE, roPT, roPX.T, roXE.T, roTE.P, roXE.P and roBF, and their significance levels based on permutations.</td>
</tr>
<tr>
<td>matrices</td>
<td>The matrices produced for the functions, see details.</td>
</tr>
<tr>
<td>FunRao</td>
<td>Rao quadratic entropy within each community, considering trait distance.</td>
</tr>
<tr>
<td>weights</td>
<td>Weight for each trait.</td>
</tr>
</tbody>
</table>

**Note**

The function calculates the correlations despite the lack of one of the matrices, provided that community data had been entered. Correlations including unspecified matrices will appear with NA.

**IMPORTANT:** The sequence of species in the community data matrix MUST be the same as that in the phylogenetic distance matrix and in traits matrix. Similarly, the sequence of communities in the community data matrix MUST be the same as that in the environmental data. See details and `organize.syncsa`.

**Author(s)**

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**References**


See Also

organize.syncsa, matrix.t, matrix.x, matrix.p, optimal, rao.diversity, cor.matrix, var.type, var.dummy

Examples

data(ADRS)
synsca(ADRS$community, ADRS$traits, ADRS$phylo, ADRS$envir, permutations = 99)
data(flona)
put.together<-list(c("fol", "sem" ), c("tam", "red" ))
put.together
res<.synsca(flona$community, flona$traits, envir = flona$environment, put.together = put.together, permutations = 99)
res$weights

var.dummy

Generate dummy variable

Description

Function to expand factor variables in dummy variables in a data.frame. See details.

Usage

var.dummy(data)

Arguments

data A data.frame.

Details

Variables in the data.frame of class factor is expanded in dummy variables. Each level of factor produce a new column in the dataframe, with presence (1) or absence (0) of level. The name of columns is a combination of orginal variable name plus the level separate by underscore (_). Ordered factor and character class are not expanded.
Description

Function extracted (with small changes) of the function `gowdis` to check the type of variables in a data.frame or matrix.

Usage

```r
var.type(data)
```

Arguments

- `data` A data.frame or matrix.

Value

A vector with the variable types, where 'c' is continuous/numeric, 'o' is ordinal, 'b' is binary, 'n' is nominal and 'f' is factor.

Author(s)

Vanderlei Julio Debastiani <vanderleidebastiani@yahoo.com.br>

See Also

`syncsa`, `organize.syncsa`, `var.type`
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