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Suggests MASS, mgcv, lattice, cluster, scatterplot3d, rgl, tcltk

Description Ordination methods, diversity analysis and other functions for community and vegetation ecologists.

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### Description

The vegan package provides tools for descriptive community ecology. It has most basic functions of diversity analysis, community ordination and dissimilarity analysis. Most of its multivariate tools can be used for other data types as well.
Details

The functions in the vegan package contain tools for diversity analysis (see vignette vegandocs("diversity")), ordination and analysis of dissimilarities (see vignette vegandocs("intro")). Together with the labdsv package, the vegan package provides most standard tools of descriptive community analysis. Package ade4 provides an alternative comprehensive package, and several other packages complement vegan and provide tools for deeper analysis in specific fields. Package BiodiversityR provides a GUI for a large subset of vegan functionality.

The vegan package is developed at R-Forge (http://vegan.r-forge.r-project.org). The R-Forge provides up-to-date information and mailing lists for help queries and bug reports. Bug reports can also be emailed to the function authors or to the package maintainers.

The vegan documents can be read with vegandocs function. In addition to vignettes of basic usage, you can read NEWS on the new features and bug fixes in the release version (vegandocs("NEWS")), and more technical and fine grained ChangeLog (vegandocs("Change")). Several frequently asked questions really are answered in the vegan FAQ (vegandocs("FAQ")). The discussion on design decisions can be read with vegandocs("decision"). A tutorial of the package at http://cc.oulu.fi/~jarioksa/opetus/metodi/vegantutor.pdf provides a more thorough introduction to the package.

To see the preferable citation of the package, type citation("vegan").

Author(s)

The vegan development team is Jari Oksanen, F. Guillaume Blanchet, Roeland Kindt, Pierre Legendre, R. B. O’Hara, Gavin L. Simpson, Peter Solymos, M. Henry H. Stevens, Helene Wagner. Many other people have contributed to individual functions: see credits in function help pages.

The maintainers at the R-Forge are Jari Oksanen <jari.oksanen@oulu.fi> and Gavin Simpson <gavin.simpson@ucl.ac.uk>.

Examples

```r
### Example 1: Unconstrained ordination
## NMDS
data(varespec)
data(varechem)
ord <- metaMDS(varespec)
plot(ord, type = "t")
## Fit environmental variables
ef <- envfit(ord, varechem)
plot(ef, p.max = 0.05)
### Example 2: Constrained ordination (RDA)
## The example uses formula interface to define the model
data(dune)
data(dune.env)
## No constraints: PCA
mod0 <- rda(dune ~ 1, dune.env)
plot(mod0)
## All environmental variables: Full model
mod1 <- rda(dune ~ ., dune.env)
mod1
```
add1.cca

Add or Drop Single Terms to a Constrained Ordination Model

Description
Compute all single terms that can be added or dropped from a constrained ordination model.

Usage
## S3 method for class 'cca'
add1(object, scope, test = c("none", "permutation"),
     pstep = 100, perm.max = 200, ...)
## S3 method for class 'cca'
drop1(object, scope, test = c("none", "permutation"),
       pstep = 100, perm.max = 200, ...)

Arguments

object
A constrained ordination object from \texttt{cca}, \texttt{rda} or \texttt{capscale}.

scope
A formula giving the terms to be considered for adding or dropping; see \texttt{add1}
for details.

test
Should a permutation test added using \texttt{anova.cca}.
Number of permutations in one step, passed as argument `step` to `anova.cca`.

Maximum number of permutation in `anova.cca`.

Other arguments passed to `add1.default`, `drop1.default`, and `anova.cca`.

**Details**

With argument `test = "none"` the functions will only call `add1.default` or `drop1.default`. With argument `test = "permutation"` the functions will add test results from `anova.cca`. Function `drop1.cca` will call `anova.cca` with argument `by = "margin"`. Function `add1.cca` will implement a test for single term additions that is not directly available in `anova.cca`.

Functions are used implicitly in `step` and `ordistep`. The `deviance.cca` and `deviance.rda` used in `step` have no firm basis, and setting argument `test = "permutation"` may help in getting useful insight into validity of model building. Function `ordistep` calls alternately `drop1.cca` and `add1.cca` with argument `test = "permutation"` and selects variables by their permutation $P$-values. Meticulous use of `add1.cca` and `drop1.cca` will allow more judicious model building.

The default `perm.max` is set to a low value, because permutation tests can take a long time. It should be sufficient to give an impression on the significances of the terms, but higher values of `perm.max` should be used if $P$ values really are important.

**Value**

Returns a similar object as `add1` and `drop1`.

**Author(s)**

Jari Oksanen

**See Also**

`add1`, `drop1` and `anova.cca` for basic methods. You probably need these functions with `step` and `link{ordistep}`. Functions `deviance.cca` and `extractAIC.cca` are used to produce the other arguments than test results in the output. Functions `cca`, `rda` and `capscale` produce result objects for these functions.

**Examples**

```r
data(dune)
data(dune.env)
## Automatic model building based on AIC but with permutation tests
step(cca(dune ~ 1, dune.env), reformulate(names(dune.env)), test="perm")
## The same, but based on permutation $P$-values
ordistep(cca(dune ~ 1, dune.env), reformulate(names(dune.env)), perm.max=200)
## Manual model building
## -- define the maximal model for scope
mbig <- rda(dune ~ ., dune.env)
## -- define an empty model to start with
m0 <- rda(dune ~ 1, dune.env)
## -- manual selection and updating
```
adipart

Additive Diversity Partitioning and Hierarchical Null Model Testing

Description

In additive diversity partitioning, mean values of alpha diversity at lower levels of a sampling hierarchy are compared to the total diversity in the entire data set (gamma diversity). In hierarchical null model testing, a statistic returned by a function is evaluated according to a nested hierarchical sampling design (hiersimu).

Usage

adipart(formula, data, index=c("richness", "shannon", "simpson"), weights=c("unif", "prop"), relative = FALSE, nsimul=99, ...)
hiersimu(formula, data, FUN, location = c("mean", "median"), relative = FALSE, drop.highest = FALSE, nsimul=99, ...)
## S3 method for class 'adipart'
print(x, ...)
## S3 method for class 'hiersimu'
print(x, ...)

Arguments

formula A two sided model formula in the form y ~ x, where y is the community data matrix with samples as rows and species as column. Right hand side (x) must contain factors referring to levels of sampling hierarchy, terms from right to left will be treated as nested (first column is the lowest, last is the highest level). These variables must be factors in order to unambiguous handling. Interaction terms are not allowed.

data A data frame where to look for variables defined in the right hand side of formula. If missing, variables are looked in the global environment.

index Character, the diversity index to be calculated (see Details).

weights Character, "unif" for uniform weights, "prop" for weighting proportional to sample abundances to use in weighted averaging of individual alpha values within strata of a given level of the sampling hierarchy.

relative Logical, if TRUE then alpha and beta diversity values are given relative to the value of gamma for function adipart.
nsimul  Number of permutation to use if `matr` is not of class 'permat'. If `nsimul = 0`, only the `FUN` argument is evaluated. It is thus possible to reuse the statistic values without using a null model.

FUN  A function to be used by `hiersimu`. This must be fully specified, because currently other arguments cannot be passed to this function via . . .

location  Character, identifies which function (mean or median) is to be used to calculate location of the samples.

drop.highest  Logical, to drop the highest level or not. When `FUN` evaluates only arrays with at least 2 dimensions, highest level should be dropped, or not selected at all.

x  An object to print.

. . .  Other arguments passed to functions, e.g. base of logarithm for Shannon diversity, or `method, thin` or `burnin` arguments for `oecosimu`.

Details

Additive diversity partitioning means that mean alpha and beta diversity adds up to gamma diversity, thus beta diversity is measured in the same dimensions as alpha and gamma (Lande 1996). This additive procedure is than extended across multiple scales in a hierarchical sampling design with \( i = 1, 2, 3, \ldots, m \) levels of sampling (Crist et al. 2003). Samples in lower hierarchical levels are nested within higher level units, thus from \( i = 1 \) to \( i = m \) grain size is increasing under constant survey extent. At each level \( i \), \( \alpha_i \) denotes average diversity found within samples.

At the highest sampling level, the diversity components are calculated as

\[
\beta_m = \gamma - \alpha_m
\]

For each lower sampling level as

\[
\beta_i = \alpha_{i+1} - \alpha_i
\]

Then, the additive partition of diversity is

\[
\gamma = \alpha_1 + \sum_{i=1}^{m} \beta_i
\]

Average alpha components can be weighted uniformly (weight = "unif") to calculate it as simple average, or proportionally to sample abundances (weight = "prop") to calculate it as weighted average as follows

\[
\alpha_i = \sum_{j=1}^{n_i} D_{ij} w_{ij}
\]

where \( D_{ij} \) is the diversity index and \( w_{ij} \) is the weight calculated for the \( j \)th sample at the \( i \)th sampling level.

The implementation of additive diversity partitioning in `adipart` follows Crist et al. 2003. It is based on species richness (`S`, not `S - 1`), Shannon’s and Simpson’s diversity indices stated as the index argument.

The expected diversity components are calculated `nsimul` times by individual based randomisation of the community data matrix. This is done by the "r2dtable" method in `oecosimu` by default.
hiersimu works almost the same as adipart, but without comparing the actual statistic values returned by FUN to the highest possible value (cf. gamma diversity). This is so, because in most of the cases, it is difficult to ensure additive properties of the mean statistic values along the hierarchy.

Value

An object of class ‘adipart’ or ‘hiersimu’ with same structure as ‘oecosimu’ objects.

Author(s)

Péter Sólymos, <solymos@ualberta.ca>

References


See Also

See oecosimu for permutation settings and calculating $p$-values.

Examples

data(mite)
data(mite.xy)
data(mite.env)
## Function to get equal area partitions of the mite data
cutter <- function (x, cut = seq(0, 10, by = 2.5)) {
  out <- rep(1, length(x))
  for (i in 2:(length(cut) - 1))
    out[which(x > cut[i] & x <= cut[(i + 1)])] <- i
  return(as.factor(out))
}
## The hierarchy of sample aggregation
levsm <- data.frame(
  l1=as.factor(1:nrow(mite)),
  l2=cutter(mite.xy$y, cut = seq(0, 10, by = 2.5)),
  l3=cutter(mite.xy$y, cut = seq(0, 10, by = 5)),
  l4=cutter(mite.xy$y, cut = seq(0, 10, by = 10)))
## Let's see in a map
par(mfrow=c(1,3))
plot(mite.xy, main="l1", col=as.numeric(levsm$l1)+1)
plot(mite.xy, main="l2", col=as.numeric(levsm$l2)+1)
plot(mite.xy, main="l3", col=as.numeric(levsm$l3)+1)
par(mfrow=c(1,1))
## Additive diversity partitioning
adipart(mite ~., levsm, index="richness", nsimul=20)
## Hierarchical null model testing
## diversity analysis (similar to adipart)
hiersimu(mite ~., levsm, diversity, relative=TRUE, nsimul=25)
## Hierarchical testing with the Morisita index
morfun <- function(x) dispindmorisita(x)$imst
hiersimu(mite ~., levsm, morfun, drop.highest=TRUE, nsimul=25)

---

### Description

Analysis of variance using distance matrices — for partitioning distance matrices among sources of variation and fitting linear models (e.g., factors, polynomial regression) to distance matrices; uses a permutation test with pseudo-F ratios.

### Usage

```r
adonis(formula, data, permutations = 999, method = "bray",
       strata = NULL, contr.unordered = "contr.sum",
       contr.ordered = "contr.poly", ...)
```

### Arguments

- **formula**: a typical model formula such as `Y ~ A + B*C`, but where `Y` is either a dissimilarity object (inheritng from class "dist") or data frame or a matrix; `A`, `B`, and `C` may be factors or continuous variables. If a dissimilarity object is supplied, no species coefficients can be calculated (see Value below).
- **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).
- **method**: the name of any method used in `vegdist` to calculate pairwise distances if the left hand side of the `formula` was a data frame or a matrix.
- **strata**: groups (strata) within which to constrain permutations.
- **contr.unordered**, **contr.ordered**: contrasts used for the design matrix (default in R is dummy or treatment contrasts for unordered factors).
- **...**: Other arguments passed to `vegdist`.

### Details

`adonis` is a function for the analysis and partitioning sums of squares using semimetric and metric distance matrices. Insofar as it partitions sums of squares of a multivariate data set, it is directly analogous to MANOVA (multivariate analysis of variance). M.J. Anderson (McArdle and Anderson 2001, Anderson 2001) refers to the method as “permutational manova” (formerly “nonparametric manova”). Further, as its inputs are linear predictors, and a response matrix of an arbitrary number of columns (2 to millions), it is a robust alternative to both parametric MANOVA and to ordination...
methods for describing how variation is attributed to different experimental treatments or uncontrolled covariates. It is also analogous to redundancy analysis (Legendre and Anderson 1999).

Typical uses of adonis include analysis of ecological community data (samples X species matrices) or genetic data where we might have a limited number of samples of individuals and thousands or millions of columns of gene expression data (e.g. Zapala and Schork 2006).

adonis is an alternative to AMOVA (nested analysis of molecular variance, Excoffier, Smouse, and Quattro, 1992; amova in the ade4 package) for both crossed and nested factors.

If the experimental design has nestedness, then use strata to test hypotheses. For instance, imagine we are testing the whether a plant community is influenced by nitrate amendments, and we have two replicate plots at each of two levels of nitrate (0, 10 ppm). We have replicated the experiment in three fields with (perhaps) different average productivity. In this design, we would need to specify strata = field so that randomizations occur only within each field and not across all fields. See example below.

Like AMOVA (Excoffier et al. 1992), adonis relies on a long-understood phenomenon that allows one to partition sums of squared deviations from a centroid in two different ways (McArdle and Anderson 2001). The most widely recognized method, used, e.g., for ANOVA and MANOVA, is to first identify the relevant centroids and then to calculated the squared deviations from these points.

For a centered $n \times p$ response matrix $Y$, this method uses the $p \times p$ inner product matrix $Y'Y$. The less appreciated method is to use the $n \times n$ outer product matrix $YY'$. Both AMOVA and adonis use this latter method. This allows the use of any semimetric (e.g. Bray-Curtis, aka Steinhaus, Czekanowski, and Sørensen) or metric (e.g. Euclidean) distance matrix (McArdle and Anderson 2001). Using Euclidean distances with the second method results in the same analysis as the first method.

Significance tests are done using $F$-tests based on sequential sums of squares from permutations of the raw data, and not permutations of residuals. Permutations of the raw data may have better small sample characteristics. Further, the precise meaning of hypothesis tests will depend upon precisely what is permuted. The strata argument keeps groups intact for a particular hypothesis test where one does not want to permute the data among particular groups. For instance, strata = B causes permutations among levels of A but retains data within levels of B (no permutation among levels of B). See permutations for additional details on permutation tests in Vegan.

The default contrasts are different than in R in general. Specifically, they use “sum” contrasts, sometimes known as ‘ANOVA’ contrasts. See a useful text (e.g. Crawley, 2002) for a transparent introduction to linear model contrasts. This choice of contrasts is simply a personal pedagogical preference. The particular contrasts can be set to any contrasts specified in R, including Helmert and treatment contrasts.

Rules associated with formulae apply. See "An Introduction to R" for an overview of rules.

print.adonis shows the aov.tab component of the output.

Value

This function returns typical, but limited, output for analysis of variance (general linear models).

aov.tab Typical AOV table showing sources of variation, degrees of freedom, sequential sums of squares, mean squares, $F$ statistics, partial R-squared and $P$ values, based on $N$ permutations.
coefficients matrix of coefficients of the linear model, with rows representing sources of variation and columns representing species; each column represents a fit of a species abundance to the linear model. These are what you get when you fit one species to your predictors. These are NOT available if you supply the distance matrix in the formula, rather than the site x species matrix.

coeff.sites matrix of coefficients of the linear model, with rows representing sources of variation and columns representing sites; each column represents a fit of sites distances (from all other sites) to the linear model. These are what you get when you fit distances of one site to your predictors.

f.perms an N by m matrix of the null F statistics for each source of variation based on N permutations of the data.

model.matrix The model.matrix for the right hand side of the formula.

terms The terms component of the model.

Author(s)

Martin Henry H. Stevens <HStevens@muohio.edu>, adapted to vegan by Jari Oksanen.

References


See Also

mrpp, anosim, mantel, varpart.

Examples

data(dune)
data(dune.env)
adonis(dune ~ Management*A1, data=dune.env, permutations=99)

### Example of use with strata, for nested (e.g., block) designs.
dat <- expand.grid(rep=gl(2,1), NO3=factor(c(0,10)),field=gl(3,1) )
dat
Agropyron <- with(dat, as.numeric(field) + as.numeric(NO3)+2) +rnorm(12)/2
Schizachyrium <- with(dat, as.numeric(field) - as.numeric(NO3)+2) +rnorm(12)/2
total <- Agropyron + Schizachyrium
library(lattice)
dotplot(total ~ NO3, dat, jitter.x=TRUE, groups=field, 
  type=c('p','a'), xlab="NO3", auto.key=list(columns=3, lines=TRUE) )

Y <- data.frame(Agropyron, Schizachyrium)
mod <- metaMDS(Y)
plot(mod)
### Hulls show treatment
ordihull(mod, group=dat$NO3, show="0")
ordihull(mod, group=dat$NO3, show="10", col=3)
### Spider shows fields
ordispider(mod, group=dat$field, lty=3, col="red")

### Correct hypothesis test (with strata)
adonis(Y ~ NO3, data=dat, strata=dat$field, perm=1e3)

### Incorrect (no strata)
adonis(Y ~ NO3, data=dat, perm=1e3)

---

### anosim

**Analysis of Similarities**

**Description**

Analysis of similarities (ANOSIM) provides a way to test statistically whether there is a significant difference between two or more groups of sampling units.

**Usage**

anosim(dat, grouping, permutations = 999, distance = "bray", strata)

**Arguments**

- **dat**
  - Data matrix or data frame in which rows are samples and columns are response variable(s), or a dissimilarity object or a symmetric square matrix of dissimilarities.

- **grouping**
  - Factor for grouping observations.

- **permutations**
  - Number of permutation to assess the significance of the ANOSIM statistic.

- **distance**
  - Choice of distance metric that measures the dissimilarity between two observations. See vegdist for options. This will be used if dat was not a dissimilarity structure or a symmetric square matrix.

- **strata**
  - An integer vector or factor specifying the strata for permutation. If supplied, observations are permuted only within the specified strata.
Details

Analysis of similarities (ANOSIM) provides a way to test statistically whether there is a significant
difference between two or more groups of sampling units. Function `anosim` operates directly on a
dissimilarity matrix. A suitable dissimilarity matrix is produced by functions `dist` or `vegdist`.
The method is philosophically allied with NMDS ordination (isoMDS), in that it uses only the rank
order of dissimilarity values.

If two groups of sampling units are really different in their species composition, then compositional
dissimilarities between the groups ought to be greater than those within the groups. The `anosim`
statistic $R$ is based on the difference of mean ranks between groups ($r_B$) and within groups ($r_W$):

$$ R = \frac{(r_B - r_W)}{(N(N - 1)/4)} $$

The divisor is chosen so that $R$ will be in the interval $-1 \ldots + 1$, value 0 indicating completely
random grouping.

The statistical significance of observed $R$ is assessed by permuting the grouping vector to obtain
the empirical distribution of $R$ under null-model. See `permutations` for additional details on
permutation tests in Vegan.

The function has `summary` and `plot` methods. These both show valuable information to assess
the validity of the method: The function assumes that all ranked dissimilarities within groups have
about equal median and range. The `plot` method uses `boxplot` with options `notch=TRUE` and
`varwidth=TRUE`.

Value

The function returns a list of class "anosim" with following items:

- `call` Function call.
- `statistic` The value of ANOSIM statistic $R$
- `signif` Significance from permutation.
- `perm` Permutation values of $R$
- `class.vec` Factor with value `Between` for dissimilarities between classes and class name
  for corresponding dissimilarity within class.
- `dis.rank` Rank of dissimilarity entry.
- `dissimilarity` The name of the dissimilarity index: the "method" entry of the `dist` object.

Note

I don’t quite trust this method. Somebody should study its performance carefully. The function
returns a lot of information to ease further scrutiny. Most `anosim` models could be analysed with
`adonis` which seems to be a more robust alternative.

Author(s)

Jari Oksanen, with a help from Peter R. Minchin.
References


See Also

`mrpp` for a similar function using original dissimilarities instead of their ranks. `dist` and `vegdist` for obtaining dissimilarities, and `rank` for ranking real values. For comparing dissimilarities against continuous variables, see `mantel`. Function `adonis` is a more robust alternative that should preferred.

Examples

data(dune)
data(dune.env)
dune.dist <- vegdist(dune)
attach(dune.env)
dune.ano <- anosim(dune.dist, Management)
summary(dune.ano)
plot(dune.ano)

---

**anova.cca**

*Permutation Test for Constrained Correspondence Analysis, Redundancy Analysis and Constrained Analysis of Principal Coordinates*

Description

The function performs an ANOVA like permutation test for Constrained Correspondence Analysis (`cca`), Redundancy Analysis (`rda`) or Constrained Analysis of Principal Coordinates (`capscale`) to assess the significance of constraints.

Usage

```r
## S3 method for class 'cca'
anova(object, alpha=0.05, beta=0.01, step=100, perm.max=9999,
          by = NULL, ...)
```

```r
permutest(x, ...)
```

```r
## S3 method for class 'cca'
permutest(x, permutations = 99,
          model = c("reduced", "direct", "full"),
          first = FALSE, strata, ...)
```
Arguments

object, x A result object from cca.
alpha Targeted Type I error rate.
beta Accepted Type II error rate.
step Number of permutations during one step.
perm.max Maximum number of permutations.
by Setting by = "axis" will assess significance for each constrained axis, and setting by = "terms" will assess significance for each term (sequentially from first to last), and setting by = "margin" will assess the marginal effects of the terms (each marginal term analysed in a model with all other variables).

... Parameters passed to other functions. anova.cca passes all arguments to permutest.cca. In anova with by = "axis" you can use argument cutoff (defaults 1) which stops permutations after exceeding the given level.

permutations Number of permutations for assessing significance of constraints.
model Permutation model (partial match).
first Assess only the significance of the first constrained eigenvalue; will be passed from anova.cca.
strata An integer vector or factor specifying the strata for permutation. If supplied, observations are permuted only within the specified strata.

Details

Functions anova.cca and permutest.cca implement an ANOVA like permutation test for the joint effect of constraints in cca, rda or capscale. Functions anova.cca and permutest.cca differ in printout style and in interface. Function permutest.cca is the proper workhorse, but anova.cca passes all parameters to permutest.cca.

In anova.cca the number of permutations is controlled by targeted “critical” P value (alpha) and accepted Type II or rejection error (beta). If the results of permutations differ from the targeted alpha at risk level given by beta, the permutations are terminated. If the current estimate of P does not differ significantly from alpha of the alternative hypothesis, the permutations are continued with step new permutations (at the first step, the number of permutations is step - 1). However, with by = "terms" a fixed number of permutations will be used, and this is given by argument permutations, or if this is missing, by step.

The function permutest.cca implements a permutation test for the “significance” of constraints in cca, rda or capscale. Community data are permuted with choice model = "direct", residuals after partial CCA/RDA/CAP with choice model = "reduced" (default), and residuals after CCA/RDA/CAP under choice model = "full". If there is no partial CCA/RDA/CAP stage, model = "reduced" simply permutes the data and is equivalent to model = "direct".

The test statistic is “pseudo-F”, which is the ratio of constrained and unconstrained total Inertia (Chi-squares, variances or something similar), each divided by their respective ranks. If there are no conditions (“partial” terms), the sum of all eigenvalues remains constant, so that pseudo-F and eigenvalues would give equal results. In partial CCA/RDA/CAP, the effect of conditioning variables (“covariables” is removed before permutation, and these residuals are added to the non-permuted fitted values of partial CCA (fitted values of X ~ Z). Consequently, the total Chi-square is not
fixed, and test based on pseudo-$F$ would differ from the test based on plain eigenvalues. CCA is a weighted method, and environmental data are re-weighted at each permutation step using permuted weights.

The default test is for the sum of all constrained eigenvalues. Setting `first = TRUE` will perform a test for the first constrained eigenvalue. Argument `first` can be set either in `anova.cca` or in `permutest.cca`. It is also possible to perform significance tests for each axis or for each term (constraining variable) using argument `by` in `anova.cca`. Setting `by = "axis"` will perform separate significance tests for each constrained axis. All previous constrained axes will be used as conditions (“partialled out”) and a test for the first constrained eigenvalues is performed. You can stop permutation tests after exceeding a given significance level with argument `cutoff` to speed up calculations in large models. Setting `by = "terms"` will perform separate significance test for each term (constraining variable). Setting `by = "margin"` will perform separate significance test for each marginal term in a model with all other terms. The marginal test also accepts a `scope` argument for the drop scope which can be a character vector of term labels that are analysed, or a fitted model of lower scope. The marginal effects are also known as “Type III” effects, but the current function only evaluates marginal terms. It will, for instance, ignore main effects that are included in interaction terms. In calculating pseudo-$F$, all terms are compared to the same residual of the full model. Permutations for all axes or terms will start from the same .Random.seed, and the seed will be advanced to the value after the longest permutation at the exit from the function.

