Package ‘betaper’

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R topics documented:

adonis_pertables ........................................ 2
Amazonia ................................................. 4
cca_pertables ........................................... 5
HCP ......................................................... 7
mantel_pertables ......................................... 7
pertables ................................................... 10
rda_pertables ............................................. 12
Index ....................................................... 15
Function to assess the effects of taxonomic uncertainty on permutational multivariate analysis of variance using distance matrices

Description

This function assesses the effects of taxonomic uncertainty on the R2 coefficients and the p-values of a permutational multivariate analysis of variance using distance matrices.

Usage

`adonis_pertables(formula = X ~ ., data, permutations = 5, method = "bray")`

## S3 method for class 'adonis_pertables'
plot(x, ...)

Arguments

- **formula**: A typical model formula such as `Y ~ A + B*C`, but where `Y` is a pertables object (i.e. a list of simulated community data matrices obtained with `pertables`; 'A', 'B', and 'C' may be factors or continuous variables.
- **data**: The data frame from which 'A', 'B', and 'C' would be drawn.
- **permutations**: Number of replicate permutations used for the hypothesis tests (F tests) for each simulated community data matrices obtained with `pertables`.
- **method**: The name of any method used in `vegdist` to calculate pairwise distances.
- **x**: `adonis_pertables` object to plot.
- **...**: Additional graphical parameters passed to plot.

Value

`adonis_pertables` returns an object of class `adonis_pertables`, basically a list with the following components:

- **raw**: An object of class `adonis`, i.e. the results of applying `mantel` to the original biological data table without the unidentified species. This includes p-values for each explanatory variable showing the probability of obtaining the same F statistic under different scenarios of taxonomic uncertainty.

- **simulation**: A list with the results of the simulation: F, i.e. a data.frame with all the simulated pseudo-F (columns) for each explanatory variable (rows); R2, i.e. a data.frame with all the simulated R2 coefficients (columns) for each explanatory variable (rows); pvalue, i.e. a data.frame with all the simulated p-values (columns) for each explanatory variable (rows); R2.quant, i.e. a data.frame with the summary of R2 by quantiles; p.quant, i.e. a data.frame with the summary of pvalue by quantiles.

The objects of class `adonis_pertables` have print and plot S3 methods for a simple access to results. See the examples.
Author(s)
Luis Cayuela and Marcelino de la Cruz

References

See Also
pertables, adonis

Examples

data(Amazonia)
data(soils)

# Define a new index that includes the terms used in the Amazonia dataset to define 
# undetermined taxa at different taxonomic levels
index.Amazon <- c(paste("sp.", rep(1:20, sep=""), "Indet.", "indet."))

# Generate a pertables object (i.e. a list of biological data tables simulated from taxonomic 
# uncertainty)
## Not run:
Amazonia100 <- pertables(Amazonia, index=index.Amazon, nsim=100)
# Assess the effects of taxonomic uncertainty on a PERMANOVA (i.e., adonis) test:
Amazonia.adonis <- adonis_pertables(Amazonia100 ~ Ca + K + Mg + Na, data=soils)
Amazonia.adonis
plot(Amazonia.adonis)

## End(Not run)
# Fast example for Rcheck
# For example
Amazonia4.p2 <- pertables.p2(Amazonia[1:50,], index=index.Amazon, nsim=4, ncl=2, iseed=4)
set.seed(2)
Amazonia.adonis <- adonis_pertables(Amazonia4.p2 ~ Ca + K + Mg + Na, data=soils)

Amazonia.adonis
plot(Amazonia.adonis)
Amazonia

Tree abundance and soil data in Western Amazonia

Description

The Amazonia data frame has tree counts in nine 0.16-hectare inventory plots in Western Amazonia. soils contains data on soil cations at each location.

Usage

data(Amazonia)
data(soils)

Format

Amazonia is a data frame with 1188 observations (species) and 12 columns (taxonomic description and sites). The three first columns refer to family, genus and specific species Latin names. Columns 4 to 12 have tree abundance data for nine inventory plots.

soils is a data frame with 9 observations (inventory plots) and 4 columns (variables). Soil variables (Ca, K, Mg, Na) are given in cmol/kg.