Value

Function `permutest.cca` returns an object of class "permutest.cca", which has its own print method. The function `anova.cca` calls `permutest.cca`, fills an `anova` table and uses `print.anova` for printing.

Note

Some cases of `anova` need access to the original data on constraints (at least `by = "term"` and `by = "margin"`), and they may fail if data are unavailable.

The default permutation model changed from "direct" to "reduced" in `vegan` version 1.14-11 (release version 1.15-0), and you must explicitly set `model = "direct"` for compatibility with the old version.

Tests `by = "terms"` and `by = "margin"` are consistent only when `model = "direct"`.

Author(s)

Jari Oksanen

References

See Also

`cca`, `rda`, `capscale` to get something to analyse. Function `drop1.cca` calls `anova.cca` with `by = "margin"`, and `add1.cca` an analysis for single terms additions, which can be used in automatic or semiautomatic model building (see `deviance.cca`).

Examples

data(varespec)
data(varechem)
vare.cca <- cca(varespec ~ Al + P + K, varechem)
## overall test
anova(vare.cca)
## Test for axes
anova(vare.cca, by="axis", perm.max=500)
## Sequential test for terms
anova(vare.cca, by="terms", permu=200)
## Marginal or Type III effects
anova(vare.cca, by="margin")
## Marginal test knows 'scope'
anova(vare.cca, by = "m", scope="P")

as.mlm.cca

Refit Constrained Ordination as a Multiple Response Linear Model

Description

Functions refit results of constrained ordination (`cca`, `rda`, `capscale`) as a multiple response linear model (`lm`). This allows finding influence statistics (`influence.measures`). This also allows deriving several other statistics, but most of these are biased and misleading, since refitting ignores a major component of variation in constrained ordination.

Usage

`as.mlm(x)`

Arguments

- `x` Constrained ordination result.

Details

Popular algorithm for constrained ordination is based on iteration with regression where weighted averages of sites are used as dependent variables and constraints as independent variables. Statistics of linear regression are a natural by-product in this algorithm. Constrained ordination in `vegan` uses different algorithm, but to obtain linear regression statistics you can refit an ordination result as a multiple response linear model (`lm`). This regression ignores residual unconstrained variation in the data, and therefore estimates of standard error are strongly biased and much too low. You can get statistics like `t`-values of coefficients, but you should not use these because of this bias.
Some useful information you can get with refitted models are statistics for detecting influential observations (influence.measures including cooks.distance, hatvalues).

Value

Function returns an object of multiple response linear model of class "mlm" documented with lm.

Note

You can use these functions to find t-values of coefficients using summary.mlm, but you should not do this because the method ignores unconstrained residual variation. You also can find several other statistics for (multiple response) linear models with similar bias. This bias is not a unique feature in vegan implementation, but also applies to implementations in other software.

Some statistics of linear models can be found without using these functions: coef.cca gives the regression coefficients, spenvcor the species-environment correlation, intersetcor the interset correlation, vif.cca the variance inflation factors.

Author(s)

Jari Oksanen

See Also

cca, rda, capscale, cca.object, lm, summary.mlm, influence.measures.

Examples

data(varespec)
data(varechem)
mod <- cca(varespec ~ Al + P + K, data=varechem)
lmod <- as.mlm(mod)
## Coefficients
lmod
coef(mod)
## Influential observations
influence.measures(lmod)
plot(mod, type = "n")
points(mod, cex = 10*hatvalues(lmod), pch=16, xpd = TRUE)
text(mod, display = "bp", col = "blue")

BCI

Barro Colorado Island Tree Counts

Description

Tree counts in 1-hectare plots in the Barro Colorado Island.
Usage

```r
data(BCI)
```

Format

A data frame with 50 plots (rows) of 1 hectare with counts of trees on each plot with total of 225 species (columns). Full Latin names are used for tree species.

Details

Data give the numbers of trees at least 10 cm in diameter at breast height (1.3 m above the ground) in each one hectare square of forest. Within each one hectare square, all individuals of all species were tallied and are recorded in this table.

The data frame contains only the Barro Colorado Island subset of the original data. The quadrats are located in a regular grid. See `examples` for the coordinates.

Source

[http://www.sciencemag.org/cgi/content/full/295/5555/666/DC1](http://www.sciencemag.org/cgi/content/full/295/5555/666/DC1)

References


See Also

`BCI.env` in `BiodiversityR` package for environmental data (coordinates are given below in the examples).

Examples

```r
data(BCI)
## UTM Coordinates (in metres)
UTM.EW <- rep(seq(625754, 626654, by=100), each=5)  
UTM.NS <- rep(seq(1011569, 1011969, by=100), len=50)
```

Beals Smoothing and Degree of Absence

Description

Beals smoothing replaces each entry in the community data with a probability of a target species occurring in that particular site, based on the joint occurrences of the target species with the species that actually occur in the site. Swan’s (1970) degree of absence applies Beals smoothing to zero items so long that all zeros are replaced with smoothed values.
Usage

```r
beals(x, species = NA, reference = x, type = 0, include = TRUE)
swan(x)
```

Arguments

- `x`: Community data frame or matrix.
- `species`: Column index used to compute Beals function for a single species. The default (NA) indicates that the function will be computed for all species.
- `reference`: Community data frame or matrix to be used to compute joint occurrences. By default, `x` is used as reference to compute the joint occurrences.
- `type`: Numeric. Specifies if and how abundance values have to be used in function `beals`. See details for more explanation.
- `include`: This logical flag indicates whether the target species has to be included when computing the mean of the conditioned probabilities. The original Beals (1984) definition is equivalent to `include=TRUE`, while the formulation of Münzbergová and Herben is equal to `include=FALSE`.

Details

Beals smoothing is the estimated probability \( p_{ij} \) that species \( j \) occurs at site \( i \). It is defined as \( p_{ij} = \frac{1}{S_i} \sum_k \frac{N_{jk} I_{ik}}{N_k} \), where \( S_i \) is the number of species at site \( i \), \( N_{jk} \) is the number of joint occurrences of species \( j \) and \( k \), \( N_k \) is the number of occurrences of species \( k \), and \( I \) is the incidence (0 or 1) of species (this last term is usually omitted from the equation, but it is necessary). As \( N_{jk} \) can be interpreted as a mean of conditional probability, the `beals` function can be interpreted as a mean of conditioned probabilities (De Cáceres & Legendre 2008). The present function is generalized to abundance values (De Cáceres & Legendre 2008).

The `type` argument specifies if and how abundance values have to be used. `type = 0` presence/absence mode. `type = 1` abundances in `reference` (or `x`) are used to compute conditioned probabilities. `type = 2` abundances in `x` are used to compute weighted averages of conditioned probabilities. `type = 3` abundances are used to compute both conditioned probabilities and weighted averages.

Beals smoothing was originally suggested as a method of data transformation to remove excessive zeros (Beals 1984, McCune 1994). However, it is not a suitable method for this purpose since it does not maintain the information on species presences: a species may have a higher probability of occurrence at a site where it does not occur than at sites where it occurs. Moreover, it regularizes data too strongly. The method may be useful in identifying species that belong to the species pool (Ewald 2002) or to identify suitable unoccupied patches in metapopulation analysis (Münzbergová & Herben 2004). In this case, the function should be called with `include = FALSE` for cross-validation smoothing for species; argument `species` can be used if only one species is studied.

Swan (1970) suggested replacing zero values with degrees of absence of a species in a community data matrix. Swan expressed the method in terms of a similarity matrix, but it is equivalent to applying Beals smoothing to zero values, at each step shifting the smallest initially non-zero item to value one, and repeating this so many times that there are no zeros left in the data. This is actually very similar to extended dissimilarities (implemented in function `stepacross`), but very rarely used.
Value

The function returns a transformed data matrix or a vector if Beals smoothing is requested for a single species.

Author(s)

Miquel De Cáceres and Jari Oksanen

References


See Also

decostand for proper standardization methods, specpool for an attempt to assess the size of species pool. Function indpower assesses the power of each species to estimate the probabilities predicted by beals.

Examples

data(dune)
## Default
x <- beals(dune)
## Remove target species
x <- beals(dune, include = FALSE)
## Smoothed values against presence or absence of species
pa <- decostand(dune, "pa")
boxplot(as.vector(x) ~ unlist(pa), xlab="Presence", ylab="Beals")
## Remove the bias of tarbet species: Yields lower values.
beals(dune, type =3, include = FALSE)
## Uses abundance information.
## Vector with beals smoothing values corresponding to the first species in dune.
beals(dune, species=1, include=TRUE)
betadisper

Multivariate homogeneity of groups dispersions (variances)

Description

Implements Marti Anderson’s PERMDISP2 procedure for the analysis of multivariate homogeneity of group dispersions (variances). betadisper is a multivariate analogue of Levene’s test for homogeneity of variances. Non-euclidean distances between objects and group centroids are handled by reducing the original distances to principal coordinates. This procedure has latterly been used as a means of assessing beta diversity. There are anova, scores, plot and boxplot methods. TukeyHSD.betadisper creates a set of confidence intervals on the differences between the mean distance-to-centroid of the levels of the grouping factor with the specified family-wise probability of coverage. The intervals are based on the Studentized range statistic, Tukey’s ‘Honest Significant Difference’ method.

Usage

betadisper(d, group, type = c("median","centroid"))

## S3 method for class 'betadisper'
anova(object, ...)

## S3 method for class 'betadisper'
scores(x, display = c("sites","centroids"),
      choices = c(1,2), ...)

## S3 method for class 'betadisper'
plot(x, axes = c(1,2), cex = 0.7, hull = TRUE,
     ylab, xlab, main, sub, ...)

## S3 method for class 'betadisper'
boxplot(x, ylab = "Distance to centroid", ...)

## S3 method for class 'betadisper'
TukeyHSD(x, which = "group", ordered = FALSE,
         conf.level = 0.95, ...)

Arguments

d  a distance structure such as that returned by dist, betadiver or vegdist.
group  vector describing the group structure, usually a factor or an object that can be coerced to a factor using as.factor. Can consist of a factor with a single level (i.e., one group).
type  the type of analysis to perform. Use the spatial median or the group centroid? The spatial median is now the default.
display  character; partial match to access scores for "sites" or "species".
details

One measure of multivariate dispersion (variance) for a group of samples is to calculate the average
distance of group members to the group centroid or spatial median (both referred to as 'centroid'
from now on unless stated otherwise) in multivariate space. To test if the dispersions (variances) of
one or more groups are different, the distances of group members to the group centroid are subject
to ANOVA. This is a multivariate analogue of Levene's test for homogeneity of variances if the
distances between group members and group centroids is the Euclidean distance.

However, better measures of distance than the Euclidean distance are available for ecological data.
These can be accommodated by reducing the distances produced using any dissimilarity coefficient
to principal coordinates, which embeds them within a Euclidean space. The analysis then proceeds
by calculating the Euclidean distances between group members and the group centroid on the basis
of the principal coordinate axes rather than the original distances.

Non-metric dissimilarity coefficients can produce principal coordinate axes that have negative Eigen-
values. These correspond to the imaginary, non-metric part of the distance between objects. If
negative Eigenvalues are produced, we must correct for these imaginary distances.

The distance to its centroid of a point is

$$z_{ij}^2 = \sqrt{\Delta^2(u_{ij}^+, c_i^+) - \Delta^2(u_{ij}^-, c_i^-)},$$

where $\Delta^2$ is the squared Euclidean distance between $u_{ij}$, the principal coordinate for the $j^{th}$
point in the $i^{th}$ group, and $c_i$, the coordinate of the centroid for the $i^{th}$ group. The super-scripted + and
− indicate the real and imaginary parts respectively. This is equation (3) in Anderson (2006). If the
imaginary part is greater in magnitude than the real part, then we would be taking the square root
of a negative value, resulting in NaN. From vegan 1.12-12 betadisper takes the absolute value
of the real distance minus the imaginary distance, before computing the square root. This is in line
with the behaviour of Marti Anderson's PERMDISP2 programme.

To test if one or more groups is more variable than the others, ANOVA of the distances to group
centroids can be performed and parametric theory used to interpret the significance of F. An al-
ternative is to use a permutation test. permute.betadisper permutes model residuals to
generate a permutation distribution of F under the Null hypothesis of no difference in dispersion
between groups.
Pairwise comparisons of group mean dispersions can also be performed using `permutest.betadisper`. An alternative to the classical comparison of group dispersions, is to calculate Tukey's Honest Significant Differences between groups, via `TukeyHSD.betadisper`. This is a simple wrapper to `TukeyHSD.aov`. The user is directed to read the help file for `TukeyHSD` before using this function. In particular, note the statement about using the function with unbalanced designs.

The results of the analysis can be visualised using the `plot` and `boxplot` methods.

One additional use of these functions is in assessing beta diversity (Anderson et al 2006). Function `betadiver` provides some popular dissimilarity measures for this purpose.

**Value**

The `anova` method returns an object of class "anova" inheriting from class "data.frame".

The `scores` method returns a list with one or both of the components "sites" and "centroids".

The `plot` function invisibly returns an object of class "ordiplot", a plotting structure which can be used by `identify.ordiplot` (to identify the points) or other functions in the `ordiplot` family.

The `boxplot` function invisibly returns a list whose components are documented in `boxplot`.

`TukeyHSD.betadisper` returns a list. See `TukeyHSD` for further details.

`betadisper` returns a list of class "betadisper" with the following components:

- **eig** numeric; the eigenvalues of the principal coordinates analysis.
- **vectors** matrix; the eigenvectors of the principal coordinates analysis.
- **distances** numeric; the Euclidean distances in principal coordinate space between the samples and their respective group centroid.
- **group** factor; vector describing the group structure
- **centroids** matrix; the locations of the group centroids on the principal coordinates.
- **call** the matched function call.

**Warning**

Stewart Schultz noticed that the permutation test for `type = "centroid"` had the wrong type I error and was anti-conservative. As such, the default for `type` has been changed to "median", which uses the spatial median as the group centroid. Tests suggests that the permutation test for this type of analysis gives the correct error rates.

**Note**

If `group` consists of a single level or group, then the `anova` and `permutest` methods are not appropriate and if used on such data will stop with an error.

Missing values in either `d` or `group` will be removed prior to performing the analysis.

**Author(s)**

Gavin L. Simpson
References


See Also

`permutest.betadisper`, `anova.lm`, `scores`, `boxplot`, `TukeyHSD`. Further measure of beta diversity can be found in `betadiver`.

Examples

data(varespec)

## Bray-Curtis distances between samples
dis <- vegdist(varespec)

## First 16 sites grazed, remaining 8 sites ungrazed
groups <- factor(c(rep(1,16), rep(2,8)), labels = c("grazed","ungrazed"))

## Calculate multivariate dispersions
mod <- betadisper(dis, groups)

## Perform test
anova(mod)

## Permutation test for F
permutest(mod, pairwise = TRUE)

## Tukey’s Honest Significant Differences
(mod.HSD <- TukeyHSD(mod))
plot(mod.HSD)

## Plot the groups and distances to centroids on the
## first two PCoA axes
plot(mod)

## can also specify which axes to plot, ordering respected
plot(mod, axes = c(3,1))

## Draw a boxplot of the distances to centroid for each group
boxplot(mod)

## simulate missing values in 'd' and 'group'
groups[c(2,20)] <- NA
dis[c(2, 20)] <- NA
mod2 <- betadisper(dis, groups) ## warnings
mod2
permutest(mod2, control = permControl(nperm = 100))
## betadiver

### Indices of beta Diversity

**Description**

The function estimates any of the 24 indices of beta diversity reviewed by Koleff et al. (2003). Alternatively, it finds the co-occurrence frequencies for triangular plots (Koleff et al. 2003).

**Usage**

```r
betadiver(x, index = NA, order = FALSE, help = FALSE, ...)
## S3 method for class 'betadiver'
plot(x, ...)
## S3 method for class 'betadiver'
scores(x, triangular = TRUE, ...)
```

**Arguments**

- `x` Community data matrix, or the `betadiver` result for `plot` and `scores` functions.
- `index` The index of beta diversity as defined in Koleff et al. (2003), Table 1. You can use either the subscript of $\beta$ or the number of the index. See argument `help` below.
- `order` Order sites by increasing number of species. This will influence the configuration in the triangular plot and non-symmetric indices.
- `help` Show the numbers, subscript names and the defining equations of the indices and exit.
- `triangular` Return scores suitable for triangular plotting of proportions. If `FALSE`, returns a 3-column matrix of raw counts.
- `...` Other arguments to functions.
Details

The most commonly used index of beta diversity is \( \beta_w = \frac{S}{\alpha} - 1 \), where \( S \) is the total number of species, and \( \alpha \) is the average number of species per site (Whittaker 1960). A drawback of this model is that \( S \) increases with sample size, but the expectation of \( \alpha \) remains constant, and so the beta diversity increases with sample size. A solution to this problem is to study the beta diversity of pairs of sites. If we denote the number of species shared between two sites as \( a \) and the numbers of unique species (not shared) as \( b \) and \( c \), then \( S = a + b + c \) and \( \alpha = \frac{(2a + b + c)}{2} \) so that \( \beta_w = \frac{b + c}{2a + b + c} \). This is the Sørensen dissimilarity as defined in \texttt{vegan} function \texttt{vegdist} with argument \texttt{binary} = \texttt{TRUE}. Many other indices are dissimilarity indices as well.

Function \texttt{betadiver} finds all indices reviewed by Koleff et al. (2003). All these indices could be found with function \texttt{designdist} which uses different notation, but the current function provides a conventional shortcut. The function only finds the indices. The proper analysis must be done with functions such as \texttt{betadisper}, \texttt{adonis} or \texttt{mantel}.

The indices are directly taken from Table 1 of Koleff et al. (2003), and they can be selected either by the index number or the subscript name used by Koleff et al. The numbers, names and defining equations can be seen using \texttt{betadiver(help = TRUE)}. In all cases where there are two alternative forms, the one with the term \(-1\) is used. There are several duplicate indices, and the number of distinct alternatives is much lower than 24 formally provided. The formulations used in functions differ occasionally from those in Koleff et al. (2003), but they are still mathematically equivalent. With \texttt{index} = \texttt{NA}, no index is calculated, but instead an object of class \texttt{betadisper} is returned. This is a list of elements \( a \), \( b \) and \( c \). Function \texttt{plot} can be used to display the proportions of these elements in triangular plot as suggested by Koleff et al. (2003), and \texttt{scores} extracts the triangular coordinates or the raw scores. Function \texttt{plot} returns invisibly the triangular coordinates as an "ordiplot" object.

Value

With \texttt{index} = \texttt{NA}, the function returns an object of class "betadisper" with elements \( a, b, \) and \( c \). If \texttt{index} is specified, the function returns a "dist" object which can be used in any function analysing dissimilarities. For beta diversity, particularly useful functions are \texttt{betadisper} to study the betadiversity in groups, \texttt{adonis} for any model, and \texttt{mantel} to compare beta diversities to other dissimilarities or distances (including geographical distances). Although \texttt{betadiver} returns a "dist" object, some indices are similarities and cannot be used as such in place of dissimilarities, but that is a severe user error. Functions 10 ("j") and 11 ("sor") are two such similarity indices.

Warning

Some indices return similarities instead of dissimilarities.

Author(s)

Jari Oksanen

References


See Also
designdist for an alternative to implement all these functions, vegdist for some canned alternatives, and betadisper, adonis, mantel for analysing beta diversity objects.

Examples

```r
## Raw data and plotting
data(sipoo)
m <- betadiver(sipoo)
plot(m)
## The indices
betadiver(help=TRUE)
## The basic Whittaker index
d <- betadiver(sipoo, "w")
## This should be equal to Sorensen index (binary Bray-Curtis in
## vegan)
range(d - vegdist(sipoo, binary=TRUE))
```

---

**bgdispersal**

*Coefficients of Biogeographical Dispersal Direction*

**Description**

This function computes coefficients of dispersal direction between geographically connected areas, as defined by Legendre and Legendre (1984), and also described in Legendre and Legendre (1998, section 13.3.4).

**Usage**

```r
bgdispersal(mat, PAonly = FALSE, abc = FALSE)
```

**Arguments**

- **mat**: Data frame or matrix containing a community composition data table (species presence-absence or abundance data).
- **PAonly**: FALSE if the four types of coefficients, DD1 to DD4, are requested; TRUE if DD1 and DD2 only are sought (see Details).
- **abc**: If TRUE, return tables a, b and c used in DD1 and DD2.
Details

The signs of the DD coefficients indicate the direction of dispersal, provided that the asymmetry is significant. A positive sign indicates dispersal from the first (row in DD tables) to the second region (column); a negative sign indicates the opposite. A McNemar test of asymmetry is computed from the presence-absence data to test the hypothesis of a significant asymmetry between the two areas under comparison.

In the input data table, the rows are sites or areas, the columns are taxa. Most often, the taxa are species, but the coefficients can be computed from genera or families as well. DD1 and DD2 only are computed for presence-absence data. The four types of coefficients are computed for quantitative data, which are converted to presence-absence for the computation of DD1 and DD2. 

PAonly = FALSE indicates that the four types of coefficients are requested. PAonly = TRUE if DD1 and DD2 only are sought.

Value

Function bgdispersal returns a list containing the following matrices:

\[
DD1[j,k] = \frac{(a \times (b - c))/((a + b + c)^2)}
\]

\[
DD2[j,k] = \frac{(2 \times a \times (b - c))/((2 \times a + b + c) \times (a + b + c))}{\text{where } a, b, \text{ and } c
\]

have the same meaning as in the computation of binary similarity coefficients.

\[
DD3[j,k] = W \times (A - B)/((A + B - W)^2)
\]

\[
DD4[j,k] = 2 \times W \times (A - B)/((A + B) \times (A + B - W)) \text{ where } W = \text{sum(pmin(vector1, vector2)), } A = \text{sum(vector1), } B = \text{sum(vector2)}
\]

McNemar McNemar chi-square statistic of asymmetry (Sokal and Rohlf 1995): 
\[
2 \times (b \times log(b) + c \times log(c) - (b + c) \times log((b + c)/2))/q \text{ where } q = 1 + 1/(2 \times (b + c))
\]
(Williams correction for continuity)

prob.McNemar probabilities associated with McNemar statistics, chi-square test. H0: no asymmetry in \((b - c)\).

Note

The function uses a more powerful alternative for the McNemar test than the classical formula. The classical formula was constructed in the spirit of Pearson’s Chi-square, but the formula in this function was constructed in the spirit of Wilks Chi-square or the \(G\) statistic. Function mcnemar.test uses the classical formula. The new formula was introduced in vegan version 1.10-11, and the older implementations of bgdispersal used the classical formula.

Author(s)

Pierre Legendre, Departement de Sciences Biologiques, Universite de Montreal

References

**bioenv**


**Examples**

```r
mat <- matrix(c(32,15,14,10,70,30,100,4,10,30,25,0,18,0,40,
               0,0,20,0,0,0,4,0,30,20,0,0,0,25,74,42,1,45,89,5,16,16,20),
               4, 10, byrow=TRUE)
bgdispersal(mat)
```

---

**bioenv**

Best Subset of Environmental Variables with Maximum (Rank) Correlation with Community Dissimilarities

**Description**

Function finds the best subset of environmental variables, so that the Euclidean distances of scaled environmental variables have the maximum (rank) correlation with community dissimilarities.

**Usage**

```r
## Default S3 method:
bioenv(comm, env, method = "spearman", index = "bray",
       upto = ncol(env), trace = FALSE, partial = NULL, ...)
## S3 method for class 'formula'
bioenv(formula, data, ...)
```

**Arguments**

- `comm` Community data frame.
- `env` Data frame of continuous environmental variables.
- `method` The correlation method used in `cor`.
- `index` The dissimilarity index used for community data in `vegdist`.
- `upto` Maximum number of parameters in studied subsets.
- `formula, data` Model `formula` and data.
- `trace` Trace the advance of calculations
- `partial` Dissimilarities partialled out when inspecting variables in `env`.
- `...` Other arguments passed to `cor`.
bioenv

Details

The function calculates a community dissimilarity matrix using `vegdist`. Then it selects all possible subsets of environmental variables, `scales` the variables, and calculates Euclidean distances for this subset using `dist`. Then it finds the correlation between community dissimilarities and environmental distances, and for each size of subsets, saves the best result. There are \(2^p - 1\) subsets of \(p\) variables, and an exhaustive search may take a very, very, very long time (parameter `upto` offers a partial relief).

The function can be called with a model `formula` where the LHS is the data matrix and RHS lists the environmental variables. The formula interface is practical in selecting or transforming environmental variables.

With argument `partial` you can perform “partial” analysis. The partializing item must be a dissimilarity object of class `dist`. The `partial` item can be used with any correlation method, but it is strictly correct only for Pearson.

Clarke & Ainsworth (1993) suggested this method to be used for selecting the best subset of environmental variables in interpreting results of nonmetric multidimensional scaling (NMDS). They recommended a parallel display of NMDS of community dissimilarities and NMDS of Euclidean distances from the best subset of scaled environmental variables. They warned against the use of Procrustes analysis, but to me this looks like a good way of comparing these two ordinations.