Details

Data from Western Amazonia includes tree inventories at nine lowland sites (approximately 100-150 m above sea level) near Iquitos, Peru. The sites were selected to represent regional variations in geology and were distributed along a soil nutrient gradient ranging from poor loamy soils to richer clayey soils. Each inventory consisted of 20 x 20 m plots (0.16 ha total area) distributed along 1.3-km transects. At each site, K. Ruokolainen and colleagues identified to species or morphospecies all woody, free-standing stems of > 2.5 cm dbh. The full inventories sampled 3980 individuals from 1188 species or morphospecies.

References


Examples

data(Amazonia)
data(soils)
Description

This function assesses the effects of taxonomic uncertainty on two widely used parameters of a [Partial] Constrained Correspondence Analysis, i.e. the 'percentage explained variance' (sometimes referred to as R-squared) and the pseudo-F.

Usage

```r
cca_pertables(formula, data, scale = FALSE, ...
## S3 method for class 'cca_pertables'
plot(x, pch = 18, ...)
```

Arguments

- `formula`: Model formula, where the left hand side gives a `pertables` object (i.e. a list of simulated community data matrices obtained with `pertables`, right hand side gives the constraining variables, and conditioning variables can be given within a special function `Condition`.
- `data`: Data frame containing the variables on the right hand side of the model formula.
- `scale`: Scale species to unit variance (like correlations).
- `x`: `cca_pertables` object to plot.
- `pch`: Plotting 'character', i.e., symbol to use in the CCA plot. See `points` for examples of use of this graphical argument.
- `...`: Additional graphical parameters passed to `plot`.

Details

This function is a wrapper to submit a `pertables` object to `cca` function of the `vegan` package. See the documentation of `cca` for details about formula and Condition use.

Value

- `cca_pertables` returns an object of class `cca_pertables`, basically a list with the following components:
  - `raw`: An object of class `cca`. The results of applying `cca` to the original biological data table without the unidentified species.
  - `simulation`: A list with the results of the simulation: results, i.e. a data.frame with all the simulated R-squared and pseudo-F values; `cca.quant`, i.e. a data.frame with the summary of results by quantiles; `sites`, i.e. a list with the scores of the sites of all the simulated data tables and `biplot`, i.e. a list with the scores of the environmental data in all the analyses.
The objects of class `cca_pertables` have `print` and `plot` S3 methods for a simple access to results. See the examples.

**Author(s)**

Luis Cayuela and Marcelino de la Cruz

**References**


**See Also**

`pertables`, `cca`

**Examples**

```r
# Define a new index that includes the terms used in the `Amazonia` dataset to define # undetermined taxa at different taxonomic levels
index.Amazon <- c(paste("sp.", rep(1:20), sep=""), "Indet.", "indet."")

# Generate a pertables object (i.e. a list of biological data tables simulated from taxonomic # uncertainty)
## Not run:
Amazonia100 <- pertables(Amazonia, index=index.Amazon, nsim=100)
# Assess the effects of taxonomic uncertainty on a CCA analysis of biological data explained # by all the environmental variables of the soil data:
Amazonia.cca <- cca_pertables(Amazonia100 ~., data=soils)
Amazonia.cca
plot(Amazonia.cca)
## End(Not run)

# Fast example for Rcheck
Amazonia4.p2 <- pertables.p2(Amazonia[1:50,], index=index.Amazon, nsim=4, ncl=2, iseed=4)
set.seed(2)
Amazonia.cca <- cca_pertables(Amazonia4.p2 ~., data=soils)
Amazonia.cca
```

**HCP**

Tree counts in tropical montane forest fragments

**Description**

HCP has tree abundance data from 16 forest fragments located in the Highlands of Chiapas, southern Mexico. HCP.coords contains the geographical UTM coordinates for the 16 forest fragments’ centroids.

**Usage**

```r
data(HCP)
data(HCP.coords)
```

**Format**

HCP is a data frame with 231 observations and 19 variables. The three first columns contain family, genus and specific species Latin names. Columns 4 to 19 have tree abundance data for the 16 forest fragments. HCP.coords is a data frame with two columns and 16 rows.

**References**


**Examples**

```r
data(HCP)
data(HCP.coords)
```

**mantel_pertables**

*Function to assess the effects of taxonomic uncertainty on Mantel tests*

**Description**

This function assesses the effects of taxonomic uncertainty on the coefficient of correlation and the p-values of a Mantel test.
mantel_pertables

Usage

mantel_pertables(pertab, env, dist.method = "bray", binary = FALSE, cor.method = "pearson", permutations = 100)

## S3 method for class 'mantel_pertables'
plot(x, xlab = "Environmental distance", ylab = "Sørensen's similarity index", pch = 19, ...)

Arguments

- **pertab**: A pertables object (i.e. a list of simulated community data matrices obtained with pertables).
- **env**: Data frame with the environmental variables.
- **dist.method**: Method to compute the dissimilarity matrices from the biological and environmental data tables. One of the methods described in function vegdist of the package vegan.
- **binary**: Value for the argument binary in the function vegdist of the package vegan.
- **cor.method**: Correlation method, as accepted by cor: "pearson", "spearman" or "kendall".
- **permutations**: Number of permutations in assessing significance.
- **x**: mantel_pertables object to plot.
- **xlab**: Label to name x-axis
- **ylab**: Label to name y-axis
- **pch**: Plotting 'character', i.e., symbol to use in the distance decay plot. See points for examples of use of this graphical argument.
- **...**: Additional graphical parameters passed to plot.

Value

mantel_pertables returns an object of class mantel_pertables, basically a list with the following components:

- **mantel**: A list with two components: mantel.raw, an object of class 'mantel', i.e. the results of applying mantel to the original biological data table without the unidentified species, and ptax, a p-value showing the probability of obtaining the same mantel statistic under different scenarios of taxonomic uncertainty.
- **simulation**: A list with the results of the simulation: results, i.e. a data.frame with all the simulated mantel statistics and p-values; mantel.quant, i.e. a data.frame with the summary of results by quantiles; vegdist, i.e. a list with all the dissimilarity matrices employed.

The objects of class mantel_pertables have print and plot S3 methods for a simple access to results. See the examples.

Author(s)

Luis Cayuela and Marcelino de la Cruz
References


See Also

`pertables`, `mantel`

Examples

```r
data(Amazonia)
data(soils)

# Define a new index that includes the terms used in the `Amazonia` dataset to define undetermined taxa at different taxonomic levels
index.Amazon <- c(paste("sp.", rep(1:20), sep=""), "Indet.", "indet."

## Not run:
# Generate a pertables object (i.e. a list of biological data tables simulated from taxonomic uncertainty)
Amazonia100 <- pertables(Amazonia, index=index.Amazon, nsim=100)

# Assess the effects of taxonomic uncertainty on a Mantel test of biological dissimilarity correlated to soil dissimilarity among sites:
Amazonia.mantel <- mantel_pertables(pertab=Amazonia100, env=soils, dist.method = "bray")
Amazonia.mantel
plot(Amazonia.mantel)

## End(Not run)
# Fast example for Rcheck
# Define a new index that includes the terms used in the `Amazonia` dataset to define undetermined taxa at different taxonomic levels
index.Amazon <- c(paste("sp.", rep(1:10), sep=""), "Indet.", "indet."

# Generate a pertables object (i.e. a list of biological data tables simulated from taxonomic uncertainty)
Amazonia100 <- pertables(Amazonia[1:50,], index=index.Amazon, nsim=4, ncl=2, iseed=4)
set.seed(2)
Amazonia.mantel <- mantel_pertables(pertab=Amazonia100, env=soils, dist.method = "bray")
Amazonia.mantel
plot(Amazonia.mantel)
```
Function to incorporate the effect of taxonomic uncertainty on multivariate analyses of ecological data.