Clarke & Ainsworth wrote a computer program BIO-ENV giving the name to the current function. Presumably BIO-ENV was later incorporated in Clarke’s PRIMER software (available for Windows). In addition, Clarke & Ainsworth suggested a novel method of rank correlation which is not available in the current function.

Value

The function returns an object of class `bioenv` with a `summary` method.

Note

If you want to study the ‘significance’ of `bioenv` results, you can use function `mantel` or `mantel.partial` which use the same definition of correlation. However, `bioenv` standardizes environmental variables to unit standard deviation using function `scale` and you must do the same in `mantel` for comparable results. Further, `bioenv` selects variables to maximize the Mantel correlation, and significance tests based on `a priori` selection of variables are biased.

Author(s)

Jari Oksanen

References

See Also

vegdist, dist, cor for underlying routines, isoMDS for ordination, procrustes for Procrustes analysis, protest for an alternative, and rankindex for studying alternatives to the default Bray-Curtis index.

Examples

# The method is very slow for large number of possible subsets.
# Therefore only 6 variables in this example.
data(varespec)
data(varechem)
sol <- bioenv(wisconsin(varespec) ~ log(N) + P + K + Ca + pH + Al, varechem)
sol
summary(sol)

biplot.rda

PCA biplot

Description

Draws a PCA biplot with species scores indicated by biplot arrows.

Usage

## S3 method for class 'rda'
biplot(x, choices = c(1, 2), scaling = 2,
       display = c("sites", "species"), type, xlim, ylim, col = c(1,2),
       const, ...)

Arguments

x
choices
scaling
display
type

A rda result object.
Ares
Scaling for species and site scores. Either species (2) or site (1) scores are scaled by eigenvalues, and the other set of scores is left unscaled, or with 3 both are scaled symmetrically by square root of eigenvalues. With negative scaling values in rda, species scores are divided by standard deviation of each species and multiplied with an equalizing constant. Unscaled raw scores stored in the result can be accessed with scaling = 0.
Scores shown. These must some of the alternatives "species" for species scores, and/or "sites" for site scores.
Type of plot: partial match to text for text labels, points for points, and none for setting frames only. If omitted, text is selected for smaller data sets, and points for larger. Can be of length 2 (e.g. type = c("text", "points")), in which case the first element describes how species scores are handled, and the second how site scores are drawn.
xlim, ylim  the x and y limits (min, max) of the plot.
col   Colours used for sites and species (in this order). If only one colour is given, it is used for both.
const  General scaling constant for scores.rda.
...   Other parameters for plotting functions.

Details

Produces a plot or biplot of the results of a call to rda. It is common for the "species" scores in a PCA to be drawn as biplot arrows that point in the direction of increasing values for that variable. The biplot.rda function provides a wrapper to plot.cca to allow the easy production of such a plot.

biplot.rda is only suitable for unconstrained models. If used on an ordination object with constraints, an error is issued.

If species scores are drawn using "text", the arrows are drawn from the origin to 0.85 * species score, whilst the labels are drawn at the species score. If the type used is "points", then no labels are drawn and therefore the arrows are drawn from the origin to the actual species score.

Value

The plot function returns invisibly a plotting structure which can be used by identify.ordiplot to identify the points or other functions in the ordiplot family.

Author(s)

Gavin Simpson, based on plot.cca by Jari Oksanen.

See Also

plot.cca, rda for something to plot, ordiplot for an alternative plotting routine and more support functions, and text, points and arrows for the basic routines.

Examples

data(dune)
mod <- rda(dune, scale = TRUE)
biplot(mod, scaling = 3)

## different type for species and site scores
biplot(mod, scaling = 3, type = c("text", "points"))
Description

Constrained Analysis of Principal Coordinates (CAP) is an ordination method similar to Redundancy Analysis (rda), but it allows non-Euclidean dissimilarity indices, such as Manhattan or Bray–Curtis distance. Despite this non-Euclidean feature, the analysis is strictly linear and metric. If called with Euclidean distance, the results are identical to rda, but capscale will be much more inefficient. Function capscale is a constrained version of metric scaling, a.k.a. principal coordinates analysis, which is based on the Euclidean distance but can be used, and is more useful, with other dissimilarity measures. The function can also perform unconstrained principal coordinates analysis, optionally using extended dissimilarities.

Usage

capscale(formula, data, distance = "euclidean", sqrt.dist = FALSE, 
comm = NULL, add = FALSE, dfun = vegdist, metaMDSdist = FALSE, 
na.action = na.fail, subset = NULL, ...)

Arguments

formula Model formula. The function can be called only with the formula interface. Most usual features of formula hold, especially as defined in cca and rda. The LHS must be either a community data matrix or a dissimilarity matrix, e.g., from vegdist or dist. If the LHS is a data matrix, function vegdist will be used to find the dissimilarities. The RHS defines the constraints. The constraints can be continuous variables or factors, they can be transformed within the formula, and they can have interactions as in a typical formula. The RHS can have a special term Condition that defines variables to be “partialled out” before constraints, just like in rda or cca. This allows the use of partial CAP.

data Data frame containing the variables on the right hand side of the model formula.

distance The name of the dissimilarity (or distance) index if the LHS of the formula is a data frame instead of dissimilarity matrix.

sqrt.dist Take square roots of dissimilarities. See section Notes below.

comm Community data frame which will be used for finding species scores when the LHS of the formula was a dissimilarity matrix. This is not used if the LHS is a data frame. If this is not supplied, the “species scores” are the axes of initial metric scaling (cmdscale) and may be confusing.

add Logical indicating if an additive constant should be computed, and added to the non-diagonal dissimilarities such that all eigenvalues are non-negative in the underlying Principal Co-ordinates Analysis (see cmdscale for details). This implements “correction method 2” of Legendre & Legendre (1998, p. 434). The negative eigenvalues are caused by using semi-metric or non-metric dissimilarities with basically metric cmdscale. They are harmless and ignored in capscale, but you also can avoid warnings with this option.
dfun: Distance or dissimilarity function used. Any function returning standard "dist" and taking the index name as the first argument can be used.

metaMDSdist: Use metaMDSdist similarly as in metaMDS. This means automatic data transformation and using extended flexible shortest path dissimilarities (function stepacross) when there are many dissimilarities based on no shared species.

na.action: Handling of missing values in constraints or conditions. The default (na.fail) is to stop with missing values. Choices na.omit and na.exclude delete rows with missing values, but differ in representation of results. With na.omit only non-missing site scores are shown, but na.exclude gives NA for scores of missing observations. Unlike in rda, no WA scores are available for missing constraints or conditions.

subset: Subset of data rows. This can be a logical vector which is TRUE for kept observations, or a logical expression which can contain variables in the working environment, data or species names of the community data (if given in the formula or as comm argument).

...: Other parameters passed to rda or to metaMDSdist.

Details

Canonical Analysis of Principal Coordinates (CAP) is simply a Redundancy Analysis of results of Metric (Classical) Multidimensional Scaling (Anderson & Willis 2003). Function capscale uses two steps: (1) it ordinates the dissimilarity matrix using cmdscale and (2) analyses these results using rda. If the user supplied a community data frame instead of dissimilarities, the function will find the needed dissimilarity matrix using vegdist with specified distance. However, the method will accept dissimilarity matrices from vegdist, dist, or any other method producing similar matrices. The constraining variables can be continuous or factors or both, they can have interaction terms, or they can be transformed in the call. Moreover, there can be a special term Condition just like in rda and cca so that “partial” CAP can be performed.

The current implementation differs from the method suggested by Anderson & Willis (2003) in three major points which actually make it similar to distance-based redundancy analysis (Legendre & Anderson 1999):

1. Anderson & Willis used the orthonormal solution of cmdscale, whereas capscale uses axes weighted by corresponding eigenvalues, so that the ordination distances are the best approximations of original dissimilarities. In the original method, later “noise” axes are just as important as first major axes.

2. Anderson & Willis take only a subset of axes, whereas capscale uses all axes with positive eigenvalues. The use of subset is necessary with orthonormal axes to chop off some “noise”, but the use of all axes guarantees that the results are the best approximation of original dissimilarities.

3. Function capscale adds species scores as weighted sums of (residual) community matrix (if the matrix is available), whereas Anderson & Willis have no fixed method for adding species scores.

With these definitions, function capscale with Euclidean distances will be identical to rda in eigenvalues and in site, species and biplot scores (except for possible sign reversal). However, it makes no sense to use capscale with Euclidean distances, since direct use of rda is much more efficient. Even with non-Euclidean dissimilarities, the rest of the analysis will be metric and linear.
The function can be also used to perform ordinary metric scaling a.k.a. principal coordinates analysis by using a formula with only a constant on the left hand side, or `comm ~ 1`. With `metaMDSdist = TRUE`, the function can do automatic data standardization and use extended dissimilarities using function `stepacross` similarly as in non-metric multidimensional scaling with `metaMDS`.

**Value**

The function returns an object of class `capscale` which is identical to the result of `rda`. At the moment, `capscale` does not have specific methods, but it uses `cca` and `rda` methods `plot.cca`, `scores.rda` etc. Moreover, you can use `anova.cca` for permutation tests of “significance” of the results.

**Note**

The function produces negative eigenvalues with non-Euclidean dissimilarity indices. The non-Euclidean component of inertia is given under the title `Imaginary` in the printed output. The `Total` inertia is the sum of all eigenvalues, but the sum of all non-negative eigenvalues is given as `Real Total` (which is higher than the `Total`). The ordination is based only on the real dimensions with positive eigenvalues, and therefore the proportions of inertia components only apply to the `Real Total` and ignore the `Imaginary` component. Permutation tests with `anova.cca` use only the real solution of positive eigenvalues. Function `adonis` gives similar significance tests, but it also handles the imaginary dimensions (negative eigenvalues) and therefore its results may differ from permutation test results of `capscale`.

If the negative eigenvalues are disturbing, you can use argument `add = TRUE` passed to `cmdscale`, or, preferably, a distance measure that does not cause these warnings. Alternatively, after square root transformation of distances (argument `sqrt.dist = TRUE`) many indices do not produce negative eigenvalues.

The inertia is named after the dissimilarity index as defined in the dissimilarity data, or as `unknown distance` if such an information is missing. Function `rda` usually divides the ordination scores by number of sites minus one. In this way, the inertia is variance instead of sum of squares, and the eigenvalues sum up to variance. Many dissimilarity measures are in the range 0 to 1, so they have already made a similar division. If the largest original dissimilarity is less than or equal to 4 (allowing for `stepacross`), this division is undone in `capscale` and original dissimilarities are used. Keyword `mean` is added to the inertia in cases where division was made, e.g. in Euclidean and Manhattan distances. Inertia is based on squared index, and keyword `squared` is added to the name of distance, unless data were square root transformed (argument `sqrt.dist = TRUE`). If an additive constant was used, keyword `euclidified` is added to the the name of inertia (argument `add = TRUE`).

**Author(s)**

Jari Oksanen

**References**


See Also

`rda`, `cca`, `plot.cca`, `anova.cca`, `vegdist`, `dist`, `cmdscale`.

Examples

```r
data(varespec)
data(varechem)
## Basic Analysis
care.cap <- capscale(varespec ~ N + P + K + Condition(Al), varechem, dist="bray")
care.cap
plot(care.cap)
anova(care.cap)
## Avoid negative eigenvalues with additive constant
capscale(varespec ~ N + P + K + Condition(Al), varechem, dist="bray", add =TRUE)
## Avoid negative eigenvalues by taking square roots of dissimilarities
capscale(varespec ~ N + P + K + Condition(Al), varechem, dist = "bray", sqrt.dist = TRUE)
## Principal coordinates analysis with extended dissimilarities
capscale(varespec ~ l, dist="bray", metaMDS = TRUE)
```

---

**cascadeKM**

*K*-means partitioning using a range of values of *K*

**Description**

This function is a wrapper for the `kmeans` function. It creates several partitions forming a cascade from a small to a large number of groups.

**Usage**

```r
cascadeKM(data, inf.gr, sup.gr, iter = 100, criterion = "calinski")
cIndexKM(y, x, index = "all")
```

## S3 method for class 'cascadeKM'

```r
plot(x, min.g, max.g, grpmts.plot = TRUE, sortg = FALSE, gridcol = NA, ...)
```
Arguments

data The data matrix. The objects (samples) are the rows.

inf.gr The number of groups for the partition with the smallest number of groups of the cascade (min).
sup.gr The number of groups for the partition with the largest number of groups of the cascade (max).
iter The number of random starting configurations for each value of $K$.
criterion The criterion that will be used to select the best partition. The default value is "calinski", which refers to the Calinski-Harabasz (1974) criterion. The simple structure index ("ssi") is also available. Other indices are available in function clustIndex (package cclust). In our experience, the two indices that work best and are most likely to return their maximum value at or near the optimal number of clusters are "calinski" and "ssi".
y Object of class "kmeans" returned by a clustering algorithm such as kmeans
x Data matrix where columns correspond to variables and rows to observations, or the plotting object in plot
index The available indices are: "calinski" and "ssi". Type "all" to obtain both indices. Abbreviations of these names are also accepted.
min.g, max.g The minimum and maximum numbers of groups to be displayed.
grpmts.plot Show the plot (TRUE or FALSE).
sortg Sort the objects as a function of their group membership to produce a more easily interpretable graph. See Details. The original object names are kept; they are used as labels in the output table x, although not in the graph. If there were no row names, sequential row numbers are used to keep track of the original order of the objects.
gridcol The colour of the grid lines in the plots. NA, which is the default value, removes the grid lines.
... Other parameters to the functions (ignored).

Details

The function creates several partitions forming a cascade from a small to a large number of groups formed by kmeans. Most of the work is performed by function cIndex which is based on the clustIndex function (package cclust). Some of the criteria were removed from this version because computation errors were generated when only one object was found in a group.

The default value is "calinski", which refers to the well-known Calinski-Harabasz (1974) criterion. The other available index is the simple structure index "ssi" (Dolnicar et al. 1999). In the case of groups of equal sizes, "calinski" is generally a good criterion to indicate the correct number of groups. Users should not take its indications literally when the groups are not equal in size. Type "all" to obtain both indices. The indices are defined as:

calinski: \( \frac{SSB/(K-1)}{SSW/(n-K)} \), where $n$ is the number of data points and $K$ is the number of clusters. $SSW$ is the sum of squares within the clusters while $SSB$ is the sum of squares among the clusters. This index is simply an $F$ (ANOVA) statistic.
**ssi:** the “Simple Structure Index” multiplicatively combines several elements which influence the interpretability of a partitioning solution. The best partition is indicated by the highest SSI value.

In a simulation study, Milligan and Cooper (1985) found that the Calinski-Harabasz criterion recovered the correct number of groups the most often. We recommend this criterion because, if the groups are of equal sizes, the maximum value of "calinski" usually indicates the correct number of groups. Another available index is the simple structure index "ssi". Users should not take the indications of these indices literally when the groups are not equal in size and explore the groups corresponding to other values of $K$.

Function `cascadeKM` has a plot method. Two plots are produced. The graph on the left has the objects in abscissa and the number of groups in ordinate. The groups are represented by colours. The graph on the right shows the values of the criterion ("calinski" or "ssi") for determining the best partition. The highest value of the criterion is marked in red. Points marked in orange, if any, indicate partitions producing an increase in the criterion value as the number of groups increases; they may represent other interesting partitions.

If `sortg=TRUE`, the objects are reordered by the following procedure: (1) a simple matching distance matrix is computed among the objects, based on the table of K-means assignments to groups, from $K = \text{min.g}$ to $K = \text{max.g}$. (2) A principal coordinate analysis (PCoA, Gower 1966) is computed on the centred distance matrix. (3) The first principal coordinate is used as the new order of the objects in the graph. A simplified algorithm is used to compute the first principal coordinate only, using the iterative algorithm described in Legendre & Legendre (1998, Table 9.10). The full distance matrix among objects is never computed; this avoids the problem of storing it when the number of objects is large. Distance values are computed as they are needed by the algorithm.

**Value**

Function `cascadeKM` returns an object of class `cascadeKM` with items:

- **partition** Table with the partitions found for different numbers of groups $K$, from $K = \text{inf.gr}$ to $K = \text{sup.gr}$.
- **results** Values of the criterion to select the best partition.
- **criterion** The name of the criterion used.
- **size** The number of objects found in each group, for all partitions (columns).

Function `cIndex` returns a vector with the index values. The maximum value of these indices is supposed to indicate the best partition. These indices work best with groups of equal sizes. When the groups are not of equal sizes, one should not put too much faith in the maximum of these indices, and also explore the groups corresponding to other values of $K$.

**Author(s)**

Marie-Helene Ouellette <Marie-Helene.Ouellette@UMontreal.ca>, Sebastien Durand <Sebastien.Durand@UMontreal.ca> and Pierre Legendre <Pierre.Legendre@UMontreal.ca>. Edited for vegan by Jari Oksanen.
References


Weingessel, A., Dimitriadou, A. and Dolnicar, S. *An Examination Of Indexes For Determining The Number Of Clusters In Binary Data Sets*, http://www.wu-wien.ac.at/am/wp99.htm

See Also

kmeans, clustIndex.

Examples

```r
# Partitioning a (10 x 10) data matrix of random numbers
mat <- matrix(runif(100),10,10)
res <- cascadeKM(mat, 2, 5, iter = 25, criterion = 'calinski')
toto <- plot(res)

# Partitioning an autocorrelated time series
vec <- sort(matrix(runif(30),30,1))
res <- cascadeKM(vec, 2, 5, iter = 25, criterion = 'calinski')
toto <- plot(res)

# Partitioning a large autocorrelated time series
# Note that we remove the grid lines
vec <- sort(matrix(runif(1000),1000,1))
res <- cascadeKM(vec, 2, 7, iter = 10, criterion = 'calinski')
toto <- plot(res, gridcol=NA)
```

Description

Function `cca` performs correspondence analysis, or optionally constrained correspondence analysis (a.k.a. canonical correspondence analysis), or optionally partial constrained correspondence analysis. Function `rda` performs redundancy analysis, or optionally principal components analysis. These are all very popular ordination techniques in community ecology.
Usage

```r
## S3 method for class 'formula'
cca(formula, data, na.action = na.fail, subset = NULL, ...)
## Default S3 method:
cca(X, Y, Z, ...)
## S3 method for class 'formula'
rda(formula, data, scale=FALSE, na.action = na.fail, subset = NULL, ...)
## Default S3 method:
rda(X, Y, Z, scale=FALSE, ...)
```

Arguments

- **formula**: Model formula, where the left hand side gives the community data matrix, 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.
- **X**: Community data matrix.
- **Y**: Constraining matrix, typically of environmental variables. Can be missing.
- **Z**: Conditioning matrix, the effect of which is removed (‘partialled out’) before next step. Can be missing.
- **scale**: Scale species to unit variance (like correlations).
- **na.action**: Handling of missing values in constraints or conditions. The default (`na.fail`) is to stop with missing value. Choice `na.omit` removes all rows with missing values. Choice `na.exclude` keeps all observations but gives `NA` for results that cannot be calculated. The WA scores of rows may be found also for missing values in constraints. Missing values are never allowed in dependent community data.
- **subset**: Subset of data rows. This can be a logical vector which is `TRUE` for kept observations, or a logical expression which can contain variables in the working environment, `data` or species names of the community data.
- **...**: Other arguments for `print` or `plot` functions (ignored in other functions).

Details

Since their introduction (ter Braak 1986), constrained, or canonical, correspondence analysis and its spin-off, redundancy analysis, have been the most popular ordination methods in community ecology. Functions `cca` and `rda` are similar to popular proprietary software Canoco, although the implementation is completely different. The functions are based on Legendre & Legendre’s (1998) algorithm: in `cca` Chi-square transformed data matrix is subjected to weighted linear regression on constraining variables, and the fitted values are submitted to correspondence analysis performed via singular value decomposition (`svd`). Function `rda` is similar, but uses ordinary, unweighted linear regression and unweighted SVD.
The functions can be called either with matrix-like entries for community data and constraints, or with formula interface. In general, the formula interface is preferred, because it allows a better control of the model and allows factor constraints.

In the following sections, \( X \), \( Y \) and \( Z \), although referred to as matrices, are more commonly data frames.

In the matrix interface, the community data matrix \( X \) must be given, but the other data matrices may be omitted, and the corresponding stage of analysis is skipped. If matrix \( Z \) is supplied, its effects are removed from the community matrix, and the residual matrix is submitted to the next stage. This is called ‘partial’ correspondence or redundancy analysis. If matrix \( Y \) is supplied, it is used to constrain the ordination, resulting in constrained or canonical correspondence analysis, or redundancy analysis. Finally, the residual is submitted to ordinary correspondence analysis (or principal components analysis). If both matrices \( Z \) and \( Y \) are missing, the data matrix is analysed by ordinary correspondence analysis (or principal components analysis).

Instead of separate matrices, the model can be defined using a model formula. The left hand side must be the community data matrix \( (X) \). The right hand side defines the constraining model. The constraints can contain ordered or unordered factors, interactions among variables and functions of variables. The defined contrasts are honoured in factor variables. The constraints can also be matrices (but not data frames). The formula can include a special term Condition for conditioning variables (“covariables”) “partialled out” before analysis. So the following commands are equivalent: \( \text{cca}(X, Y, Z) \), \( \text{cca}(X \sim Y + \text{Condition}(Z)) \), where \( Y \) and \( Z \) refer to constraints and conditions matrices respectively.

Constrained correspondence analysis is indeed a constrained method: CCA does not try to display all variation in the data, but only the part that can be explained by the used constraints. Consequently, the results are strongly dependent on the set of constraints and their transformations or interactions among the constraints. The shotgun method is to use all environmental variables as constraints. However, such exploratory problems are better analysed with unconstrained methods such as correspondence analysis (\texttt{decorana}, \texttt{corresp}) or non-metric multidimensional scaling (\texttt{isoMDS}) and environmental interpretation after analysis (\texttt{envfit}, \texttt{ordisurf}). CCA is a good choice if the user has clear and strong a priori hypotheses on constraints and is not interested in the major structure in the data set.

CCA is able to correct the curve artefact commonly found in correspondence analysis by forcing the configuration into linear constraints. However, the curve artefact can be avoided only with a low number of constraints that do not have a curvilinear relation with each other. The curve can reappear even with two badly chosen constraints or a single factor. Although the formula interface makes easy to include polynomial or interaction terms, such terms often produce curved artefacts (that are difficult to interpret), these should probably be avoided.

According to folklore, \texttt{rda} should be used with “short gradients” rather than \texttt{cca}. However, this is not based on research which finds methods based on Euclidean metric as uniformly weaker than those based on Chi-squared metric. However, standardized Euclidean distance may be an appropriate measures (see Hellinger standardization in \texttt{decostand} in particular).

Partial CCA (pCCA; or alternatively partial RDA) can be used to remove the effect of some conditioning or “background” or “random” variables or “covariables” before CCA proper. In fact, pCCA compares models \( \text{cca}(X \sim Z) \) and \( \text{cca}(X \sim Y + Z) \) and attributes their difference to the effect of \( Y \) cleansed of the effect of \( Z \). Some people have used the method for extracting “components of variance” in CCA. However, if the effect of variables together is stronger than sum of both separately, this can increase total Chi-square after “partialling out” some variation, and give negative
“components of variance”. In general, such components of “variance” are not to be trusted due to interactions between two sets of variables.

The functions have `summary` and `plot` methods which are documented separately (see `plot.cca`, `summary.cca`).

**Value**

Function `cca` returns a huge object of class `cca`, which is described separately in `cca.object`. Function `rda` returns an object of class `rda` which inherits from class `cca` and is described in `cca.object`. The scaling used in `rda` scores is described in a separate vignette with this package.

**Author(s)**

The responsible author was Jari Oksanen, but the code borrows heavily from Dave Roberts (http://labdsv.nr.usu.edu/).

**References**

The original method was by ter Braak, but the current implementations follows Legendre and Legendre. 

**See Also**

There is a special documentation for `plot.cca` and `summary.cca` functions with their helper functions (`text.cca`, `points.cca`, `scores.cca`). Function `anova.cca` provides an ANOVA like permutation test for the “significance” of constraints. Automatic model building (dangerous!) is discussed in `deviance.cca`. Diagnostic tools, prediction and adding new points in ordination are discussed in `goodness.cca` and `predict.cca`. Function `cca` (library `ade4`) provide alternative implementations of CCA (these are internally quite different). Function `capscale` is a non-Euclidean generalization of `rda`. The result object is described in `cca.object`. You can use `as.mlm` to refit ordination result as a multiple response linear model to find some descriptive statistics. Design decisions are explained in vignette ‘decision-vegan’ which also can be accessed with `vegandocs`.

**Examples**

```r
data(varespec)
data(varechem)
## Common but bad way: use all variables you happen to have in your
## environmental data matrix
vare.cca <- cca(varespec, varechem)
```
```r
vare.cca
plot(vare.cca)
## Formula interface and a better model
vare.cca <- cca(varespec ~ Al + P*(K + Baresoil), data=varechem)
plot(vare.cca)
## 'Partialling out' and 'negative components of variance'
cca(varespec ~ Ca, varechem)
## RDA
cca(varespec ~ Ca + Condition(pH), varechem)
## For further documentation:
## Not run:
vegandocs("decision")
## End(Not run)
```

---

**cca.object**

*Result Object from Constrained Ordination with cca, rda or capscale*

**Description**

Ordination methods `cca`, `rda` and `capscale` return similar result objects. Function `capscale` inherits from `rda` and `rda` inherits from `cca`. This inheritance structure is due to historic reasons: `cca` was the first of these implemented in vegan. Hence the nomenclature in `cca.object` reflects `cca`. This help page describes the internal structure of the `cca.object` for programmers.

**Value**

A `cca` object has the following elements:

- **call** the function call.
- **colsum, rowsum, rowsum.excluded** Column and row sums in `cca`. In `rda`, item `colsum` contains standard deviations of species and `rowsum` is NA. If some data were removed in `na.action`, the row sums of excluded observations are in item `rowsum.excluded` in `cca` (but not in `rda`). The `rowsum.excluded` add to the total (one) of `rowsum`.
- **grand.total** Grand total of community data in `cca` and NA in `rda`.
- **inertia** Text used as the name of inertia.
- **method** Text used as the name of the ordination method.
- **terms** The `terms` component of the `formula`. This is missing if the ordination was not called with `formula`.  

Further information on terms with three subitems: terms which is like the terms component above, but lists conditions and constraints similarly; xlev which lists the factor levels, and ordered which is TRUE to ordered factors. This is produced by vegan internal function ordiTerminfo, and it is needed in predict.cca with newdata. This is missing if the ordination was not called with formula.

tot.chi
Total inertia or the sum of all eigenvalues.

na.action
The result of na.action if missing values in constraints were handled by na.omit or na.exclude (or NULL if there were no missing values). This is a vector of indices of missing value rows in the original data and a class of the action, usually either "omit" or "exclude".

pCCA, CCA, CA
Actual ordination results for conditioned (partial), constrained and unconstrained components of the model. Any of these can be NULL if there is no corresponding component. Items pCCA, CCA and CA have similar structure, and contain following items:

- alias: The names of the aliased constraints or conditions. Function alias.cca does not access this item directly, but it finds the aliased variables and their defining equations from the QR item.
- biplot: Biplot scores of constraints. Only in CCA.
- centroids: (Weighted) centroids of factor levels of constraints. Only in CCA. Missing if the ordination was not called with formula.
- eig: Eigenvalues of axes. In CCA and CA.
- envcentre: (Weighted) means of the original constraining or conditioning variables. In pCCA and in CCA.
- Fit: The fitted values of standardized data matrix after fitting conditions. Only in pCCA.
- QR: The QR decomposition of explanatory variables as produced by qr. The constrained ordination algorithm is based on QR decomposition of constraints and conditions (environmental data). The environmental data are first centred in rda or weighted and centred in cca. The QR decomposition is used in many functions that access cca results, and it can be used to find many items that are not directly stored in the object. For examples, see coef.cca, coef.rda, vif.cca, permutest.cca, predict.cca, predict.rda, calibrate.cca. For possible uses of this component, see qr. In pCCA and CCA.
- rank: The rank of the ordination component.
- qrank: The rank of the constraints which is the difference of the ranks of QR decompositions in pCCA and CCA components. Only in CCA.
- tot.chi: Total inertia or the sum of all eigenvalues of the component.
- imaginary.chi, imaginary.rank: The sum and rank (number) of negative eigenvalues in capscale. Only in CA and only if negative eigenvalues were found in capscale.
- u: (Weighted) orthonormal site scores. Please note that scaled scores are not stored in the cca object, but they are made when the object is accessed with functions like scores.cca, summary.cca or plot.cca, or their rda
variants. Only in CCA and CA. In the CCA component these are the so-called linear combination scores.

\texttt{u.eig} \texttt{u} scaled by eigenvalues. There is no guarantee that any \texttt{.eig} variants of scores will be kept in the future releases.

\texttt{v} (Weighted) orthonormal species scores. If missing species were omitted from the analysis, this will contain attribute \texttt{na.action} that lists the omitted species. Only in CCA and CA.

\texttt{v.eig} \texttt{v} weighted by eigenvalues.

\texttt{wa} Site scores found as weighted averages (\texttt{cca}) or weighted sums (\texttt{rda}) of \texttt{v} with weights \texttt{Xbar}, but the multiplying effect of eigenvalues removed. These often are known as WA scores in \texttt{cca}. Only in CCA.

\texttt{wa.eig} The direct result of weighted averaging or weighted summation (matrix multiplication) with the resulting eigenvalue inflation.

\texttt{wa.excluded}, \texttt{u.excluded} WA scores for rows removed by \texttt{na.action} = \texttt{na.exclude} in CCA and CA components if these could be calculated.

\texttt{Xbar} The standardized data matrix after previous stages of analysis. In CCA this is after possible \texttt{pCCA} or after partialling out the effects of conditions, and in CA after both \texttt{pCCA} and CCA. In \texttt{cca} the standardization is Chi-square, and in \texttt{rda} centring and optional scaling by species standard deviations using function \texttt{scale}.

**NA Action and Subset**

If the constraints had missing values or subsets, and \texttt{na.action} was set to \texttt{na.exclude} or \texttt{na.omit}, the result will have some extra items:

\texttt{subset} subset evaluated as a logical vector (TRUE for included cases).

\texttt{na.action} The object returned by \texttt{na.action} which is a named vector of indices of removed items. The class of the vector is either "omit" or "exclude" as set by \texttt{na.action}. The \texttt{na.action} is applied after \texttt{subset} so that the indices refer to the subset data.

\texttt{residuals.zombie} A zombie vector of the length of number of rows in the residual ordination.

Some standard \texttt{R} functions find the number of valid observations from this vector, and it is provided for their use although this is useless in \texttt{vegan}.

\texttt{rowsum.excluded} Row sums of removed observations. Only in \texttt{cca}.

\texttt{CCA$wa.excluded} The WA scores for sites (found from community data) in constrained ordination if \texttt{na.action} was \texttt{na.exclude} and the scores could be calculated. The scores cannot be found for \texttt{capscale} and in partial ordination.

\texttt{CA$u.excluded} Row scores for sites in unconstrained ordination with identical conditions as above.

**capscale**

Function \texttt{capscale} may add some items depending on its arguments:

\texttt{metaMDSdist} The data set name if \texttt{metaMDSdist} = TRUE.

\texttt{ac} Additive constant used if \texttt{add} = TRUE.

\texttt{adjust} Adjustment of dissimilarities: see \texttt{capscale}, section “Notes”. 

Author(s)

Jari Oksanen

References


See Also

The description here provides a hacker’s interface. For more user friendly access to the `cca` object see `alias.cca`, `coef.cca`, `deviance.cca`, `predict.cca`, `scores.cca`, `summary.cca`, `vif.cca`, `weights.cca`, `spenvcor` or `rda` variants of these functions. You can use `as.mlm` to cast a `cca.object` into result of multiple response linear model (`lm`) in order to more easily find some statistics (which in principle could be directly found from the `cca.object` as well).

Examples

```r
# Some species will be missing in the analysis, because only a subset of sites is used below.
data(dune)
data(dune.env)
mod <- cca(dune[1:15,] ~ ., dune.env[1:15,])
# Look at the names of missing species
attr(mod$CCA$v, "na.action")
# Look at the names of the aliased variables:
mod$CCA$alias
# Access directly constrained weighted orthonormal species and site scores, constrained eigenvalues and margin sums.
spec <- mod$CCA$v
sites <- mod$CCA$u
eig <- mod$CCA$eig
rsum <- mod$srowsum
csum <- mod$scolsum
```

---

**CCorA**

*Canonical Correlation Analysis*

Description

Canonical correlation analysis, following Brian McArdrle’s unpublished graduate course notes, plus improvements to allow the calculations in the case of very sparse and collinear matrices, and permutation test of Pillai’s trace statistic.

Usage

```r
CCorA(Y, X, stand.Y=FALSE, stand.X=FALSE, nperm = 0, ...)
```

# S3 method for class 'CCorA'
biplot(x, plot.type="ov", xlabs, plot.axes = 1:2, int=0.5,
col.Y="red", col.X="blue", cex=c(0.7,0.9), ...)

Arguments

Y  
Left matrix (object class: matrix or data.frame).

X  
Right matrix (object class: matrix or data.frame).

stand.Y  
Logical; should Y be standardized?

stand.X  
Logical; should X be standardized?

nperm  
Numeric; number of permutations to evaluate the significance of Pillai’s trace, e.g. nperm=99 or nperm=999.

x  
CCoAR result object.

plot.type  
A character string indicating which of the following plots should be produced: "objects", "variables", "ov" (separate graphs for objects and variables), or "biplots". Any unambiguous subset containing the first letters of these names can be used instead of the full names.

xlabels  
Row labels. The default is to use row names, NULL uses row numbers instead, and NA suppresses plotting row names completely.

plot.axes  
A vector with 2 values containing the order numbers of the canonical axes to be plotted. Default: first two axes.

int  
Radius of the inner circles plotted as visual references in the plots of the variables. Default: int=0.5. With int=0, no inner circle is plotted.

col.Y  
Color used for objects and variables in the first data table (Y) plots. In biplots, the objects are in black.

col.X  
Color used for objects and variables in the second data table (X) plots.

cex  
A vector with 2 values containing the size reduction factors for the object and variable names, respectively, in the plots. Default values: cex=c(0.7,0.9).

...  
Other arguments passed to these functions. The function biplot.CCorA passes graphical arguments to biplot and biplot.default. CCorA currently ignores extra arguments.

Details

Canonical correlation analysis (Hotelling 1936) seeks linear combinations of the variables of Y that are maximally correlated to linear combinations of the variables of X. The analysis estimates the relationships and displays them in graphs. Pillai’s trace statistic is computed and tested parametrically (F-test); a permutation test is also available.

Algorithmic note – The blunt approach would be to read the two matrices, compute the covariance matrices, then the matrix $S_{12} \%\% \text{inv}(S_{22}) \%\% t(S_{12}) \%\% \text{inv}(S_{11})$. Its trace is Pillai’s trace statistic. This approach may fail, however, when there is heavy multicollinearity in very sparse data matrices. The safe approach is to replace all data matrices by their PCA object scores.

The function can produce different types of plots depending on the option chosen: "objects" produces two plots of the objects, one in the space of Y, the second in the space of X; "variables" produces two plots of the variables, one of the variables of Y in the space of Y, the second of the variables of X in the space of X; "ov" produces four plots, two of the objects and two of the variables; "biplots" produces two biplots, one for the first matrix (Y) and one for second matrix (X) solutions. For biplots, the function passes all arguments to biplot.default; consult its help page for configuring biplots.
Value

Function CCorA returns a list containing the following elements:

**Pillai**
Pillai’s trace statistic = sum of the canonical eigenvalues.

**Eigenvalues**
Canonical eigenvalues. They are the squares of the canonical correlations.

**CanCorr**
Canonical correlations.

**Mat.ranks**
Ranks of matrices Y and X.

**RDA.Rsquares**
Bimultivariate redundancy coefficients (R-squares) of RDAs of Y|X and X|Y.

**RDA.adj.Rsq**
RDA.Rsquares adjusted for n and the number of explanatory variables.

**nperm**
Number of permutations.

**p.Pillai**
Parametric probability value associated with Pillai’s trace.

**p.perm**
Permutational probability associated with Pillai’s trace.

**Cy**
Object scores in Y biplot.

**Cx**
Object scores in X biplot.

**corr.Y.Cy**
Scores of Y variables in Y biplot, computed as cor(Y,Cy).

**corr.X.Cx**
Scores of X variables in X biplot, computed as cor(X,Cx).

**corr.Y.Cx**
cor(Y,Cy) available for plotting variables Y in space of X manually.

**corr.X.Cy**
cor(X,Cx) available for plotting variables X in space of Y manually.

**call**
Call to the CCorA function.

Author(s)

Pierre Legendre, Departement de Sciences Biologiques, Universite de Montreal. Implemented in vegan with the help of Jari Oksanen.

References


Examples

```
# Example using two mite groups. The mite data are available in vegan
data(mite)
# Two mite species associations (Legendre 2005, Fig. 4)

group.1 <- c(1,2,4:8,10:15,17,19:22,24,26:30)
group.2 <- c(3,9,16,18,23,25,31:35)

# Separate Hellinger transformations of the two groups of species
mite.hel.1 <- decostand(mite[,group.1], "hel")
mite.hel.2 <- decostand(mite[,group.2], "hel")
rownames(mite.hel.1) = paste("S",1:nrow(mite),sep="")
```
rownames(mite.hel.2) = paste("S",1:nrow(mite),sep="")
out <- CCorA(mite.hel.1, mite.hel.2)
out
biplot(out, "ob") # Two plots of objects
biplot(out, "v", cex=c(0.7,0.6)) # Two plots of variables
biplot(out, "ov", cex=c(0.7,0.6)) # Four plots (2 for objects, 2 for variables)
biplot(out, "b", cex=c(0.7,0.6)) # Two biplots
biplot(out, xlabs = NA, plot.axes = c(3,5)) # Plot axes 3, 5. No object names
biplot(out, plot.type="biplots", xlabs = NULL) # Replace object names by numbers

# Example using random numbers. No significant relationship is expected
mat1 <- matrix(rnorm(60),20,3)
mat2 <- matrix(rnorm(100),20,5)
out2 = CCorA(mat1, mat2, nperm=99)
out2
biplot(out2, "b")

---

**contribdiv**  

**Contribution Diversity Approach**

**Description**

The contribution diversity approach is based in the differentiation of within-unit and among-unit diversity by using additive diversity partitioning and unit distinctiveness.

**Usage**

```r
contribdiv(comm, index = c("richness", "simpson"),
           relative = FALSE, scaled = TRUE, drop.zero = FALSE)
```

# S3 method for class 'contribdiv'
plot(x, sub, xlab, ylab, ylim, col, ...)

**Arguments**

- **comm**
  The community data matrix with samples as rows and species as column.

- **index**
  Character, the diversity index to be calculated.

- **relative**
  Logical, if TRUE then contribution diversity values are expressed as their signed deviation from their mean. See details.

- **scaled**
  Logical, if TRUE then relative contribution diversity values are scaled by the sum of gamma values (if index = "richness") or by sum of gamma values times the number of rows in comm (if index = "simpson"). See details.

- **drop.zero**
  Logical, should empty rows dropped from the result? If empty rows are not dropped, their corresponding results will be NAs.

- **x**
  An object of class "contribdiv".

- **sub**, **xlab**, **ylab**, **ylim**, **col**
  Graphical arguments passed to plot.

- **...**
  Other arguments passed to plot.
Details

This approach was proposed by Lu et al. (2007). Additive diversity partitioning (see adipart for more references) deals with the relation of mean alpha and the total (gamma) diversity. Although alpha diversity values often vary considerably. Thus, contributions of the sites to the total diversity are uneven. This site specific contribution is measured by contribution diversity components. A unit that has e.g. many unique species will contribute more to the higher level (gamma) diversity than another unit with the same number of species, but all of which common.

Distinctiveness of species \( j \) can be defined as the number of sites where it occurs \( (n_j) \), or the sum of its relative frequencies \( (p_j) \). Relative frequencies are computed sitewise and \( \sum_j p_{ij} \) at site \( i \) sum up to 1.

The contribution of site \( i \) to the total diversity is given by \( alpha_i = \sum_j (1/n_{ij}) \) when dealing with richness and \( alpha_i = \sum(p_{ij}*(1-p_{ij})) \) for the Simpson index.

The unit distinctiveness of site \( i \) is the average of the species distinctiveness, averaging only those species which occur at site \( i \). For species richness: \( alpha_i = mean(n_i) \) (in the paper, the second equation contains a typo, \( n \) is without index). For the Simpson index: \( alpha_i = mean(n_i) \).

The Lu et al. (2007) gives an in-depth description of the different indices.

Value

An object of class "contribdiv" in heriting from data frame.

Returned values are alpha, beta and gamma components for each sites (rows) of the community matrix. The "diff.coef" attribute gives the differentiation coefficient (see Examples).

Author(s)

Péter Sólymos, <solymos@ualberta.ca>

References


See Also

adipart, diversity

Examples

```r
## Artificial example given in
## Table 2 in Lu et al. 2007
x <- matrix(c(1/3,1/3,1/3,0,0,0,
0,0,1/3,1/3,1/3,0,
0,0,0,1/3,1/3,1/3),
3, 6, byrow = TRUE,
dimnames = list(LETTERS[1:3],letters[1:6]))
x
## Compare results with Table 2
```
contribdiv(x, "richness")
contribdiv(x, "simpson")
## Relative contribution (C values), compare with Table 2
(cd1 <- contribdiv(x, "richness", relative = TRUE, scaled = FALSE))
(cd2 <- contribdiv(x, "simpson", relative = TRUE, scaled = FALSE))
## Differentiation coefficients
attr(cd1, "diff.coef") # D_ST
attr(cd2, "diff.coef") # D_DT
## BCI data set
data(BCI)
opar <- par(mfrow=c(2,2))
plot(contribdiv(BCI, "richness"), main = "Absolute")
plot(contribdiv(BCI, "richness", relative = TRUE), main = "Relative")
plot(contribdiv(BCI, "simpson"))
plot(contribdiv(BCI, "simpson", relative = TRUE))
par(opar)

decorana **Detrended Correspondence Analysis and Basic Reciprocal Averaging**

**Description**

Performs detrended correspondence analysis and basic reciprocal averaging or orthogonal correspondence analysis.

**Usage**

```r
decorana(veg, iweigh=0, iresc=4, ira=0, mk=26, short=0,
        before=NULL, after=NULL)
```

## S3 method for class 'decorana'

```r
plot(x, choices=c(1,2), origin=TRUE,
     display=c("both","sites","species","none"),
     cex = 0.8, cols = c(1,2), type, xlim, ylim, ...)
```

## S3 method for class 'decorana'

```r
text(x, display = c("sites", "species"), labels,
     choices = 1:2, origin = TRUE, select, ...)
```

## S3 method for class 'decorana'

```r
points(x, display = c("sites", "species"),
       choices=1:2, origin = TRUE, select, ...)
```

## S3 method for class 'decorana'

```r
summary(object, digits=3, origin=TRUE,
        display=c("both", "species","sites","none"), ...)
```

## S3 method for class 'summary.decorana'

```r
```
print(x, head = NA, tail = head, ...)  

downweight(veg, fraction = 5)  

## S3 method for class 'decorana'  
scores(x, display=c("sites","species"), choices=1:4,  
origin=TRUE, ...)  

Arguments

veg: Community data, a matrix-like object.  
iweigh: Downweighting of rare species (0: no).  
iresc: Number of rescaling cycles (0: no rescaling).  
ira: Type of analysis (0: detrended, 1: basic reciprocal averaging).  
mk: Number of segments in rescaling.  
short: Shortest gradient to be rescaled.  
before: Hill’s piecewise transformation: values before transformation.  
after: Hill’s piecewise transformation: values after transformation – these must correspond to values in before.  
x, object: A decorana result object.  
choices: Axes shown.  
origin: Use true origin even in detrended correspondence analysis.  
display: Display only sites, only species, both or neither.  
cex: Plot character size.  
cols: Colours used for sites and species.  
type: Type of plots, partial match to "text", "points" or "none".  
labels: Optional text to be used instead of row names.  
select: Items to be displayed. This can either be a logical vector which is TRUE for displayed items or a vector of indices of displayed items.  
xlim, ylim: the x and y limits (min,max) of the plot.  
digits: Number of digits in summary output.  
head, tail: Number of rows printed from the head and tail of species and site scores. Default NA prints all.  
fraction: Abundance fraction where downweighting begins.  
...: Other arguments for plot function.

Details

In late 1970s, correspondence analysis became the method of choice for ordination in vegetation science, since it seemed better able to cope with non-linear species responses than principal components analysis. However, even correspondence analysis can produce an arc-shaped configuration of a single gradient. Mark Hill developed detrended correspondence analysis to correct two assumed
‘faults’ in correspondence analysis: curvature of straight gradients and packing of sites at the ends of the gradient.

The curvature is removed by replacing the orthogonalization of axes with detrending. In orthogonalization successive axes are made non-correlated, but detrending should remove all systematic dependence between axes. Detrending is performed using a five-segment smoothing window with weights (1,2,3,2,1) on mk segments — which indeed is more robust than the suggested alternative of detrending by polynomials. The packing of sites at the ends of the gradient is undone by rescaling the axes after extraction. After rescaling, the axis is supposed to be scaled by ‘SD’ units, so that the average width of Gaussian species responses is supposed to be one over whole axis. Other innovations were the piecewise linear transformation of species abundances and downweighting of rare species which were regarded to have an unduly high influence on ordination axes.

It seems that detrending actually works by twisting the ordination space, so that the results look non-curved in two-dimensional projections (‘lolly paper effect’). As a result, the points usually have an easily recognized triangular or diamond shaped pattern, obviously an artefact of detrending. Rescaling works differently than commonly presented, too. decorana does not use, or even evaluate, the widths of species responses. Instead, it tries to equalize the weighted variance of species scores on axis segments (parameter mk has only a small effect, since decorana finds the segment number from the current estimate of axis length). This equalizes response widths only for the idealized species packing model, where all species initially have unit width responses and equally spaced modes.

The summary method prints the ordination scores, possible prior weights used in downweighting, and the marginal totals after applying these weights. The plot method plots species and site scores. Classical decorana scaled the axes so that smallest site score was 0 (and smallest species score was negative), but summary, plot and scores use the true origin, unless origin = FALSE.

In addition to proper eigenvalues, the function also reports ‘decorana values’ in detrended analysis. These ‘decorana values’ are the values that the legacy code of decorana returns as ‘eigenvalues’. They are estimated internally during iteration, and it seems that detrending interferes the estimation so that these values are generally too low and have unclear interpretation. Moreover, ‘decorana values’ are estimated before rescaling which will change the eigenvalues. The proper eigenvalues are estimated after extraction of the axes and they are the ratio of biased weighted variances of site and species scores even in detrended and rescaled solutions. The ‘decorana values’ are provided only for the compatibility with legacy software, and they should not be used.

Value

decorana returns an object of class "decorana", which has print, summary and plot methods.

Note

decorana uses the central numerical engine of the original Fortran code (which is in the public domain), or about 1/3 of the original program. I have tried to implement the original behaviour, although a great part of preparatory steps were written in R language, and may differ somewhat from the original code. However, well-known bugs are corrected and strict criteria used (Oksanen & Minchin 1997).

Please note that there really is no need for piecewise transformation or even downweighting within decorana, since there are more powerful and extensive alternatives in R, but these options are
decorana

included for compliance with the original software. If a different fraction of abundance is needed in downweighting, function downweight must be applied before decorana. Function downweight indeed can be applied prior to correspondence analysis, and so it can be used together with cca, too.

The function finds only four axes: this is not easily changed.

Author(s)

Mark O. Hill wrote the original Fortran code, the R port was by Jari Oksanen.

References


See Also

For unconstrained ordination, non-metric multidimensional scaling in isoMDS may be more robust (see also metaMDS). Constrained (or ‘canonical’) correspondence analysis can be made with cca. Orthogonal correspondence analysis can be made with corresp, or with decorana or cca, but the scaling of results vary (and the one in decorana corresponds to scaling = -1 in cca.). See predict.decorana for adding new points to an ordination.

Examples

data(varespec)
vare.dca <- decorana(varespec)
vare.dca
summary(vare.dca)
plot(vare.dca)

### the detrending rationale:
gaussresp <- function(x,u) exp(-(x-u)^2/2)
x <- seq(0,6,length=15)  ## The gradient
u <- seq(-2,8,len=23)  ## The optima
pack <- outer(x,u,gaussresp)
matplot(x, pack, type="l", main="Species packing")
opar <- par(mfrow=c(2,2))
plot(scores(prcomp(pack)), asp=1, type="b", main="PCA")
plot(scores(decorana(pack, ira=1)), asp=1, type="b", main="CA")
plot(scores(decorana(pack)), asp=1, type="b", main="DCA")
plot(scores(cca(pack ~ x), dis="sites"), asp=1, type="b", main="CCA")

### Let's add some noise:
noisy <- (0.5 + runif(length(pack))) * pack
par(mfrow=c(2,1))
matplot(x, pack, type="l", main="Ideal model")
matplot(x, noisy, type="l", main="Noisy model")
par(mfrow=c(2,2))
plot(scores(prcomp(noisy)), type="b", main="PCA", asp=1)
plot(scores(decorana(noisy, ira=1)), type="b", main="CA", asp=1)
plot(scores(decorana(noisy)), type="b", main="DCA", asp=1)
plot(scores(cca(noisy ~ x), dis="sites"), asp=1, type="b", main="CCA")
par(opar)

decostand  

Standardization Methods for Community Ecology

Description

The function provides some popular (and effective) standardization methods for community ecologists.

Usage

decostand(x, method, MARGIN, range.global, logbase = 2, na.rm=FALSE, ...)

wisconsin(x)

Arguments

x  Community data, a matrix-like object.
method  Standardization method. See Details for available options.
MARGIN  Margin, if default is not acceptable. 1 = rows, and 2 = columns of x.
range.global  Matrix from which the range is found in method = "range". This allows using same ranges across subsets of data. The dimensions of MARGIN must match with x.
logbase  The logarithm base used in method = "log".
na.rm  Ignore missing values in row or column standardizations.
...  Other arguments to the function (ignored).