Description

This function implements a permutational method to incorporate taxonomic uncertainty on multivariate analyses typically used in the analysis of ecological data. The procedure is based on iterative randomizations that randomly re-assign non identified species in each site to any of the other species found in the remaining sites.

Usage

pertables(data, index = NULL, nsim = 100)
pertables.p2(data, index = NULL, nsim = 100, ncl=2, iseed = NULL)

Arguments

data Community data matrix. The three first columns are factors referring to the family, genus and species specific names. The remaining columns are numeric vectors indicating species abundances at each site.

index List of additional parameters to determine the level at which species have been identified. Default values include ’Indet’, ’indet’, ’sp’, ’sp1’ to ’sp100’, ’sp 1’ to ’sp 100’, ”, and ’ ’.

nsim Number of simulations of species’ identities, i.e., number of data tables to simulate.

ncl Number of clusters for parallel simulation.

iseed An integer to be supplied to clusterSetRNGStream, or NULL not to set reproducible seeds.

Details

The procedure is implemented in two sequential steps:

Step 1. Morphospecies identified only to genus are randomly re-assigned with the same probability within the group of species and morphospecies that share the same genus, provided they are not found in the same sites. In the re-assignment of the species identity, the species considered can also receive its own identity. For instance, let’s assume we have three floristic inventories. In site A we have *Eugenia sp1* and *E. nesiotica*. In site B we have *Eugenia nesiotica*, *E. principium* and *E. salamensis*. In site C we have *Eugenia sp2* and *E. salamensis*. *Eugenia sp1* can be thus re-identified with equal probability as *Eugenia sp2*, *E. principium*, *E. salamensis* or just maintain its own identity (*Eugenia sp1*). In the latter case, this means that we assume that *E. sp1* is a completely different species, although we do not know its true identity. On the contrary, we cannot re-identify *E. sp1* as *E. nesiotica* because they were found in the same site, so we are quite certain that *E. sp1* is different from *E. nesiotica*. The same is applied to species identified only to family and fully unidentified species. Note that when collating inventories from different researchers, we must rename all unidentified species. This is because two researchers can use the same label, e.g. *Eugenia*
sp1, even though this name does not necessarily refer to the same species. For a verification of the biological identity of *Eugenia sp1* one would need to cross-check the vouchers bearing the same name.

Step 2. Step 1 is iterated *nsim* times. As a result, *nsim* matrices are obtained, all of which contain the same number of sites but variable number of species depending on the resulting re-assignment of morphospecies. The process can be time-consuming if community data matrices are large.

Function `pertables.p2` implements a parallelized version which considerably reduces computation time.

**Value**

The function return a list of class `pertables` with the following components

- `taxunc`: Summary of the number of species fully identified (0), identified to genus (1), identified to family (2), or fully undetermined (3).
- `pertables`: A list with all the simulated data matrices.
- `raw`: The raw data matrix, without the unidentified species.

**Author(s)**

Luis Cayuela and Marcelino de la Cruz

**References**


**Examples**

```r
data(Amazonia)
data(soils)

# Define a new index that includes the terms used in the Amazonia dataset to define undetermined taxa at different taxonomic levels
index.Amazon <- c(paste("sp.", rep(1:20), sep=""), "Indet.", "indet."

# Generate a pertables object (i.e. a list of biological data tables simulated from taxonomic uncertainty)

## Not run:
# compare performance of pertables and pertables.p2
nsim <- 100
ncl <- 2
gc()
t0 <- Sys.time()
Amazonia100 <- pertables(Amazonia, index=index.Amazon, nsim=nsim)
Sys.time() - t0
```
rda_pertables

Function to assess the effects of taxonomic uncertainty on [Partial] Redundance Analysis

Description

This function assesses the effects of taxonomic uncertainty on two widely used parameters of a [Partial] Redundance Analysis, i.e. the 'percentage explained variance' (sometimes referred to as $R^{2}$) and the 'pseudo-$F$'.