Details

The function offers following standardization methods for community data:

- total: divide by margin total (default MARGIN = 1).
- max: divide by margin maximum (default MARGIN = 2).
- freq: divide by margin maximum and multiply by the number of non-zero items, so that the average of non-zero entries is one (Oksanen 1983; default MARGIN = 2).
- normalize: make margin sum of squares equal to one (default MARGIN = 1).
- range: standardize values into range 0…1 (default MARGIN = 2). If all values are constant, they will be transformed to 0.
- standardize: scale x to zero mean and unit variance (default MARGIN = 2).
• **pa**: scale \( x \) to presence/absence scale (0/1).

• **chi.square**: divide by row sums and square root of column sums, and adjust for square root of matrix total (Legendre & Gallagher 2001). When used with the Euclidean distance, the distances should be similar to the Chi-square distance used in correspondence analysis. However, the results from `cmdscale` would still differ, since CA is a weighted ordination method (default `MARGIN = 1`).

• **hellinger**: square root of `method = "total"` (Legendre & Gallagher 2001).

• **log**: logarithmic transformation as suggested by Anderson et al. (2006): \( \log_b(x) + 1 \) for \( x > 0 \), where \( b \) is the base of the logarithm; zeros are left as zeros. Higher bases give less weight to quantities and more to presences, and \( \log_{base} = \infty \) gives the presence/absence scaling. Please note this is not \( \log(x + 1) \). Anderson et al. (2006) suggested this for their (strongly) modified Gower distance, but the standardization can be used independently of distance indices.

Standardization, as contrasted to transformation, means that the entries are transformed relative to other entries.

All methods have a default margin. `MARGIN=1` means rows (sites in a normal data set) and `MARGIN=2` means columns (species in a normal data set).

Command `wisconsin` is a shortcut to common Wisconsin double standardization where species (`MARGIN=2`) are first standardized by maxima (`max`) and then sites (`MARGIN=1`) by site totals (`tot`).

Most standardization methods will give nonsense results with negative data entries that normally should not occur in the community data. If there are empty sites or species (or constant with `method = "range"`), many standardization will change these into NaN.

**Value**

Returns the standardized data frame, and adds an attribute "decostand" giving the name of applied standardization "method".

**Note**

Common transformations can be made with standard \( \mathbb{R} \) functions.

**Author(s)**

Jari Oksanen and Etienne Laliberté (method = "log").

**References**


Examples

data(varespec)
strans <- decostand(varespec, "max")
apply(strans, 2, max)
strans <- wisconsin(varespec)

## Chi-square: PCA similar but not identical to CA.
## Use wcmdscale for weighted analysis and identical results.
strans <- decostand(varespec, "chi.square")
plot(procrustes(rda(strans), cca(varespec)))

---

designist  
**Design your own Dissimilarities**

Description

You can define your own dissimilarities using terms for shared and total quantities, number of rows and number of columns. The shared and total quantities can be binary, quadratic or minimum terms. In binary terms, the shared component is number of shared species, and totals are numbers of species on sites. The quadratic terms are cross-products and sums of squares, and minimum terms are sums of parallel minima and row totals.

Usage

designist(x, method = "(A+B-2*J)/(A+B)",
    terms = c("binary", "quadratic", "minimum"),
    abcd = FALSE, name)

Arguments

x Input data.

method Equation for your dissimilarities. This can use terms J for shared quantity, A and B for totals, N for the number of rows (sites) and P for the number of columns (species). The equation can also contain any R functions that accepts vector arguments and returns vectors of the same length.

terms How shared and total components are found. For vectors x and y the "quadratic" terms are J = sum(x*y), A = sum(x^2), B = sum(y^2), and "minimum" terms are J = sum(pmin(x,y)), A = sum(x) and B = sum(y), and "binary" terms are either of these after transforming data into binary form (shared number of species, and number of species for each row).

abcd Use 2x2 contingency table notation for binary data: a is the number of shared species, b and c are the numbers of species occurring only one of the sites but not in both, and d is the number of species that occur on neither of the sites.

name The name you want to use for your index. The default is to combine the method equation and terms argument.
Most popular dissimilarity measures in ecology can be expressed with the help of terms $J$, $A$ and $B$, and some also involve matrix dimensions $N$ and $P$. Some examples you can define in `designdist` are:

- $A+B-2*J$ "quadratic" squared Euclidean
- $A+B-2*J$ "minimum" Manhattan
- $(A+B-2*J)/(A+B)$ "minimum" Bray-Curtis
- $(A+B-2*J)/(A+B)$ "binary" Sørensen
- $(A+B-2*J)/(A+B-J)$ "minimum" Ružička
- $(A+B-2*J)/(A+B-J)$ "quadratic" (dis)similarity ratio
- $1-\text{phyper}(J-1, A, P-A, B)$ "binary" Ochiai
- $1-\text{phyper}(J-1, A, P-A, B)$ "binary" Raup-Crick

The function `designdist` can implement most dissimilarity indices in `vegdist` or elsewhere, and it can also be used to implement many other indices, amongst them, most of those described in Legendre & Legendre (1998). It can also be used to implement all indices of beta diversity described in Koleff et al. (2003), but there also is a specific function `betadiver` for the purpose.

If you want to implement binary dissimilarities based on the 2x2 contingency table notation, you can set `abcd = TRUE`. In this notation $a = J$, $b = A-J$, $c = B-J$, $d = P-A-B+J$. This notation is often used instead of the more more tangible default notation for reasons that are opaque to me.

**Value**

`designdist` returns an object of class `dist`.

**Note**

`designdist` does not use compiled code, and may be slow or use plenty of memory in large data sets. It is very easy to make errors when defining a function by hand. If an index is available in a function using compiled code, it is better to use the canned alternative.

**Author(s)**

Jari Oksanen

**References**


**See Also**

`vegdist, betadiver, dist`. 
Examples

## Arrhenius dissimilarity: the value of z in the species-area model
## S = c*A^z when combining two sites of equal areas, where S is the
## number of species, A is the area, and c and z are model parameters.
## The A below is not the area (which cancels out), but number of
## species in one of the sites, as defined in designdist().
data(BCI)
dis <- designdist(BCI, "(log(A+B-J)-log(A+B)+log(2))/log(2)")
## This can be used in clustering or ordination...
ordiplot(cmdscale(dis))
## ... or in analysing beta diversity (without gradients)
summary(dis)

deviance.cca

Statistics Resembling Deviance and AIC for Constrained Ordination

Description

The functions extract statistics that resemble deviance and AIC from the result of constrained correspondence analysis cca or redundancy analysis rda. These functions are rarely needed directly, but they are called by step in automatic model building. Actually, cca and rda do not have AIC and these functions are certainly wrong.

Usage

## S3 method for class 'cca'
deviance(object, ...)

## S3 method for class 'cca'
extractAIC(fit, scale = 0, k = 2, ...)

Arguments

object the result of a constrained ordination (cca or rda).
fit fitted model from constrained ordination.
scale optional numeric specifying the scale parameter of the model, see scale in step.
k numeric specifying the "weight" of the equivalent degrees of freedom (=:edf) part in the AIC formula.
... further arguments.
Details

The functions find statistics that resemble \texttt{deviance} and \texttt{AIC} in constrained ordination. Actually, constrained ordination methods do not have a log-Likelihood, which means that they cannot have AIC and deviance. Therefore you should not use these functions, and if you use them, you should not trust them. If you use these functions, it remains as your responsibility to check the adequacy of the result.

The deviance of \texttt{cca} is equal to the Chi-square of the residual data matrix after fitting the constraints. The deviance of \texttt{rda} is defined as the residual sum of squares. The deviance function of \texttt{rda} is also used for \texttt{capscale}. Function \texttt{extractAIC} mimics \texttt{extractAIC.lm} in translating deviance to AIC.

There is little need to call these functions directly. However, they are called implicitly in \texttt{step} function used in automatic selection of constraining variables. You should check the resulting model with some other criteria, because the statistics used here are unfounded. In particular, the penalty \( k \) is not properly defined, and the default \( k = 2 \) is not justified theoretically. If you have only continuous covariates, the \texttt{step} function will base the model building on magnitude of eigenvalues, and the value of \( k \) only influences the stopping point (but the variables with the highest eigenvalues are not necessarily the most significant in permutation tests in \texttt{anova.cca}). If you also have multi-class factors, the value of \( k \) will have a capricious effect in model building. The \texttt{step} function will pass arguments to \texttt{add1.cca} and \texttt{drop1.cca}, and setting \texttt{test = "permutation"} will provide permutation tests of each deletion and addition which can help in judging the validity of the model building.

Value

The \texttt{deviance} functions return “deviance”, and \texttt{extractAIC} returns effective degrees of freedom and “AIC”.

Note

These functions are unfounded and untested and they should not be used directly or implicitly. Moreover, usual caveats in using \texttt{step} are very valid.

Author(s)

Jari Oksanen

References


See Also

\texttt{cca, rda, anova.cca, step, extractAIC, add1.cca, drop1.cca}. 
**Examples**

```r
# The deviance of correspondence analysis equals Chi-square
data(dune)
data(dune.env)
chisq.test(dune)
deviance(cca(dune))
# Backward elimination from a complete model "dune ~ ."
ord <- cca(dune ~ ., dune.env)
ord
step(ord)
# Stepwise selection (forward from an empty model "dune ~ 1")
step(cca(dune ~ 1, dune.env), scope = formula(ord))
# ANOVA: added variable + the first left out
anova(cca(dune ~ Moisture + Management, dune.env), permut=200,
     by = "terms")
```

---

**dispindmorisita**  
*Morisita index of intraspecific aggregation*

**Description**

Calculates the Morisita index of dispersion, standardized index values, and the so called clumpedness and uniform indices.

**Usage**

```r
dispindmorisita(x, unique.rm = FALSE, crit = 0.05)
```

**Arguments**

- `x`: community data matrix, with sites (samples) as rows and species as columns.
- `unique.rm`: logical, if TRUE, unique species (occurring in only one sample) are removed from the result.
- `crit`: two-sided p-value used to calculate critical Chi-squared values.

**Details**

The Morisita index of dispersion is defined as (Morisita 1959, 1962):

\[
I_{\text{mor}} = n \times \frac{\sum (x_i^2) - \sum (x_i)}{(\sum (x_i)^2 - \sum (x_i))}
\]

where \(x_i\) is the count of individuals in sample \(i\), and \(n\) is the number of samples \((i = 1, 2, \ldots, n)\).  
\(I_{\text{mor}}\) has values from 0 to \(n\). In uniform (hyperdispersed) patterns its value falls between 0 and 1, in clumped patterns it falls between 1 and \(n\). For increasing sample sizes (i.e. joining neighbouring quadrats), \(I_{\text{mor}}\) goes to \(n\) as the quadrat size approaches clump size. For random patterns, \(I_{\text{mor}} = 1\) and counts in the samples follow Poisson frequency distribution.

The deviation from random expectation can be tested using critical values of the Chi-squared distribution with \(n - 1\) degrees of freedom. Confidence interval around 1 can be calculated by the
clumped $M_{clu}$ and uniform $M_{uni}$ indices (Hairston et al. 1971, Krebs 1999) (Chi2Lower and Chi2Upper refers to e.g. 0.025 and 0.975 quantile values of the Chi-squared distribution with $n - 1$ degrees of freedom, respectively, for $\alpha = 0.05$):

$$M_{clu} = \frac{(\text{Chi2Lower} - n + \text{sum}(x_i))}{(\text{sum}(x_i) - 1)}$$

$$M_{uni} = \frac{(\text{Chi2Upper} - n + \text{sum}(x_i))}{(\text{sum}(x_i) - 1)}$$

Smith-Gill (1975) proposed scaling of Morisita index from [0, n] interval into [-1, 1], and setting up -0.5 and 0.5 values as confidence limits around random distribution with rescaled value 0. To rescale the Morisita index, one of the following four equations apply to calculate the standardized index $I_{nst}$:

(a) $Imor \geq M_{clu} > 1$: $I_{nst} = 0.5 + 0.5 \frac{(Imor - M_{clu})}{(n - M_{clu})}$,

(b) $M_{clu} > Imor \geq 1$: $I_{nst} = 0.5 \frac{(Imor - 1)}{(M_{clu} - 1)}$,

(c) $1 > Imor > M_{uni}$: $I_{nst} = -0.5 \frac{(Imor - 1)}{(M_{uni} - 1)}$,

(d) $1 > M_{uni} > Imor$: $I_{nst} = -0.5 + 0.5 \frac{(Imor - M_{uni})}{M_{uni}}$.

Value

Returns a data frame with as many rows as the number of columns in the input data, and with four columns. Columns are: imor unstandardized Morisita index, mclu the clumpedness index, muni the uniform index, imst standardized Morisita index.

Note

A common error found in several papers is that when standardizing as in the case (b), the denominator is given as $M_{uni} - 1$. This results in a hiatus in the [0, 0.5] interval of the standardized index. The root of this typo is the book of Krebs (1999), see the Errata for the book (Page 217, http://www.zoology.ubc.ca/~krebs/downloads/errors_2nd_printing.pdf).

Author(s)

Péter Sólymos, <solymos@ualberta.ca>

References


Examples

data(dune)
x <- dispindmorisita(dune)
x
y <- dispindmorisita(dune, unique.rm = TRUE)
y
dim(x) ## with unique species
dim(y) ## unique species removed

Description

Function distconnected finds groups that are connected disregarding dissimilarities that are at or above a threshold or NA. The function can be used to find groups that can be ordinated together or transformed by stepacross. Function no.shared returns a logical dissimilarity object, where TRUE means that sites have no species in common. This is a minimal structure for distconnected or can be used to set missing values to dissimilarities.

Usage

distconnected(dis, toolong = 1, trace = TRUE)

no.shared(x)

Arguments

dis Dissimilarity data inheriting from class dist or an object, such as a matrix, that can be converted to a dissimilarity matrix. Functions vegdist and dist are some functions producing suitable dissimilarity data.

toolong Shortest dissimilarity regarded as NA. The function uses a fuzz factor, so that dissimilarities close to the limit will be made NA, too. If toolong = 0 (or negative), no dissimilarity is regarded as too long.

trace Summarize results of distconnected

x Community data.

Details

Data sets are disconnected if they have sample plots or groups of sample plots which share no species with other sites or groups of sites. Such data sets cannot be sensibly ordinated by any unconstrained method because these subsets cannot be related to each other. For instance, correspondence analysis will polarize these subsets with eigenvalue 1. Neither can such dissimilarities be transformed with stepacross, because there is no path between all points, and result will contain NAs. Function distconnected will find such subsets in dissimilarity matrices. The function will return a grouping vector that can be used for sub-setting the data. If data are connected, the result
vector will be all 1s. The connectedness between two points can be defined either by a threshold toolong or using input dissimilarities with NAs.

Function no.shared returns a dist structure having value TRUE when two sites have nothing in common, and value FALSE when they have at least one shared species. This is a minimal structure that can be analysed with distconnected. The function can be used to select dissimilarities with no shared species in indices which do not have a fixed upper limit.

Function distconnected uses depth-first search (Sedgewick 1990).

Value

Function distconnected returns a vector for observations using integers to identify connected groups. If the data are connected, values will be all 1. Function no.shared returns an object of class dist.

Author(s)

Jari Oksanen

References


See Also

vegdist or dist for getting dissimilarities, stepacross for a case where you may need distconnected, and for connecting points spantree.

Examples

```r
## There are no disconnected data in vegan, and the following uses an
## extremely low threshold limit for connectedness. This is for
## illustration only, and not a recommended practice.
data(dune)
dis <- vegdist(dune)
gr <- distconnected(dis, toolong=0.4)
# Make sites with no shared species as NA in Manhattan dissimilarities
dis <- vegdist(dune, "manhattan")
is.na(dis) <- no.shared(dune)
```

| diversity | Ecological Diversity Indices and Rarefaction Species Richness |

Description

Shannon, Simpson, and Fisher diversity indices and rarefied species richness for community ecologists.
## diversity

### Usage

```r
diversity(x, index = "shannon", MARGIN = 1, base = exp(1))
rarefy(x, sample, se = FALSE, MARGIN = 1)
rrarefy(x, sample)
drarefy(x, sample)
fisher.alpha(x, MARGIN = 1, se = FALSE, ...)
specnumber(x, MARGIN = 1)
```

### Arguments

- **x**: Community data, a matrix-like object or a vector.
- **index**: Diversity index, one of "shannon", "simpson" or "invsimpson".
- **MARGIN**: Margin for which the index is computed.
- **base**: The logarithm base used in `shannon`.
- **sample**: Subsample size for rarefying community, either a single value or a vector.
- **se**: Estimate standard errors.
- **...**: Parameters passed to `nlm`

### Details

Shannon or Shannon–Weaver (or Shannon–Wiener) index is defined as

\[ H' = -\sum_i p_i \log_b p_i, \]

where \( p_i \) is the proportional abundance of species \( i \) and \( b \) is the base of the logarithm. It is most popular to use natural logarithms, but some argue for base \( b = 2 \) (which makes sense, but no real difference).

Both variants of Simpson's index are based on \( D = \sum p_i^2 \). Choice `simpson` returns \( 1 - D \) and `invsimpson` returns \( 1/D \).

Function `rarefy` gives the expected species richness in random subsamples of size `sample` from the community. The size of `sample` should be smaller than total community size, but the function will silently work for larger `sample` as well and return non-rarefied species richness (and standard error = 0). If `sample` is a vector, rarefaction of all observations is performed for each sample size separately. Rarefaction can be performed only with genuine counts of individuals. The function `rarefy` is based on Hurlbert's (1971) formulation, and the standard errors on Heck et al. (1975).

Function `rrarefy` generates one randomly rarefied community data frame or vector of given `sample` size. The `sample` can be a vector giving the sample sizes for each row, and its values must be less or equal to observed number of individuals. The random rarefaction is made without replacement so that the variance of rarefied communities is rather related to rarefaction proportion than to the size of the sample.

Function `drarefy` returns probabilities that species occur in a rarefied community of size `sample`. The `sample` can be a vector giving the sample sizes for each row.

`fisher.alpha` estimates the \( \alpha \) parameter of Fisher's logarithmic series (see `fisherfit`). The estimation is possible only for genuine counts of individuals. The function can optionally return standard errors of \( \alpha \). These should be regarded only as rough indicators of the accuracy: the confidence limits of \( \alpha \) are strongly non-symmetric and the standard errors cannot be used in Normal inference.
Function `specnumber` finds the number of species. With `MARGIN = 2`, it finds frequencies of species. The function is extremely simple, and shortcuts are easy in plain R.

Better stories can be told about Simpson’s index than about Shannon’s index, and still grander narratives about rarefaction (Hurlbert 1971). However, these indices are all very closely related (Hill 1973), and there is no reason to despise one more than others (but if you are a graduate student, don’t drag me in, but obey your Professor’s orders). In particular, the exponent of the Shannon index is linearly related to inverse Simpson (Hill 1973) although the former may be more sensitive to rare species. Moreover, inverse Simpson is asymptotically equal to rarefied species richness in sample of two individuals, and Fisher’s $\alpha$ is very similar to inverse Simpson.

**Value**

A vector of diversity indices or rarefied species richness values. With a single `sample` and `se = TRUE`, function `rarefy` returns a 2-row matrix with rarefied richness ($S$) and its standard error (`se`). If `sample` is a vector in `rarefy`, the function returns a matrix with a column for each `sample size`, and if `se = TRUE`, rarefied richness and its standard error are on consecutive lines. With option `se = TRUE`, function `fisher.alpha` returns a data frame with items for $\alpha$ (`alpha`), its approximate standard errors (`se`), residual degrees of freedom (`df.residual`), and the code returned by `nlm` on the success of estimation.

**Author(s)**

Jari Oksanen and Bob O’Hara <bob.ohara@helsinki.fi> (`fisher.alpha`).

**References**


**See Also**

Function `renyi` for generalized Rényi diversity and Hill numbers.

**Examples**

data(BCI)
H <- diversity(BCI)
simp <- diversity(BCI, "simpson")
invsimp <- diversity(BCI, "inv")
## Unbiased Simpson of Hurlbert 1971 (eq. 5):
unbias.simp <- rarefy(BCI, 2) - 1
alpha <- fisher.alpha(BCI)
pairs(cbind(H, simp, invsimp, unbias.simp, alpha), pch="+", col="blue")
## Species richness (S) and Pielou's evenness (J):
S <- specnumber(BCI) ## rowSums(BCI > 0) does the same...
J <- H/log(S)

---

**dune**

*Vegetation and Environment in Dutch Dune Meadows.*

**Description**

The dune meadow vegetation data, `dune`, has cover class values of 30 species on 20 sites. The corresponding environmental data frame `dune.env` has following entries:

**Usage**

```r
data(dune)
data(dune.env)
```

**Format**

For `dune`, a data frame of observations of 30 species at 20 sites.

For `dune.env`, a data frame of 20 observations on the following 5 variables:

- **A1**: a numeric vector of thickness of soil A1 horizon.
- **Moisture**: an ordered factor with levels: 1 < 2 < 4 < 5.
- **Management**: a factor with levels: BF (Biological farming), HF (Hobby farming), NM (Nature Conservation Management), and SF (Standard Farming).
- **Use**: an ordered factor of land-use with levels: Hayfield < Haypastu < Pasture.
- **Manure**: an ordered factor with levels: 0 < 1 < 2 < 3 < 4.

**Source**


**Examples**

```r
data(dune)
data(dune.env)
```
dune.taxon  

Taxonomic Classification of Dune Meadow Species

Description

Classification table of the species in the dune data set.

Usage

data(dune.taxon)

Format

A data frame with 30 species (rows) classified into five taxonomic levels (columns).

Details

The classification of vascular plants is adapted from AGP (2003), and that of mosses from Hill et al. (2006).

Note

The data set was made to demonstrate taxondive, and will probably be removed after a better example is found.

References


Examples

data(dune.taxon)
eigenvals

---

**eigenvals**  
*Extract Eigenvalues from an Ordination Object*

---

### Description

Function extracts eigenvalues from an object that has them. Many multivariate methods return such objects.

### Usage

```r
eigenvals(x, ...)  
# S3 method for class 'cca'
eigenvals(x, constrained = FALSE, ...)
# S3 method for class 'eigenvals'
summary(object, ...)
```

### Arguments

- `x`  
  An object from which to extract eigenvalues.
- `object`  
  An `eigenvals` result object.
- `constrained`  
  Return only constrained eigenvalues.
- `...`  
  Other arguments to the functions (usually ignored)

### Details

This is a generic function that has methods for `cca`, `cmdscale`, `pcnm`, `prcomp`, `princomp`, `dudi` (of `ade4`), and `pca` and `pco` (of `labdsv`) result objects. The default method also extracts eigenvalues if the result looks like being from `eigen` or `svd`. Functions `prcomp` and `princomp` contain square roots of eigenvalues that all called standard deviations, but `eigenvals` function returns their squares. Function `svd` contains singular values, but function `eigenvals` returns their squares. For constrained ordination methods `cca`, `rda` and `capscale` the function returns the both constrained and unconstrained eigenvalues concatenated in one vector, but the partial component will be ignored. However, with argument `constrained = TRUE` only constrained eigenvalues are returned.

The summary of `eigenvals` result returns eigenvalues, proportion explained and cumulative proportion explained. The result object can have some negative eigenvalues (of `cmdscale`, `capscale`, `pcnm`) which correspond to imaginary axes of Euclidean mapping of non-Euclidean distances (Gower 1985). In these cases, the sum of absolute values of eigenvalues is used in calculating the proportions explained, and real axes (corresponding to positive eigenvalues) will only explain a part of total variation (Mardia et al. 1979, Gower 1985).

### Value

An object of class "eigenvals" which is a vector of eigenvalues.
Author(s)

Jari Oksanen.

References


See Also

eigen, svd, prcomp, princomp, cca, rda, capscale, wcmdscale, cca.object.

Examples

```r
data(varespec)
data(varechem)
mod <- cca(varespec ~ Al + P + K, varechem)
ev <- eigenvals(mod)
ev
summary(ev)
```

envfit

*Fits an Environmental Vector or Factor onto an Ordination*

Description

The function fits environmental vectors or factors onto an ordination. The projections of points onto vectors have maximum correlation with corresponding environmental variables, and the factors show the averages of factor levels.

Usage

```r
## Default S3 method:
envfit(ord, env, permutations = 999, strata, choices=c(1,2),
       display = "sites", w = weights(ord), na.rm = FALSE, ...)
## S3 method for class 'formula'
envfit(formula, data, ...)
## S3 method for class 'envfit'
plot(x, choices = c(1,2), arrow.mul, at = c(0,0), axis = FALSE,
     p.max = NULL, col = "blue", add = TRUE, ...)
## S3 method for class 'envfit'
scores(x, display, choices, ...)
vectorfit(X, P, permutations = 0, strata, w, ...)
factorfit(X, P, permutations = 0, strata, w, ...)
```
Arguments

ord An ordination object or other structure from which the ordination scores can be extracted (including a data frame or matrix of scores).

env Data frame, matrix or vector of environmental variables. The variables can be of mixed type (factors, continuous variables) in data frames.

X Matrix or data frame of ordination scores.

P Data frame, matrix or vector of environmental variable(s). These must be continuous for vectorfit and factors or characters for factorfit.

permutations Number of permutations for assessing significance of vectors or factors. Set to 0 to skip permutations.

formula, data Model formula and data.

na.rm Remove points with missing values in ordination scores or environmental variables. The operation is case-wise: the whole row of data is removed if there is a missing value and na.rm = TRUE.

x A result object from envfit.

choices Axes to plotted.

arrow.mul Multiplier for vector lengths. The arrows are automatically scaled similarly as in plot.cca if this is not given and add = TRUE.

at The origin of fitted arrows in the plot. If you plot arrows in other places then you probably have to specify arrow.mul.

axis Plot axis showing the scaling of fitted arrows.

p.max Maximum estimated $P$ value for displayed variables. You must calculate $P$ values with setting permutations to use this option.

col Colour in plotting.

add Results added to an existing ordination plot.

strata An integer vector or factor specifying the strata for permutation. If supplied, observations are permuted only within the specified strata.

display In fitting functions these are ordinary site scores or linear combination scores ("lc") in constrained ordination (cca, rda, capscale). In scores function they are either "vectors" or "factors" (with synonyms "bp" or "cn", resp.).

w Weights used in fitting (concerns mainly cca and decorana results which have nonconstant weights).

... Parameters passed to scores.

Details

Function envfit finds vectors or factor averages of environmental variables. Function plot.envfit adds these in an ordination diagram. If X is a data.frame, envfit uses factorfit for factor variables and vectorfit for other variables. If X is a matrix or a vector, envfit uses only vectorfit. Alternatively, the model can be defined a simplified model formula, where the left hand side must be an ordination result object or a matrix of ordination scores, and right
hand side lists the environmental variables. The formula interface can be used for easier selection and/or transformation of environmental variables. Only the main effects will be analysed even if interaction terms were defined in the formula.

Functions `vectorfit` and `factorfit` can be called directly. Function `vectorfit` finds directions in the ordination space towards which the environmental vectors change most rapidly and to which they have maximal correlations with the ordination configuration. Function `factorfit` finds averages of ordination scores for factor levels. Function `factorfit` treats ordered and unordered factors similarly.

If `permutations > 0`, the ‘significance’ of fitted vectors or factors is assessed using permutation of environmental variables. The goodness of fit statistic is squared correlation coefficient ($r^2$). For factors this is defined as $r^2 = 1 - s_s w / s_s t$, where $s_s w$ and $s_s t$ are within-group and total sums of squares. See `permutations` for additional details on permutation tests in Vegan.

User can supply a vector of prior weights $w$. If the ordination object has weights, these will be used. In practise this means that the row totals are used as weights with `cca` or `decorana` results. If you do not like this, but want to give equal weights to all sites, you should set $w = NULL$. The weighted fitting gives similar results to biplot arrows and class centroids in `cca`. For complete similarity between fitted vectors and biplot arrows, you should set display = "lc" (and possibly scaling = 2).

The lengths of arrows for fitted vectors are automatically adjusted for the physical size of the plot, and the arrow lengths cannot be compared across plots. For similar scaling of arrows, you must explicitly set the `arrow.mul` argument in the `plot` command.

The results can be accessed with `scores.envfit` function which returns either the fitted vectors scaled by correlation coefficient or the centroids of the fitted environmental variables.

Value

Functions `vectorfit` and `factorfit` return lists of classes `vectorfit` and `factorfit` which have a `print` method. The result object have the following items:

- `arrows` Arrow endpoints from `vectorfit`. The arrows are scaled to unit length.
- `centroids` Class centroids from `factorfit`.
- `r` Goodness of fit statistic: Squared correlation coefficient
- `permutations` Number of permutations.
- `pvals` Empirical P-values for each variable.

Function `envfit` returns a list of class `envfit` with results of `vectorfit` and `envfit` as items.

Function `plot.envfit` scales the vectors by correlation.

Note

Fitted vectors have become the method of choice in displaying environmental variables in ordination. Indeed, they are the optimal way of presenting environmental variables in Constrained Correspondence Analysis `cca`, since there they are the linear constraints. In unconstrained ordination the relation between external variables and ordination configuration may be less linear, and therefore other methods than arrows may be more useful. The simplest is to adjust the plotting symbol sizes
(cex, symbols) by environmental variables. Fancier methods involve smoothing and regression methods that abound in R, and ordisurf provides a wrapper for some.

Author(s)

Jari Oksanen. The permutation test derives from the code suggested by Michael Scroggie.

See Also

A better alternative to vectors may be ordisurf.

Examples

data(varespec)
data(varechem)
library(MASS)
ord <- metaMDS(varespec)
(fit <- envfit(ord, varechem, perm = 999))
scores(fit, "vectors")
plot(ord)
plot(fit)
plot(fit, p.max = 0.05, col = "red")
## Adding fitted arrows to CCA. We use "lc" scores, and hope
## that arrows are scaled similarly in cca and envfit plots
ord <- cca(varespec ~ Al + P + K, varechem)
plot(ord, type="p")
fit <- envfit(ord, varechem, perm = 999, display = "lc")
plot(fit, p.max = 0.05, col = "red")
## Class variables, formula interface, and displaying the
## inter-class variability with `ordispider'
data(dune)
data(dune.env)
attach(dune.env)
ord <- cca(dune)
fit <- envfit(ord ~ Moisture + A1, dune.env)
plot(ord, type = "n")
ordispider(ord, Moisture, col="skyblue")
points(ord, display = "sites", col = as.numeric(Moisture), pch=16)
plot(fit, cex=1.2, axis=TRUE)
Usage

fisherfit(x, ...)  
## S3 method for class 'fisherfit'  
confint(object, parm, level = 0.95, ...)  
## S3 method for class 'fisherfit'  
profile(fitted, alpha = 0.01, maxsteps = 20, del = zmax/5, ...)  
prestonfit(x, tiesplit = TRUE, ...)  
prestondistr(x, truncate = -1, ...)  
## S3 method for class 'prestonfit'  
plot(x, xlab = "Frequency", ylab = "Species", bar.col = "skyblue", line.col = "red", lwd = 2, ...)  
## S3 method for class 'prestonfit'  
lines(x, line.col = "red", lwd = 2, ...)  
veiledspec(x, ...)  
as.fisher(x, ...)  
as.preston(x, tiesplit = TRUE, ...)

Arguments

x Community data vector for fitting functions or their result object for plot functions.

object, fitted Fitted model.

parm Not used.

level The confidence level required.

alpha The extend of profiling as significance.

maxsteps Maximum number of steps in profiling.

del Step length.

tiesplit Split frequencies 1, 2, 4, 8 etc between adjacent octaves.

truncate Truncation point for log-Normal model, in log2 units. Default value −1 corresponds to the left border of zero Octave. The choice strongly influences the fitting results.

xlab, ylab Labels for x and y axes.

bar.col Colour of data bars.

line.col Colour of fitted line.

lwd Width of fitted line.

... Other parameters passed to functions. Ignored in prestonfit and tiesplit passed to as.preston in prestondistr.

Details

In Fisher's logarithmic series the expected number of species \( f \) with \( n \) observed individuals is \( f_n = \alpha n^{-\alpha} / n \) (Fisher et al. 1943). The estimation follows Kempton & Taylor (1974) and uses function \texttt{nlm}. The estimation is possible only for genuine counts of individuals. The parameter \( \alpha \)
Fisher’s diversity index, $\alpha$ and its standard error can be estimated with a separate function \texttt{fisher.alpha}. The parameter $x$ is taken as a nuisance parameter which is not estimated separately but taken to be $n/(n+\alpha)$. Helper function \texttt{as.fisher} transforms abundance data into Fisher frequency table.

Function \texttt{fisherfit} estimates the standard error of $\alpha$. However, the confidence limits cannot be directly estimated from the standard errors, but you should use function \texttt{confint} based on profile likelihood. Function \texttt{confint} uses function \texttt{confint.glm} of the \texttt{MASS} package, using \texttt{profile.fisherfit} for the profile likelihood. Function \texttt{profile.fisherfit} follows \texttt{profile.glm} and finds the $\tau$ parameter or signed square root of two times log-Likelihood profile. The profile can be inspected with a \texttt{plot} function which shows the $\tau$ and a dotted line corresponding to the Normal assumption: if standard errors can be directly used in Normal inference these two lines are similar.

Preston (1948) was not satisfied with Fisher’s model which seemed to imply infinite species richness, and postulated that rare species is a diminishing class and most species are in the middle of frequency scale. This was achieved by collapsing higher frequency classes into wider and wider “octaves” of doubling class limits: 1, 2, 3–4, 5–8, 9–16 etc. occurrences. It seems that Preston regarded frequencies 1, 2, 4, etc. as “tied” between octaves (Williamson & Gaston 2005). This means that only half of the species with frequency 1 are shown in the lowest octave, and the rest are transferred to the second octave. Half of the species from the second octave are transferred to the higher one as well, but this is usually not as large a number of species. This practise makes data look more lognormal by reducing the usually high lowest octaves. This can be achieved by setting argument \texttt{tiesplit = TRUE}. With \texttt{tiesplit = FALSE} the frequencies are not split, but all ones are in the lowest octave, all twos in the second, etc. Williamson & Gaston (2005) discuss alternative definitions in detail, and they should be consulted for a critical review of log-Normal model.

Any logseries data will look like lognormal when plotted in Preston’s way. The expected frequency $f$ at abundance octave $o$ is defined by $f_o = S_0 \exp(-\log_2(o) - \mu)^2/2/\sigma^2)$, where $\mu$ is the location of the mode and $\sigma$ the width, both in log$_2$ scale, and $S_0$ is the expected number of species at mode. The lognormal model is usually truncated on the left so that some rare species are not observed. Function \texttt{prestonfit} fits the truncated lognormal model as a second degree log-polynomial to the octave pooled data using Poisson (when \texttt{tiesplit = FALSE}) or quasi-Poisson (when \texttt{tiesplit = TRUE}). error. Function \texttt{prestondistr} fits left-truncated Normal distribution to log$_2$ transformed non-pooled observations with direct maximization of log-likelihood. Function \texttt{prestondistr} is modelled after function \texttt{fitdistr} which can be used for alternative distribution models.

The functions have common \texttt{print}, \texttt{plot} and \texttt{lines} methods. The \texttt{lines} function adds the fitted curve to the octave range with line segments showing the location of the mode and the width (sd) of the response. Function \texttt{as.preston} transforms abundance data to octaves. Argument \texttt{tiesplit} will not influence the fit in \texttt{prestondistr}, but it will influence the barplot of the octaves.

The total extrapolated richness from a fitted Preston model can be found with function \texttt{veiledspec}. The function accepts results both from \texttt{prestonfit} and from \texttt{prestondistr}. If \texttt{veiledspec} is called with a species count vector, it will internally use \texttt{prestonfit}. Function \texttt{specpool} provides alternative ways of estimating the number of unseen species. In fact, Preston’s lognormal model seems to be truncated at both ends, and this may be the main reason why its result differ from lognormal models fitted in Rank–Abundance diagrams with functions \texttt{rad.lognormal}. 
Value

The function `prestonfit` returns an object with fitted coefficients, and with observed (freq) and fitted (fitted) frequencies, and a string describing the fitting method. Function `prestondistr` omits the entry fitted. The function `fisherfit` returns the result of `nlm`, where item estimate is \( \alpha \). The result object is amended with the following items:

- `df.residuals` Residual degrees of freedom.
- `nuisance` Parameter \( x \).
- `fisher` Observed data from `as.fisher`.

Author(s)

Bob O’Hara <bob.ohara@helsinki.fi> (fisherfit) and Jari Oksanen.

References


See Also

diversity, fisher.alpha, radfit, specpool. Function `fitdistr` of MASS package was used as the model for `prestondistr`. Function `density` can be used for smoothed “non-parametric” estimation of responses, and `qqplot` is an alternative, traditional and more effective way of studying concordance of observed abundances to any distribution model.

Examples

data(BCI)
mod <- fisherfit(BCI[5,])
mod
plot(profile(mod))
confint(mod)
# prestonfit seems to need large samples
mod.oct <- prestonfit(colSums(BCI))
mod.ll <- prestondistr(colSums(BCI))
mod.oct
mod.ll
plot(mod.oct)
lines(mod.oct, line.col="blue3") # Different
## Smoothed density
den <- density(log2(colSums(BCI)))
lines(den$x, ncol(BCI)*den$y, lwd=2) # Fairly similar to mod.oct
goodness.cca

## Extrapolated richness
veiledspec(mod.oct)
veiledspec(mod.ll)

goodness.cca  Diagnostic Tools for [Constrained] Ordination (CCA, RDA, DCA, CA, PCA)

Description

Functions `goodness` and `inertcomp` can be used to assess the goodness of fit for individual sites or species. Function `vif.cca` and `alias.cca` can be used to analyse linear dependencies among constraints and conditions. In addition, there are some other diagnostic tools (see 'Details').

Usage

```r
## S3 method for class 'cca'
goodness(object, display = c("species", "sites"), choices, model = c("CCA", "CA"), statistic = c("explained", "distance"), summarize = FALSE, ...)
inertcomp(object, display = c("species", "sites"), statistic = c("explained", "distance"), proportional = FALSE)
spenvcor(object)
intersetcor(object)
vif.cca(object)
## S3 method for class 'cca'
alias(object, names.only = FALSE, ...)
```

Arguments

- `object` A result object from `cca`, `rda`, `capscale` or `decorana`.
- `display` Display "species" or "sites".
- `choices` Axes shown. Default is to show all axes of the "model".
- `model` Show constrained ("CCA") or unconstrained ("CA") results.
- `statistic` Statistic used: "explained" gives the cumulative percentage accounted for, "distance" shows the residual distances. Distances are not available for sites in constrained or partial analyses.
- `summarize` Show only the accumulated total.
- `proportional` Give the inertia components as proportional for the corresponding total.
- `names.only` Return only names of aliased variable(s) instead of defining equations.
- `...` Other parameters to the functions.
Details

Function **goodness** gives the diagnostic statistics for species or sites. The alternative statistics are the cumulative proportion of inertia accounted for by the axes, and the residual distance left unaccounted for. The conditional ("partialled out") constraints are always regarded as explained and included in the statistics.

Function **inertcomp** decomposes the inertia into partial, constrained and unconstrained components for each site or species. Instead of inertia, the function can give the total dispersion or distances from the centroid for each component.

Function **spenvcor** finds the so-called "species – environment correlation" or (weighted) correlation of weighted average scores and linear combination scores. This is a bad measure of goodness of ordination, because it is sensitive to extreme scores (like correlations are), and very sensitive to overfitting or using too many constraints. Better models often have poorer correlations. Function **ordispider** can show the same graphically.

Function **intersetcor** finds the so-called “interset correlation” or (weighted) correlation of weighted averages scores and constraints. The defined contrasts are used for factor variables. This is a bad measure since it is a correlation. Further, it focuses on correlations between single contrasts and single axes instead of looking at the multivariate relationship. Fitted vectors (envfit) provide a better alternative. Biplot scores (see scores.cca) are a multivariate alternative for (weighted) correlation between linear combination scores and constraints.

Function **vif.cca** gives the variance inflation factors for each constraint or contrast in factor constraints. In partial ordination, conditioning variables are analysed together with constraints. Variance inflation is a diagnostic tool to identify useless constraints. A common rule is that values over 10 indicate redundant constraints. If later constraints are complete linear combinations of conditions or previous constraints, they will be completely removed from the estimation, and no biplot scores or centroids are calculated for these aliased constraints. A note will be printed with default output if there are aliased constraints. Function **alias** will give the linear coefficients defining the aliased constraints, or only their names with argument names.only = TRUE.

Value

The functions return matrices or vectors as is appropriate.

Note

It is a common practise to use **goodness** statistics to remove species from ordination plots, but this may not be a good idea, as the total inertia is not a meaningful concept in **cca**, in particular for rare species.

Function **vif** is defined as generic in package **car** (vif), but if you have not loaded that package you must specify the call as vif.cca. Variance inflation factor is useful diagnostic tool for detecting nearly collinear constraints, but these are not a problem with algorithm used in this package to fit a constrained ordination.

Author(s)

Jari Oksanen. The vif.cca relies heavily on the code by W. N. Venables. alias.cca is a simplified version of alias.lm.
goodness.metaMDS

References


See Also

`cca`, `rda`, `capscale`, `decorana`, `vif`.

Examples

data(dune)
data(dune.env)
mod <- cca(dune ~ A1 + Management + Condition(Moisture), data=dune.env)
goodness(mod)
goodness(mod, summ = TRUE)
# Inertia components
inertcomp(mod, prop = TRUE)
inertcomp(mod, stat="d")
# vif.cca
vif.cca(mod)
# Aliased constraints
mod <- cca(dune ~ ., data=dune.env)
mod
vif.cca(mod)
alias(mod)
with(dune.env, table(Management, Manure))
# The standard correlations (not recommended)
spenvcor(mod)
intersetcor(mod)

---

**goodness.metaMDS**

*Goodness of Fit and Shepard Plot for Nonmetric Multidimensional Scaling*

**Description**

Function `goodness.metaMDS` find goodness of fit measure for points in nonmetric multidimensional scaling, and function `stressplot` makes a Shepard diagram.

**Usage**

```r
## S3 method for class 'metaMDS'
goodness(object, dis, ...)
stressplot(object, dis, pch, p.col = "blue", l.col = "red", lwd = 2, ...
```
Arguments

- **object**: A result object from `metaMDS` or `isoMDS`.
- **dis**: Dissimilarities. Normally this should not be used with `metaMDS`, but should be always used with `isoMDS`.
- **pch**: Plotting character for points. Default is dependent on the number of points.
- **p.col, l.col**: Point and line colours.
- **lwd**: Line width.
- **...**: Other parameters to functions, e.g. graphical parameters.

Details

Function `goodness.metaMDS` finds a goodness of fit statistic for observations (points). This is defined so that sum of squared values is equal to squared stress. Large values indicate poor fit.

Function `stressplot` is a wrapper to `Shepard` function in `MASS` package. It plots ordination distances against original dissimilarities, and draws a step line of the nonlinear fit. In addition, it adds to the graph two correlation-like statistics on the goodness of fit. The nonmetric fit is based on stress $S$ and defined as $\sqrt{1 - S^2}$. The “linear fit” is the correlation between fitted values and ordination distances.

Both functions can be used both with `metaMDS` and with `isoMDS`. With `metaMDS`, the functions try to reconstruct the dissimilarities using `metaMDSredist`, and dissimilarities should not be given. With `isoMDS` the dissimilarities must be given. In either case, the functions inspect that dissimilarities are consistent with current ordination, and refuse to analyse inconsistent dissimilarities. Function `goodness.metaMDS` is generic in `vegan`, but you must spell its name completely with `isoMDS` which has no class.

Value

Function `goodness` returns a vector of values. Function `stressplot` returns invisibly a `Shepard` object.

Author(s)

Jari Oksanen.

See Also

`metaMDS, isoMDS, Shepard`.

Examples

```r
data(varespec)
mod <- metaMDS(varespec)
stressplot(mod)
gof <- goodness(mod)
gof
plot(mod, display = "sites", type = "n")
points(mod, display = "sites", cex = gof/2)
```
**humpfit**

*No-interaction Model for Hump-backed Species Richness vs. Biomass*

**Description**

Function `humpfit` fits a no-interaction model for species richness vs. biomass data (Oksanen 1996). This is a null model that produces a hump-backed response as an artifact of plant size and density.

**Usage**

```r
humpfit(mass, spno, family = poisson, start)
## S3 method for class 'humpfit'
summary(object, ...)
## S3 method for class 'humpfit'
predict(object, newdata = NULL, ...)
## S3 method for class 'humpfit'
plot(x, xlab = "Biomass", ylab = "Species Richness", lwd = 2,
     l.col = "blue", p.col = 1, type = "b", ...)
## S3 method for class 'humpfit'
points(x, ...)
## S3 method for class 'humpfit'
lines(x, segments=101, ...)
## S3 method for class 'humpfit'
profile(fitted, parm = 1:3, alpha = 0.01, maxsteps = 20, del = zmax/5, ...)
```

**Arguments**

- `mass` Biomass.
- `spno` Species richness.
- `start` Vector of starting values for all three parameters.
- `family` Family of error distribution. Any `family` can be used, but the link function is always Fisher’s diversity model, and other link functions are silently ignored.
- `x, object, fitted` Result object of `humpfit`.
- `newdata` Values of `mass` used in `predict`. The original data values are used if missing.
- `xlab, ylab` Axis labels in `plot`.
- `lwd` Line width.
- `l.col, p.col` Line and point colour in `plot`.
- `type` Type of `plot`: "p" for observed points, "l" for fitted lines, "b" for both, and "n" for only setting axes.
- `segments` Number of segments used for fitted lines.
- `parm` Profiled parameters.
alpha, maxsteps, del

Parameters for profiling range and density.

... Other parameters to functions.

Details

The no-interaction model assumes that the humped species richness pattern along biomass gradient is an artifact of plant size and density (Oksanen 1996). For low-biomass sites, it assumes that plants have a fixed size, and biomass increases with increasing number of plants. When the sites becomes crowded, the number of plants and species richness reaches the maximum. Higher biomass is reached by increasing the plant size, and then the number of plants and species richness will decrease. At biomasses below the hump, plant number and biomass are linearly related, and above the hump, plant number is proportional to inverse squared biomass. The number of plants is related to the number of species by the relationship (link function) from Fisher’s log-series (Fisher et al. 1943).

The parameters of the model are:

1. hump: the location of the hump on the biomass gradient.
2. scale: an arbitrary multiplier to translate the biomass into virtual number of plants.
3. alpha: Fisher’s $\alpha$ to translate the virtual number of plants into number of species.

The parameters scale and alpha are intermingled and this function should not be used for estimating Fisher’s $\alpha$. Probably the only meaningful and interesting parameter is the location of the hump.

Function may be very difficult to fit and easily gets trapped into local solutions, or fails with non-Poisson families, and function profile should be used to inspect the fitted models. If you have loaded package MASS, you can use functions plot.profile, pairs.profile for graphical inspection of the profiles, and confint.profile.glm for the profile based confidence intervals.

The original model intended to show that there is no need to speculate about ‘competition’ and ‘stress’ (Al-Mufii et al. 1977), but humped response can be produced as an artifact of using fixed plot size for varying plant sizes and densities.

Value

The function returns an object of class "humpfit" inheriting from class "glm". The result object has specific summary, predict, plot, points and lines methods. In addition, it can be accessed by the following methods for glm objects: AIC, extractAIC, deviance, coef, residuals.glm (except type = "partial"), fitted, and perhaps some others. In addition, function ellipse.glm (package ellipse) can be used to draw approximate confidence ellipses for pairs of parameters, if the normal assumptions look appropriate.

Note

The function is a replacement for the original GLIM4 function at the archive of Journal of Ecology. There the function was represented as a mixed glm with one non-linear parameter (hump) and a special one-parameter link function from Fisher’s log-series. The current function directly applies non-linear maximum likelihood fitting using function nlm. Some expected problems with the current approach are:
• The function is discontinuous at hump and may be difficult to optimize in some cases (the lines will always join, but the derivative jumps).

• The function does not try very hard to find sensible starting values and can fail. The user may supply starting values in argument start if fitting fails.

• The estimation is unconstrained, but both scale and alpha should always be positive. Perhaps they should be fitted as logarithmic. Fitting Gamma family models might become easier, too.

Author(s)

Jari Oksanen

References


See Also

fishefit.profile.glm, confint.glm

Examples

```r
## Data approximated from Al-Mufti et al. (1977)
##
mass <- c(140,230,310,310,400,510,610,670,860,900,1050,1160,1900,2480)
spno <- c(1, 4, 3, 9, 18, 30, 20, 14, 3, 2, 3, 2, 5, 2)
sol <- humpfit(mass, spno)
summary(sol)  # Almost infinite alpha...
plot(sol)
# confint is in MASS, and implicitly calls profile.humpfit.
# Parameter 3 (alpha) is too extreme for profile and confint, and we
# must use only "hump" and "scale".
library(MASS)
plot(profile(sol, parm=1:2))
confint(sol, parm=c(1,2))
```
Description

Indicator power calculation of Halme et al. (2009) or the congruence between indicator and target species.

Usage

\[
\text{indpower}(x, \text{type} = 0)
\]

Arguments

- \(x\): Community data frame or matrix.
- \(\text{type}\): The type of statistic to be returned. See Details for explanation.

Details

Halme et al. (2009) described an index of indicator power defined as \(IP_I = \sqrt{a \times b}\), where \(a = S/O_I\) and \(b = 1 - (O_T - S)/(N - O_I)\). \(N\) is the number of sites, \(S\) is the number of shared occurrences of the indicator (I) and the target (T) species. \(O_I\) and \(O_T\) are number of occurrences of the indicator and target species. The \(\text{type}\) argument in the function call enables to choose which statistic to return. \(\text{type} = 0\) returns \(IP_I\), \(\text{type} = 1\) returns \(a\), \(\text{type} = 2\) returns \(b\).

Total indicator power (TIP) of an indicator species is the column mean (without its own value, see examples). Halme et al. (2009) explain how to calculate confidence intervals for these statistics.

Value

A matrix with indicator species as rows and target species as columns (this is indicated by the first letters of the row/column names).