Usage

rda_pertables(fml, data, scale=FALSE, ...)

## S3 method for class 'rda_pertables'
plot(x, pch = 18, ...)

Arguments

fml         Model formula, where the left hand side gives a pertables object (i.e. a list of simulated community data matrices obtained with pertables, right hand side gives the constraining variables, and conditioning variables can be given within a special function Condition.
data        Data frame containing the variables on the right hand side of the model formula.
scale       Scale species to unit variance (like correlations).
x           rda_pertables object to plot.
pch         Plotting 'character', i.e., symbol to use in the RDA plot. See points for examples of use of this graphical argument.
...         Additional graphical parameters passed to plot.

Details

This function is a wrapper to submit a pertables object to rda function of the vegan package. See the documentation of cca for details about formula and Condition use.
**Value**

`rda_pertables` returns an object of class 'rda_pertables', basically a list with the following components:

- **raw**: An object of class 'rda'. The results of applying `rda` to the original biological data table without the unidentified species.
- **simulation**: A list with the results of the simulation: 'results', i.e. a data.frame with all the simulated R-squared and pseudo-F values; 'rda.quant', i.e. a data.frame with the summary of 'results' by quantiles; 'sites' i.e. a list with the scores of the sites of all the simulated data tables and 'biplot', i.e. a list with the scores of the environmental data in all the analyses.

The objects of class 'rda_pertables' have print and plot S3 methods for a simple access to results. See the examples.

**Author(s)**

Luis Cayuela and Marcelino de la Cruz

**References**


**See Also**

`pertables`, `cca`

**Examples**

```r
data(Amazonia)
data(soils)

# Define a new index that includes the terms used in the Amazonia dataset to define
# undetermined taxa at different taxonomic levels
index.Amazon <- c(paste("sp.", rep(1:20), sep=""), "Indet.", "indet.")

# Generate a pertables object (i.e. a list of biological data tables simulated from taxonomic
# uncertainty)
# Not run:
Amazonia100 <- pertables(Amazonia, index=index.Amazon, nsim=100)

# Assess the effects of taxonomic uncertainty on a RDA analysis of biological data explained
# by all the environmental variables of the soil data:
```
Amazonia.rda <- rda_pertables(Amazonia100 ~ ., data=soils)

Amazonia.rda

plot(Amazonia.rda)

## End(Not run)

# Fast example for Rcheck

Amazonia4.p2 <- pertables.p2(Amazonia[1:50,], index=index.Amazon, nsim=4, ncl=2, iseed=4)

set.seed(2)

Amazonia.rda <- rda_pertables(Amazonia4.p2 ~ ., data=soils)

Amazonia.rda

plot(Amazonia.rda)
Index

* **datasets**
  - Amazonia, 4
  - HCP, 7

* **multivariate**
  - adonis_pertables, 2
  - cca_pertables, 5
  - mantel_pertables, 7
  - pertables, 10
  - rda_pertables, 12

  adonis, 3
  adonis_pertables, 2
  Amazonia, 4

  cca, 5, 6, 12, 13
  cca_pertables, 5
  cor, 8

  HCP, 7

  mantel, 2, 8, 9
  mantel_pertables, 7

  pertables, 2, 5, 8, 10, 12
  plot.adonis_pertables
    (adonis_pertables), 2
  plot.cca_pertables (cca_pertables), 5
  plot.mantel_pertables
    (mantel_pertables), 7
  plot.rda_pertables (rda_pertables), 12
  points, 5, 8, 12
  print.adonis_pertables
    (adonis_pertables), 2
  print.cca_pertables (cca_pertables), 5
  print.mantel_pertables
    (mantel_pertables), 7
  print.pertables (pertables), 10
  print.rda_pertables (rda_pertables), 12

  rda_pertables, 12

  soils (Amazonia), 4

  vegdist, 8