Author(s)

Peter Solymos

References


See Also

\texttt{indval} (package \texttt{labdsv}) for the indicator species analysis of Dufrêne & Legendre. Function \texttt{beals} estimates individual cell probabilities of species occurrences.
isomap

Examples

data(dune)
## IP values
ip <- indpower(dune)
## and TIP values
diag(ip) <- NA
rowMeans(ip, na.rm=TRUE)

isomap

Isometric Feature Mapping Ordination

Description

The function performs isometric feature mapping which consists of three simple steps: (1) retain only some of the shortest dissimilarities among objects, (2) estimate all dissimilarities as shortest path distances, and (3) perform metric scaling (Tenenbaum et al. 2000).

Usage

isomap(dist, ndim=10, ...)
isomapdist(dist, epsilon, k, path = "shortest", fragmentedOK =FALSE, ...)
## S3 method for class 'isomap'
summary(object, axes = 4, ...)
## S3 method for class 'isomap'
plot(x, net = TRUE, n.col = "gray", ...)
rgl.isomap(x, web = "white", ...)

Arguments

dist Dissimilarities.
ndim Number of axes in metric scaling (argument k in cmdscale).
epsilon Shortest dissimilarity retained.
k Number of shortest dissimilarities retained for a point. If both epsilon and k are given, epsilon will be used.
p
path Method used in stepacross to estimate the shortest path, with alternatives "shortest" and "extended".
fragmentedOK What to do if dissimilarity matrix is fragmented. If TRUE, analyse the largest connected group, otherwise stop with error.
x, object An isomap result object.
axes Number of axes displayed.
net Draw the net of retained dissimilarities.
n.col Colour of drawn net segments.
web Colour of the web in rgl graphics.
... Other parameters passed to functions.
Details

The function isomap first calls function isomapdist for dissimilarity transformation, and then performs metric scaling for the result. All arguments to isomap are passed to isomapdist. The functions are separate so that the isomapdist transformation could be easily used with other functions than simple linear mapping of cmdscale.

Function isomapdist retains either dissimilarities equal or shorter to epsilon, or if epsilon is not given, at least k shortest dissimilarities for a point. Then a complete dissimilarity matrix is reconstructed using stepacross using either flexible shortest paths or extended dissimilarities (for details, see stepacross).

De’ath (1999) actually published essentially the same method before Tenenbaum et al. (2000), and De’ath’s function is available in xdiss in package mvpart. The differences are that isomap introduced the k criterion, whereas De’ath only used epsilon criterion. In practice, De’ath also retains higher proportion of dissimilarities than typical isomap.

In addition to the standard plot function, function rgl.isomap can make dynamic 3D plots that can be rotated on the screen. The functions is based on ordirgl, but it adds the connecting lines. The function passes extra arguments to scores and ordirgl functions so that you can select axes, or define colours and sizes of points.

Value

Function isomapdist returns a dissimilarity object similar to dist. Function isomap returns an object of class isomap with plot and summary methods. The plot function returns invisibly an object of class ordiplot. Function scores can extract the ordination scores.

Note

Tenenbaum et al. (2000) justify isomap as a tool of unfolding a manifold (e.g. a 'Swiss Roll'). Even with a manifold structure, the sampling must be even and dense so that dissimilarities along a manifold are shorter than across the folds. If data do not have such a manifold structure, the results are very sensitive to parameter values.

Author(s)

Jari Oksanen

References


See Also

The underlying functions that do the proper work are stepacross, distconnected and cmdscale. Package mvpart provides a parallel (but a bit different) implementation (xdiss). Moreover, vegan function metaMDS may trigger stepacross transformation, but usually only
for longest dissimilarities. The `plot` method of `vegan` minimum spanning tree function (`spantree`) has even more extreme way of isomapping things.

**Examples**

```r
## The following examples also overlay minimum spanning tree to
## the graphics in red.
op <- par(mar=c(4,4,1,1)+0.2, mfrow=c(2,2))
data(BCI)
dis <- vegdist(BCI)
tr <- spantree(dis)
pl <- ordiplot(cmdscale(dis), main="cmdscale")
lines(tr, pl, col="red")
ord <- isomap(dis, k=3)
ord
pl <- plot(ord, main="isomap k=3")
lines(tr, pl, col="red")
pl <- plot(isomap(dis, k=5), main="isomap k=5")
lines(tr, pl, col="red")
pl <- plot(isomap(dis, epsilon=0.45), main="isomap epsilon=0.45")
lines(tr, pl, col="red")
par(op)
## The following command requires user interaction
## Not run:
rgl.isomap(ord, size=4, color="hotpink")
## End(Not run)
```

---

**kendall.global**

*Kendall coefficient of concordance*

**Description**

Function `kendall.global` computes and tests the coefficient of concordance among several judges (variables, species) through a permutation test.

Function `kendall.post` carries out *a posteriori* tests of the contributions of individual judges (variables, species) to the overall concordance of their group through permutation tests.

If several groups of judges are identified in the data table, coefficients of concordance (`kendall.global`) or *a posteriori* tests (`kendall.post`) will be computed for each group separately. Use in ecology: to identify significant species associations.

**Usage**

```r
kendall.global(Y, group, nperm = 999, mult = "holm")
kendall.post(Y, group, nperm = 999, mult = "holm")
```
Arguments

- **Y**: Data file (data frame or matrix) containing quantitative or semiquantitative data. Rows are objects and columns are judges (variables). In community ecology, that table is often a site-by-species table.

- **group**: A vector defining how judges should be divided into groups. See example below. If groups are not explicitly defined, all judges in the data file will be considered as forming a single group.

- **nperm**: Number of permutations to be performed. Default is 999.

- **mult**: Correct P-values for multiple testing using the alternatives described in \(p.adjust\) and in addition "sidak" (see Details). The Bonferroni correction is overly conservative; it is not recommended. It is included to allow comparisons with the other methods.

Details

**Y** must contain quantitative data. They will be transformed to ranks within each column before computation of the coefficient of concordance.

The search for species associations described in Legendre (2005) proceeds in 3 steps:

1. Correlation analysis of the species. A possible method is to compute Ward’s agglomerative clustering of a matrix of correlations among the species. In detail: (1.1) compute a Pearson or Spearman correlation matrix (\(correl.matrix\)) among the species; (1.2) turn it into a distance matrix: \(mat.D = as.dist(1 - correl.matrix)\); (1.3) carry out Ward’s hierarchical clustering of that matrix using hclust: \(clust.ward = hclust(mat.D, "ward")\); (1.4) plot the dendrogram: \(plot(clust.ward, hang=-1)\); (1.5) cut the dendrogram in two groups, retrieve the vector of species membership: \(group.2 = cutree(clust.ward, k=2)\). (1.6) After steps 2 and 3 below, you may have to come back and try divisions of the species into \(k = 3, 4, 5, \ldots\) groups.

2. Compute global tests of significance of the 2 (or more) groups using the function \(kendall.global\) and the vector defining the groups. Groups that are not globally significant must be refined or abandoned.

3. Compute a posteriori tests of the contribution of individual species to the concordance of their group using the function \(kendall.post\) and the vector defining the groups. If some species have negative values for "Spearman.mean", this means that these species clearly do not belong to the group, hence that group is too inclusive. Go back to (1.5) and cut the dendrogram more finely. The left and right groups can be cut separately, independently of the levels along the dendrogram; write your own vector of group membership if \(cutree\) does not produce the desired groups.

The corrections used for multiple testing are applied to the list of P-values (P); they take into account the number of tests (k) carried out simultaneously (number of groups in \(kendall.global\), or number of species in \(kendall.post\)). The corrections are performed using function \(p.adjust\); see that function for the description of the correction methods. In addition, there is Šidák correction which defined as \(P_{corr} = 1 - (1 - P)^k\).

Value

A table containing the following information in rows. The columns correspond to the groups of "judges" defined in vector "group". When function \(kendall.post\) is used, there are as many tables as the number of predefined groups.
Kendall's coefficient of concordance, W.

F statistic. \( F = \frac{W(m-1)}{1-W} \) where \( m \) is the number of judges.

Probability associated with the F statistic, computed from the F distribution with
\( \nu_1 = n-1-(2/m) \) and \( \nu_2 = \nu_1*(m-1) \); \( n \) is the number of objects.

Probabilities associated with F, corrected using the method selected in parameter
mult. Shown only if there are more than one group.

Friedman’s chi-square statistic (Friedman 1937) used in the permutation test of
W.

Permutational probabilities, uncorrected.

Permutational probabilities corrected using the method selected in parameter
mult. Shown only if there are more than one group.

Mean of the Spearman correlations between the judge under test and all the other
judges in the same group.

Contribution of the judge under test to the overall concordance statistic for that
group.

Author(s)

F. Guillaume Blanchet, University of Alberta, and Pierre Legendre, Université de Montréal

References

Friedman, M. 1937. The use of ranks to avoid the assumption of normality implicit in the analysis

Kendall, M. G. and B. Babington Smith. 1939. The problem of m rankings. Annals of Mathematical

of Agricultural, Biological, and Environmental Statistics 10: 226-245.

Publications (in press).


See Also
cor, friedman.test, hclust, cutree, kmeans, cascadeKM, indval

Examples

data(mite)
mite.hel <- decostand(mite, "hel")
# Reproduce the results shown in Table 2 of Legendre (2005), a single group
mite.small <- mite.hel[c(4,9,14,22,31,34,45,53,61,69),c(13:15,23)]
kendall.global(mite.small, nperm=99)
kendall.post(mite.small, mult="holm", nperm=99)

# Reproduce the results shown in Tables 3 and 4 of Legendre (2005), 2 groups
group <- c(1,1,2,1,1,1,2,1,1,1,1,1,2,1,2,1,1,1,1,2,1,2,1,1,1,2,2,2,2,2)
kendall.global(mite.hel, group=group, nperm=99)
kendall.post(mite.hel, group=group, mult="holm", nperm=99)

# NOTE: 'nperm' argument usually needs to be larger than 99.
# It was set to this low value for demonstration purposes.

---

### linestack

**Plots One-dimensional Diagrams without Overwriting Labels**

**Description**

Function `linestack` plots vertical one-dimensional plots for numeric vectors. The plots are always labelled, but the labels are moved vertically to avoid overwriting.

**Usage**

```r
linestack(x, labels, cex = 0.8, side = "right", hoff = 2, air = 1.1,
at = 0, add = FALSE, axis = FALSE, ...)
```

**Arguments**

- `x`  
  Numeric vector to be plotted.
- `labels`  
  Text labels used instead of default (names of `x`).
- `cex`  
  Size of the labels.
- `side`  
  Put labels to the "right" or "left" of the axis.
- `hoff`  
  Distance from the vertical axis to the label in units of the width of letter “m”.
- `air`  
  Multiplier to string height to leave empty space between labels.
- `at`  
  Position of plot in horizontal axis.
- `add`  
  Add to an existing plot.
- `axis`  
  Add axis to the plot.
- `...`  
  Other graphical parameters to labels.

**Value**

The function returns invisibly the shifted positions of labels in user coordinates.

**Note**

The function always draws labelled diagrams. If you want to have unlabelled diagrams, you can use, e.g., `plot`, `stripchart` or `rug`.  

### Examples

```r
## First DCA axis
data(dune)
ord <- decorana(dune)
linestack(scores(ord, choices=1, display="sp"))
linestack(scores(ord, choices=1, display="si"), side="left", add=TRUE)
title(main="DCA axis 1")
```

### Description

A standard CEP name has four first letters of the generic name and four first letters of the specific epithet of a Latin name. The last epithet, that may be a subspecific name, is used in the current function. If the name has only one component, it is abbreviated to eight characters (see `abbreviate`). The returned names are made unique with function `make.unique` which adds numbers to the end of CEP names if needed.

### Usage

```r
make.cepnames(names)
```

### Arguments

- `names` The names to be formatted into CEP names.

### Details

Cornell Ecology Programs (CEP) used eight-letter abbreviations for species and site names. In species, the names were formed by taking four first letters of the generic name and four first letters of the specific or subspecific epithet. The CEP names were originally used, because old FORTRAN IV did not have CHARACTER data type, but text had to be stored in numerical variables, which in popular computers could hold four characters. In modern times, there is no reason for this limitation, but ecologists are used to these names, and they may be practical to avoid congestion in ordination plots.

### Value

Function returns CEP names.

### Note

The function is simpleminded and rigid. You must write a better one if you need.
Author(s)

Jari Oksanen

See Also

make.names, strsplit, substring, paste, abbreviate.

Examples

make.cepnames(c("Aa maderoi", "Poa sp.", "Cladina rangiferina", "Cladonia cornuta", "Cladonia cornuta var. groenlandica", "Cladonia rangiformis", "Bryoerythrophyllum"))
data(BCI)
colnames(BCI) <- make.cepnames(colnames(BCI))

mantel

Mantel and Partial Mantel Tests for Dissimilarity Matrices

Description

Function mantel finds the Mantel statistic as a matrix correlation between two dissimilarity matrices, and function mantel.partial finds the partial Mantel statistic as the partial matrix correlation between three dissimilarity matrices. The significance of the statistic is evaluated by permuting rows and columns of the first dissimilarity matrix.

Usage

mantel(xdis, ydis, method="pearson", permutations=999, strata)
mantel.partial(xdis, ydis, zdis, method = "pearson", permutations = 999, strata)

Arguments

xdis, ydis, zdis

Dissimilarity matrices or a dist objects.

method

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

permutations

Number of permutations in assessing significance.

strata

An integer vector or factor specifying the strata for permutation. If supplied, observations are permuted only within the specified strata.
Details

Mantel statistic is simply a correlation between entries of two dissimilarity matrices (some use cross products, but these are linearly related). However, the significance cannot be directly assessed, because there are $N(N-1)/2$ entries for just $N$ observations. Mantel developed asymptotic test, but here we use permutations of $N$ rows and columns of dissimilarity matrix. See permutations for additional details on permutation tests in Vegan.

Partial Mantel statistic uses partial correlation conditioned on the third matrix. Only the first matrix is permuted so that the correlation structure between second and first matrices is kept constant. Although mantel.partial silently accepts other methods than "pearson", partial correlations will probably be wrong with other methods.

The function uses cor, which should accept alternatives pearson for product moment correlations and spearman or kendall for rank correlations.

Value

The function returns a list of class mantel with following components:

- **Call** Function call.
- **method** Correlation method used, as returned by cor.test.
- **statistic** The Mantel statistic.
- **signif** Empirical significance level from permutations.
- **perm** A vector of permuted values.
- **permutations** Number of permutations.

Note

Legendre & Legendre (1998) say that partial Mantel correlations often are difficult to interpret.

Author(s)

Jari Oksanen

References

The test is due to Mantel, of course, but the current implementation is based on Legendre and Legendre.


See Also

cor for correlation coefficients, protest (“Procrustes test”) for an alternative with ordination diagrams, anosim and mrpp for comparing dissimilarities against classification. For dissimilarity matrices, see vegdist or dist. See bioenv for selecting environmental variables.
Examples

```r
## Is vegetation related to environment?
data(varespec)
data(varechem)
veg.dist <- vegdist(varespec) # Bray-Curtis
env.dist <- vegdist(scale(varechem), "euclid")
mantel(veg.dist, env.dist)
mantel(veg.dist, env.dist, method="spear")
```

mantel.correlog  Mantel Correlogram

Description


Usage

```r
mantel.correlog(D.eco, D.geo=NULL, XY=NULL, n.class=0, break.pts=NULL, cutoff=TRUE, r.type="pearson", nperm=999, mult="holm", progressive=TRUE)
## S3 method for class 'mantel.correlog'
plot(x, alpha=0.05, ...)
```

Arguments

- **D.eco**: An ecological distance matrix, with class either `dist` or `matrix`.
- **D.geo**: A geographic distance matrix, with class either `dist` or `matrix`. Provide either `D.geo` or `XY`. Default: `D.geo=NULL`.
- **XY**: A file of Cartesian geographic coordinates of the points. Default: `XY=NULL`.
- **n.class**: Number of classes. If `n.class=0`, the Sturges equation will be used unless break points are provided.
- **break.pts**: Vector containing the break points of the distance distribution. Provide `(n.class+1)` breakpoints, that is, a list with a beginning and an ending point. Default: `break.pts=NULL`.
- **cutoff**: For the second half of the distance classes, `cutoff = TRUE` limits the correlogram to the distance classes that include all points. If `cutoff = FALSE`, the correlogram includes all distance classes.
- **r.type**: Type of correlation in calculation of the Mantel statistic. Default: `r.type="pearson"`. Other choices are `r.type="spearman"` and `r.type="kendall"`, as in functions `cor` and `mantel`.
- **n.perm**: Number of permutations for the tests of significance. Default: `nperm=999`. For large data files, permutation tests are rather slow.

For the second half of the distance classes, cutoff = TRUE limits the correlogram to the distance classes that include all points. If cutoff = FALSE, the correlogram includes all distance classes.

Other choices are r.type=spearman and r.type=kendall, as in functions cor and mantel.

For large data files, permutation tests are rather slow.
Correct P-values for multiple testing. The correction methods are "holm" (default), "hochberg", "sidak", and other methods available in the `p.adjust` function: "bonferroni" (best known, but not recommended because it is overly conservative), "hommel", "BH", "BY", "fdr", and "none".

Default: `progressive=TRUE` for progressive correction of multiple-testing, as described in Legendre and Legendre (1998, p. 721). Test of the first distance class: no correction; second distance class: correct for 2 simultaneous tests; distance class k: correct for k simultaneous tests. `progressive=FALSE`: correct all tests for n.class simultaneous tests.

Output of `mantel.correlog`.

Significance level for the points drawn with black symbols in the correlogram. Default: `alpha=0.05`.

Other parameters passed from other functions.

A correlogram is a graph in which spatial correlation values are plotted, on the ordinate, as a function of the geographic distance classes among the study sites along the abscissa. In a Mantel correlogram, a Mantel correlation (Mantel 1967) is computed between a multivariate (e.g. multi-species) distance matrix of the user's choice and a design matrix representing each of the geographic distance classes in turn. The Mantel statistic is tested through a permutational Mantel test performed by `vegan`'s `mantel` function.

When a correction for multiple testing is applied, more permutations are necessary than in the no-correction case, to obtain significant p-values in the higher correlogram classes.

The `print.mantel.correlog` function prints out the correlogram. See examples.

A table with the distance classes as rows and the class indices, number of distances per class, Mantel statistics (computed using Pearson’s r, Spearman’s r, or Kendall’s tau), and p-values as columns. A positive Mantel statistic indicates positive spatial correlation. An additional column with p-values corrected for multiple testing is added unless `mult="none"`.

The number of distance classes.

The break points provided by the user or computed by the program.

The name of the correction for multiple testing. No correction: `mult="none"`.

A logical (TRUE, FALSE) value indicating whether or not a progressive correction for multiple testing was requested.

The number of distance classes for which Mantel tests have been computed and tested for significance.

The function call.

Pierre Legendre, Université de Montréal
References


Examples

```r
# Mite data available in "vegan"
data(mite)
data(mite.xy)
mite.hel <- decostand(mite, "hellinger")

# Detrend the species data by regression on the site coordinates
mite.hel.resid <- resid(lm(as.matrix(mite.hel) ~ ., data=mite.xy))

# Compute the detrended species distance matrix
mite.hel.D = dist(mite.hel.resid)

# Compute Mantel correlogram with cutoff, Pearson statistic
mite.correlog = mantel.correlog(mite.hel.D, XY=mite.xy, nperm=99)
summary(mite.correlog)
mite.correlog
# or: print(mite.correlog)
# or: print.mantel.correlog(mite.correlog)
plot(mite.correlog)

# Compute Mantel correlogram without cutoff, Spearman statistic
mite.correlog2 = mantel.correlog(mite.hel.D, XY=mite.xy, cutoff=FALSE, r.type="spearman", nperm=99)
summary(mite.correlog2)
mite.correlog2
plot(mite.correlog2)
```

**metaMDS**

Nonmetric Multidimensional Scaling with Stable Solution from Random Starts, Axis Scaling and Species Scores


**Description**

Function `metaMDS` uses `isoMDS` to perform Nonmetric Multidimensional Scaling (NMDS), but tries to find a stable solution using several random starts (function `initMDS`). In addition, it standardizes the scaling in the result, so that the configurations are easier to interpret (function `postMDS`), and adds species scores to the site ordination (function `wascores`).

**Usage**

```r
metaMDS(comm, distance = "bray", k = 2, trymax = 20, autotransform = TRUE,
         noshare = 0.1, wascores = TRUE, expand = TRUE, trace = 1,
         plot = FALSE, previous.best, old.wa = FALSE, ...)
## S3 method for class 'metaMDS'
plot(x, display = c("sites", "species"), choices = c(1, 2),
     type = "p", shrink = FALSE, ...)  # S3 method for class 'metaMDS'
points(x, display = c("sites", "species"),
       choices = c(1, 2), shrink = FALSE, select, ...)
## S3 method for class 'metaMDS'
text(x, display = c("sites", "species"), labels,
      choices = c(1, 2), shrink = FALSE, select, ...)
## S3 method for class 'metaMDS'
scores(x, display = c("sites", "species"), shrink = FALSE,
       choices, ...)
metaMDSdist(comm, distance = "bray", autotransform = TRUE, noshare = 0.1,
            trace = 1, commname, zerodist = "fail", distfun = vegdist, ...)
metaMDSiter(dist, k = 2, trymax = 20, trace = 1, plot = FALSE, previous.best,
             ...)
initMDS(x, k=2)
postMDS(X, dist, pc=TRUE, center=TRUE, halfchange, threshold=0.8,
        nthreshold=10, plot=FALSE, ...)  
metaMDSredist(object, ...)
metaMDSrotate(object, vec, ...)
```

**Arguments**

- `comm`: Community data. Alternatively, dissimilarities either as a `dist` structure or as a symmetric square matrix. In the latter case all other stages are skipped except random starts and centring and pc rotation of axes.
- `distance`: Dissimilarity index used in `vegdist`.
- `k`: Number of dimensions in `isoMDS`.
- `trymax`: Maximum number of dimensions in search of stable solution.
- `autotransform`: Use simple heuristics for possible data transformation (see below).
- `noshare`: Proportion of site pairs with no shared species to trigger `stepacross` to find flexible shortest paths among dissimilarities.
- `wascores`: Calculate species scores using function `wascores`.
- `expand`: Expand weighted averages of species in `wascores`. 
metaMDS

trace    Trace the function; trace = 2 or higher will be more voluminous.
plot     Graphical tracing: plot interim results. You may want to set \texttt{par(ask = TRUE)} with this option.
previous.best    Start searches from a previous solutions. Otherwise use \texttt{isoMDS} default for the starting solution.
old.wa    Use the old way of calculating WA scores for species: in vegan versions 1.12-5 and 1.11-2 WA scores were based on untransformed data even when data were transformed in analysis, but since then the similar transformation will be used in WA scores as in ordination.
x    Dissimilarity matrix for \texttt{isoMDS} or \texttt{plot} object.
choices    Axes shown.
type    Plot type: "p" for points, "t" for text, and "n" for axes only.
display    Display "sites" or "species".
shrink    Shrink back species scores if they were expanded originally.
labels    Optional test to be used instead of row names.
select    Items to be displayed. This can either be a logical vector which is \texttt{TRUE} for displayed items or a vector of indices of displayed items.
X    Configuration from multidimensional scaling.
commname    The name of \texttt{comm}: should not be given if the function is called directly.
zerodist    Handling of zero dissimilarities: either "fail" or "add" a small positive value, or "ignore".
distfun    Dissimilarity function. Any function returning a \texttt{dist} object and accepting argument \texttt{method} can be used (but some extra arguments may cause name conflicts).
dist    Dissimilarity matrix used in multidimensional scaling.
pc    Rotate to principal components.
center    Centre the configuration.
halfchange    Scale axes to half-change units. This defaults \texttt{TRUE} when dissimilarities were evaluated within \texttt{metaMDS} and the dissimilarity index has an upper limit of 1. If \texttt{FALSE}, the ordination dissimilarities are scaled to the same range as the input dissimilarities.
threshold    Largest dissimilarity used in half-change scaling.
nthreshold    Minimum number of points in half-change scaling.
object    A result object from \texttt{metaMDS}.
vec    A continuous site variable (vector).
...    Other parameters passed to functions.
Details

Non-metric Multidimensional Scaling (NMDS) is commonly regarded as the most robust unconstrained ordination method in community ecology (Minchin 1987). Functions initMDS and postMDS together with some other functions are intended to help run NMDS with isoMDS like recommended by Minchin (1987). Function metaMDS combines all recommendations into one command for a shotgun style analysis. The complete steps in metaMDS are:

1. Transformation: If the data values are larger than common class scales, the function performs a Wisconsin double standardization using wisconsin. If the values look very large, the function also performs sqrt transformation. Both of these standardization are generally found to improve the results. However, the limits are completely arbitrary (at present, data maximum 50 triggers sqrt and >9 triggers wisconsin). If you want to have a full control of the analysis, you should set autotransform = FALSE and make explicit standardization in the command.

2. Choice of dissimilarity: For a good result, you should use dissimilarity indices that have a good rank order relation to ordering sites along gradients (Faith et al. 1987). The default is Bray dissimilarity, because it often is the test winner. However, any other dissimilarity index in vegdist can be used. Function rankindex can be used for finding the test winner for you data and gradients.

3. Step-across dissimilarities: Ordination may be very difficult if a large proportion of sites have no shared species. In this case, the results may be improved with stepacross dissimilarities, or flexible shortest paths among all sites. The stepacross is triggered by option noshare. If you do not like manipulation of original distances, you should set noshare = 1.

4. NMDS with random starts: NMDS easily gets trapped into local optima, and you must start NMDS several times from random start to be confident that you have found the global solution. The default in isoMDS is to start from metric scaling (with cmdscale) which typically is close to a local optimum. The strategy in metaMDS is to first run a default isoMDS, or use the previous.best solution if supplied, and take its solution as the standard (Run 0). Then metaMDS starts isoMDS from several random starts (maximum number is given by trymax). If a solution is better (has a lower stress) than the previous standard, it is taken as the new standard. If the solution is better or close to a standard, metaMDS compares two solutions using Procrustes analysis using function procrustes with option symmetric = TRUE. If the two solutions are very similar in their Procrustes rmse and the largest residual is very small, the solutions are regarded as convergent and the best one is saved. Please note that the conditions are stringent, and you may have found good and relatively stable solutions although the function is not yet satisfied. Setting trace = TRUE will monitor the final stresses, and plot = TRUE will display Procrustes overlay plots from each comparison. This is the only step performed if input data (comm) were dissimilarities.

5. Scaling of the results: metaMDS will run postMDS for the final result. Function postMDS provides the following ways of “fixing” the indeterminacy of scaling and orientation of axes in NMDS: Centring moves the origin to the average of the axes. Principal components rotate the configuration so that the variance of points is maximized on first dimension (with function metaMDSrotate you can alternatively rotate the configuration so that the first axis is parallel to an environmental variable). Half-change scaling scales the configuration so that one unit means halving of community similarity from replicate similarity. Half-change scaling is based on closer dissimilarities where the relation between ordination distance and community dissimilarity is rather linear; the limit is controlled by parameter threshold. If there
are enough points below this threshold (controlled by the parameter `nthreshold`), dissimilarities are regressed on distances. The intercept of this regression is taken as the replicate dissimilarity, and half-change is the distance where similarity halves according to linear regression. Obviously the method is applicable only for dissimilarity indices scaled to 0...1, such as Kulczynski, Bray-Curtis and Canberra indices. If half-change scaling is not used, the ordination is scaled to the same range as the original dissimilarities.

6. Species scores: Function adds the species scores to the final solution as weighted averages using function `wasyncores` with given value of parameter `expand`. The expansion of weighted averages can be undone with `shrink = TRUE` in `plot` or `scores` functions, and the calculation of species scores can be suppressed with `wasyncores = FALSE`.

Value

Function `metaMDS` returns an object of class `metaMDS`. The final site ordination is stored in the item `points`, and species ordination in the item `species`. The other items store the information on the steps taken by the function. The object has `print`, `plot`, `points` and `text` methods. Functions `metaMDSdist` and `metaMDSredist` return `vegdist` objects. Function `initMDS` returns a random configuration which is intended to be used within `isoMDS` only. Functions `metaMDSiter` and `postMDS` returns the result of `isoMDS` with updated configuration.

Warning

The calculation of `wasyncores` for species was changed in `vegan` version 1.12-6. They are now based on the community data transformed similarly as in the ordination. Previously the species scores always were based on the original data. You can re-establish the old behaviour with argument `old.wa = TRUE`.

Note

Function `metaMDS` is a simple wrapper for `isoMDS` and some support functions. You can call these support functions separately for better control of results. Data transformation, dissimilarities and possible `stepacross` are made in function `metaMDSdist` which returns a dissimilarity result. Iterative search (with starting values from `initMDS`) is made in `metaMDSiter`. Processing of result configuration is done in `postMDS`, and species scores added by `wasyncores`. If you want to be more certain of reaching a global solution, you can compare results from several independent runs. You can also continue analysis from previous results or from your own configuration. Function does not save the used dissimilarity matrix, but `metaMDSredist` tries to reconstruct the used dissimilarities with original data transformation and possible `stepacross`.

The `metaMDS` function was designed to be used with community data. If you have other type of data, you should probably set some arguments to non-default values: probably at least `wasyncores`, `autotransform` and `noshare` should be `FALSE`. If you have negative data entries, `metaMDS` will set the previous to `FALSE` with a warning.

Author(s)

Jari Oksanen
References


See Also

`isoMDS, decostand, wascores, ordiplot`.

Examples

```r
## The recommended way of running NMDS (Minchin 1987)
##
data(dune)
library(MASS) ## isoMDS
# NMDS
sol <- metaMDS(dune)
sol
plot(sol, type="t")

## Start from previous best solution
sol2 <- metaMDS(dune, previous.best = sol)
```

mite

Oribatid Mite Data with Explanatory Variables

Description

Oribatid mite data. 70 soil cores collected by Daniel Borcard in 1989. See Borcard et al. (1992, 1994) for details.

Usage

```r
data(mite)
data(mite.env)
data(mite.pcnm)
data(mite.xy)
```

Format

There are three linked data sets: `mite` that contains the data on 35 species of Oribatid mites, `mite.env` that contains environmental data in the same sampling sites, `mite.xy` that contains geographic coordinates, and `mite.pcnm` that contains 22 PCNM base functions (columns) computed from the geographic coordinates of the 70 sampling sites (Borcard & Legendre 2002). The whole sampling area was 2.5 m x 10 m in size.

The fields in the environmental data are:
**SubsDens**  Substrate density (g/L)
**WatrCont**  Water content of the substrate (g/L)
**Substrate**  Substrate type, factor with levels Sphagn1, Sphagn2 Sphagn3 Sphagn Litter Barepeat Interface
**Shrub**  Shrub density, an ordered factor with levels $1 < 2 < 3$
**Topo**  Microtopography, a factor with levels Blanket and Hummock

**Source**
Pierre Legendre

**References**

**Examples**

```r
data(mite)
```

---

**model.matrix.cca**  Reconstruct Model Frame and Model Matrices of Constrained Ordination

**Description**
Function `model.frame.cca` reconstructs a `data.frame` with the variables used in the constrained ordination method (`cca`, `rda` or `capscale`). Function `model.matrix.cca` creates a list of design matrices used in constrained ordination. The items of the list are called `Conditions` and `Constraints`. If either partial (`Conditions`) or constrained component was missing, a single matrix is returned.

**Usage**

```r
## S3 method for class 'cca'
model.frame(formula, ...)
## S3 method for class 'cca'
model.matrix(object, ...)
```
Arguments

formula, object

A constrained ordination result object from which the needed information is extracted.

... Other arguments passed to the default method of the function.

Details

The constrained ordination method objects do not save data on model frame or design matrix, and the functions must reconstruct the information in the session. This will fail if the data sets and variables of the original model are unavailable.

Value

Returns a data frame (model.frame) or an unnamed matrix or a list of two matrices called Constraints and Conditions (model.matrix).

Author(s)

Jari Oksanen

See Also

model.frame, model.matrix.

Examples

data(dune)
data(dune.env)
mod <- cca(dune ~ poly(A1, 2) + Management + Use, dune.env)
model.frame(mod)
model.matrix(mod)

Description

Mitchell-Olds & Shaw test concerns the location of the highest (hump) or lowest (pit) value of a quadratic curve at given points. Typically, it is used to study whether the quadratic hump or pit is located within a studied interval. The current test is generalized so that it applies generalized linear models (glm) with link function instead of simple quadratic curve. The test was popularized in ecology for the analysis of humped species richness patterns (Mittelbach et al. 2001), but it is more general. With logarithmic link function, the quadratic response defines the Gaussian response model of ecological gradients (ter Braak & Looman 1986), and the test can be used for inspecting the location of Gaussian optimum within a given range of the gradient. It can also be used to replace Tokeshi’s test of “bimodal” species frequency distribution.
Usage

```r
MOStest(x, y, interval, ...)  
## S3 method for class 'MOStest'
plot(x, which = c(1,2,3,6), ...)  
fieller.MOStest(object, level = 0.95)  
## S3 method for class 'MOStest'
profile(fitted, alpha = 0.01, maxsteps = 10, del = zmax/5, ...)  
## S3 method for class 'MOStest'
confint(object, parm = 1, level = 0.95, ...)
```

Arguments

- **x**: The independent variable or plotting object in `plot`.
- **y**: The dependent variable.
- **interval**: The two points at which the test statistic is evaluated. If missing, the extremes of `x` are used.
- **which**: Subset of plots produced. Values `which`=1 and 2 define plots specific to `MOStest` (see Details), and larger values select graphs of `plot.lm` (minus 2).
- **object, fitted**: A result object from `MOStest`.
- **level**: The confidence level required.
- **alpha**: Maximum significance level allowed.
- **maxsteps**: Maximum number of steps in the profile.
- **del**: A step length parameter for the profile (see code).
- **parm**: Ignored.
- **...**: Other variables passed to functions. Function `MOStest` passes these to `glm` so that these can include `family`. The other functions pass these to underlying graphical functions.

Details

The function fits a quadratic curve $\mu = b_0 + b_1x + b_2x^2$ with given `family` and link function. If $b_2 < 0$, this defines a unimodal curve with highest point at $u = -b_2/(2b_3)$ (ter Braak & Looman 1986). If $b_2 > 0$, the parabola has a minimum at $u$ and the response is sometimes called “bimodal”.

The null hypothesis is that the extreme point $u$ is located within the interval given by points $p_1$ and $p_2$. If the extreme point $u$ is exactly at $p_1$, then $b_1 = 0$ on shifted axis $x - p_1$. In the test, origin of $x$ is shifted to the values $p_1$ and $p_2$, and the test statistic is based on the differences of deviances between the original model and model where the origin is forced to the given location using the standard `anova.glm` function (Oksanen et al. 2001). Mitchell-Olds & Shaw (1987) used the first degree coefficient with its significance as estimated by the `summary.glm` function. This give identical results with Normal error, but for other error distributions it is preferable to use the test based on differences in deviances in fitted models.

The test is often presented as a general test for the location of the hump, but it really is dependent on the quadratic fitted curve. If the hump is of different form than quadratic, the test may be insignificant.
Because of strong assumptions in the test, you should use the support functions to inspect the fit. Function `plot(..., which=1)` displays the data points, fitted quadratic model, and its approximate 95% confidence intervals (2 times SE). Function `plot` with `which = 2` displays the approximate confidence interval of the polynomial coefficients, together with two lines indicating the combinations of the coefficients that produce the evaluated points of $x$. Moreover, the cross-hair shows the approximate confidence intervals for the polynomial coefficients ignoring their correlations. Higher values of `which` produce corresponding graphs from `plot.lm`. That is, you must add 2 to the value of `which` in `plot.lm`.

Function `fieller.MOStest` approximates the confidence limits of the location of the extreme point (hump or pit) using Fieller’s theorem following ter Braak & Looman (1986). The test is based on quasideviance except if the `family` is `poisson` or `binomial`. Function `profile` evaluates the profile deviance of the fitted model, and `confint` finds the profile based confidence limits following Oksanen et al. (2001).

The test is typically used in assessing the significance of diversity hump against productivity gradient (Mittelbach et al. 2001). It also can be used for the location of the pit (deepest points) instead of the Tokeshi test. Further, it can be used to test the location of the the Gaussian optimum in ecological gradient analysis (ter Braak & Looman 1986, Oksanen et al. 2001).

**Value**

The function is based on `glm`, and it returns the result of object of `glm` amended with the result of the test. The new items in the `MOStest` are:

- `isHump` TRUE if the response is a hump.
- `isBracketed` TRUE if the hump or the pit is bracketed by the evaluated points.
- `hump` Sorted vector of location of the hump or the pit and the points where the test was evaluated.
- `coefficients` Table of test statistics and their significances.

**Note**

Function `fieller.MOStest` is based on package `optgrad` in the Ecological Archives (http://www.esapubs.org/archive/ecol/E082/015/default.htm) accompanying Oksanen et al. (2001). The Ecological Archive package `optgrad` also contains profile deviance method for the location of the hump or pit, but the current implementation of `profile` and `confint` rather follow the example of `profile.glm` and `confint.glm` in the MASS package.

**Author(s)**

Jari Oksanen

**References**


See Also

The no-interaction model can be fitted with `humpfit`.

Examples

```r
## The Al-Mufti data analysed in humpfit():
mass <- c(140,230,310,310,400,510,610,670,860,900,1050,1160,1900,2480)
spno <- c(1, 4, 3, 9, 18, 30, 20, 14, 3, 2, 3, 2, 5, 2)
mod <- MOStest(mass, spno)
## Insignificant
mod
## ... but inadequate shape of the curve
op <- par(mfrow=c(2,2), mar=c(4,4,1,1)+.1)
plot(mod)
## Looks rather like log-link with Poisson error and logarithmic biomass
mod <- MOStest(log(mass), spno, family=quasipoisson)
mod
plot(mod)
par(op)
## Confidence Limits
fieller.MOStest(mod)
confint(mod)
plot(profile(mod))
```

mrpp

Multi Response Permutation Procedure and Mean Dissimilarity Matrix

Description

Multiple Response Permutation Procedure (MRPP) provides a test of whether there is a significant difference between two or more groups of sampling units. Function `meandist` finds the mean within and between block dissimilarities.

Usage

```r
mrpp(dat, grouping, permutations = 999, distance = "euclidean",
     weight.type = 1, strata)
meandist(dist, grouping, ...)  
## S3 method for class 'meandist'
summary(object, ...)
## S3 method for class 'meandist'
plot(x, kind = c("dendrogram", "histogram"), cluster = "average",
     ylim, axes = TRUE, ...)```
Arguments

dat    data matrix or data frame in which rows are samples and columns are response variable(s), or a dissimilarity object or a symmetric square matrix of dissimilarities.
grouping  Factor or numeric index for grouping observations.
permutations Number of permutations to assess the significance of the MRPP statistic, delta.
distance  Choice of distance metric that measures the dissimilarity between two observations. See vegdist for options. This will be used if dat was not a dissimilarity structure of a symmetric square matrix.
weight.type choice of group weights. See Details below for options.
strata  An integer vector or factor specifying the strata for permutation. If supplied, observations are permuted only within the specified strata.
dist     A dist object of dissimilarities, such as produced by functions dist, vegdist or designdist..
object, x A meandist result object.
kind     Draw a dendrogram or a histogram; see Details.
cluster  A clustering method for the hclust function for kind = "dendrogram". Any hclust method can be used, but perhaps only "average" and "single" make sense.
ylim     Limits for vertical axes (optional).
axes     Draw scale for the vertical axis.
...      Further arguments passed to functions.

Details

Multiple Response Permutation Procedure (MRPP) provides a test of whether there is a significant difference between two or more groups of sampling units. This difference may be one of location (differences in mean) or one of spread (differences in within-group distance). Function mrpp operates on a data.frame matrix where rows are observations and responses data matrix. The response(s) may be uni- or multivariate. The method is philosophically and mathematically allied with analysis of variance, in that it compares dissimilarities within and among groups. If two groups of sampling units are really different (e.g. in their species composition), then average of the within-group compositional dissimilarities ought to be less than the average of the dissimilarities between two random collection of sampling units drawn from the entire population.

The mrpp statistic $\delta$ is the overall weighted mean of within-group means of the pairwise dissimilarities among sampling units. The choice of group weights is currently not clear. The mrpp function offers three choices: (1) group size ($n$), (2) a degrees-of-freedom analogue ($n-1$), and (3) a weight that is the number of unique distances calculated among $n$ sampling units ($n(n-1)/2$).

The mrpp algorithm first calculates all pairwise distances in the entire dataset, then calculates $\delta$. It then permutes the sampling units and their associated pairwise distances, and recalculates $\delta$ based on the permuted data. It repeats the permutation step permutations times. The significance test is the fraction of permuted deltas that are less than the observed delta, with a small sample correction. The function also calculates the change-corrected within-group agreement $A = 1 - \delta/E(\delta)$, where $E(\delta)$ is the expected $\delta$ assessed as the average of dissimilarities.
If the first argument `dat` can be interpreted as dissimilarities, they will be used directly. In other cases the function treats `dat` as observations, and uses `vegdist` to find the dissimilarities. The default distance is Euclidean as in the traditional use of the method, but other dissimilarities in `vegdist` also are available.

Function `meandist` calculates a matrix of mean within-cluster dissimilarities (diagonal) and between-cluster dissimilarities (off-diagonal elements), and an attribute `n` of grouping counts. Function `summary` finds the within-class, between-class and overall means of these dissimilarities, and the MRPP statistics with all `weight.type` options and the Classification Strength, CS (Van Sickle and Hughes, 2000). CS is defined for dissimilarities as $\bar{B} - \bar{W}$, where $\bar{B}$ is the mean between cluster dissimilarity and $\bar{W}$ is the mean within cluster dissimilarity with `weight.type = 1`. The function does not perform significance tests for these statistics, but you must use `mrpp` with appropriate `weight.type`. There is currently no significance test for CS, but `mrpp` with `weight.type = 1` gives the correct test for $\bar{W}$ and a good approximation for CS. Function `plot` draws a dendrogram or a histogram of the result matrix based on the within-group and between group dissimilarities. The dendrogram is found with the method given in the `cluster` argument using function `hclust`. The terminal segments hang to within-cluster dissimilarity. If some of the clusters are more heterogeneous than the combined class, the leaf segment are reversed. The histograms are based on dissimilarities, but are otherwise similar to those of Van Sickle and Hughes (2000): horizontal line is drawn at the level of mean between-cluster dissimilarity and vertical lines connect within-cluster dissimilarities to this line.

**Value**

The function returns a list of class `mrpp` with following items:

- `call`  
  Function call.
- `delta`  
  The overall weighted mean of group mean distances.
- `E.delta`  
  expected delta, under the null hypothesis of no group structure. This is the mean of original dissimilarities.
- `CS`  
  Classification strength (Van Sickle and Hughes, 2000). Currently not implemented and always `NA`.
- `n`  
  Number of observations in each class.
- `classdelta`  
  Mean dissimilarities within classes. The overall $\delta$ is the weighted average of these values with given `weight.type`.
- `Pvalue`  
  Significance of the test.
- `A`  
  A chance-corrected estimate of the proportion of the distances explained by group identity; a value analogous to a coefficient of determination in a linear model.
- `distance`  
  Choice of distance metric used; the "method" entry of the `dist` object.
- `weight.type`  
  The choice of group weights used.
- `boot.deltas`  
  The vector of "permuted deltas," the deltas calculated from each of the permuted datasets.
- `permutations`  
  The number of permutations used.
Note

This difference may be one of location (differences in mean) or one of spread (differences in within-group distance). That is, it may find a significant difference between two groups simply because one of those groups has a greater dissimilarities among its sampling units. Most mrpp models can be analysed with adonis which seems not suffer from the same problems as mrpp and is a more robust alternative.

Author(s)

M. Henry H. Stevens <HStevens@muohio.edu> and Jari Oksanen.

References


See Also

anosim for a similar test based on ranks, and mantel for comparing dissimilarities against continuous variables, and vegdist for obtaining dissimilarities, adonis is a more robust alternative in most cases.

Examples

data(dune)
data(dune.env)
dune.mrpp <- mrpp(dune, dune.env$Management)
dune.mrpp

# Save and change plotting parameters
def.par <- par(no.readonly = TRUE)
layout(matrix(1:2,nr=1))

ggplot(dune.ord <- metaMDS(dune), type="text", display="sites" )
ordihull(dune.ord, dune.env$Management)

with(dune.mrpp, {
fig.dist <- hist(boot.deltas, xlim=range(c(delta,boot.deltas)),
main="Test of Differences Among Groups")
abline(v=delta);
text(delta, 2*mean(fig.dist$counts), adj = -0.5,
expression(bold(delta)), cex=1.5 )
})
par(def.par)
## meandist
dune.md <- with(dune.env, meandist(vegdist(dune), Management))
The function `mso` adds an attribute `vario` to an object of class "cca" that describes the spatial partitioning of the `cca` object and performs an optional permutation test for the spatial independence of residuals. The function `plot.mso` creates a diagnostic plot of the spatial partitioning of the "cca" object.

**Usage**

```r
mso(object.cca, object.xy, grain = 1, round.up = FALSE, permutations = FALSE)
msoplot(x, alpha = 0.05, explained = FALSE, ...)
```

**Arguments**

- `object.cca`: An object of class `cca`, created by the `cca` or `rda` function.
- `object.xy`: A vector, matrix or data frame with the spatial coordinates of the data represented by `object.cca`. Must have the same number of rows as `object.cca$CA$Xbar` (see `cca.object`).
- `grain`: Interval size for distance classes.
- `round.up`: Determines the choice of breaks. If false, distances are rounded to the nearest multiple of grain. If true, distances are rounded to the upper multiple of grain.
- `permutations`: If false, suppresses the permutation test. If an integer, determines the number of permutations for the Mantel test of spatial independence of residual inertia.
- `x`: A result object of `mso`.
- `alpha`: Significance level for the two-sided permutation test of the Mantel statistic for spatial independence of residual inertia and for the point-wise envelope of the variogram of the total variance. A Bonferroni-type correction can be achieved by dividing the overall significance value (e.g. 0.05) by the number of distance classes.
- `explained`: If false, suppresses the plotting of the variogram of explained variance.
- `...`: Other arguments passed to functions.
Details

The Mantel test is an adaptation of the function `mantel` of the `vegan` package to the parallel testing of several distance classes. It compares the mean inertia in each distance class to the pooled mean inertia of all other distance classes.

If there are explanatory variables (RDA, CCA, pRDA, pCCA) and a significance test for residual autocorrelation was performed when running the function `mso`, the function `plot.mso` will print an estimate of how much the autocorrelation (based on significant distance classes) causes the global error variance of the regression analysis to be underestimated.

Value

The function `mso` returns an amended `cca` or `rda` object with the additional attributes `grain`, `H`, `H.test`, and `vario`.

- **grain**: The grain attribute defines the interval size of the distance classes.
- **H**: `H` is an object of class `’dist’` and contains the geographic distances between observations.
- **H.test**: `H.test` contains a set of dummy variables that describe which pairs of observations (rows = elements of `object$H`) fall in which distance class (columns).
- **vario**: The vario attribute is a data frame that contains some or all of the following components for the rda case (cca case in brackets):
  - `H`: Distance class as multiples of grain.
  - `Dist`: Average distance of pairs of observations in distance class `H`.
  - `n`: Number of unique pairs of observations in distance class `H`.
  - `All`: Empirical (chi-square) variogram of total variance (inertia).
  - `Sum`: Sum of empirical (chi-square) variograms of explained and residual variance (inertia).
  - `CA`: Empirical (chi-square) variogram of residual variance (inertia).
  - `CCA`: Empirical (chi-square) variogram of explained variance (inertia).
  - `pCCA`: Empirical (chi-square) variogram of conditioned variance (inertia).
  - `se`: Standard error of the empirical (chi-square) variogram of total variance (inertia).
  - `CA.signif`: P-value of permutation test for spatial independence of residual variance (inertia).

Note

The function is based on the code published in the Ecological Archives E085-006 (http://www.esapubs.org/archive/ecol/E085/006/default.htm).

Author(s)

The responsible author was Helene Wagner.
References


See Also

Function `cca` and `rda.cca.object`.

Examples

```r
## Reconstruct worked example of Wagner (submitted):
X <- matrix(c(1, 2, 3, 2, 1, 0), 3, 2)
Y <- c(3, -1, -2)
tmat <- c(1:3)
## Canonical correspondence analysis (cca):
Example.cca <- cca(X, Y)
Example.cca <- mso(Example.cca, tmat)
msoplot(Example.cca)
Example.cca$vario

## Correspondence analysis (ca):
Example.ca <- mso(cca(X), tmat)
msoplot(Example.ca)

## Unconstrained ordination with test for autocorrelation
## using oribatid mite data set as in Wagner (2004)
data(mite)
data(mite.env)
data(mite.xy)

mite.cca <- cca(log(mite + 1))
mite.cca <- mso(mite.cca, mite.xy, grain = 1, permutations = 100)
msoplot(mite.cca)
mite.cca

## Constrained ordination with test for residual autocorrelation
## and scale-invariance of species-environment relationships
mite.cca <- cca(log(mite + 1) ~ SubsDens + WatrCont + Substrate + Shrub + Topo, mite.env)
mite.cca <- mso(mite.cca, mite.xy, permutations = 100)
msoplot(mite.cca)
mite.cca
```

### Multiplicative Diversity Partitioning

**Description**

In multiplicative diversity partitioning, mean values of alpha diversity at lower levels of a sampling hierarchy are compared to the total diversity in the entire data set or the pooled samples (gamma diversity).
Usage

multipart(formula, data, index=c("renyi", "tsallis"), scales = 1,
          global = FALSE, relative = FALSE, nsimul=99, ...)

## S3 method for class 'multipart'

print(x, ...)

Arguments

formula A two sided model formula in the form \( y \sim x \), where \( y \) is the community data
matrix with samples as rows and species as column. Right hand side (\( x \)) must
contain factors referring to levels of sampling hierarchy, terms from right to left
will be treated as nested (first column is the lowest, last is the highest level).
These variables must be factors in order to unambiguous handling. Interaction
terms are not allowed.

data A data frame where to look for variables defined in the right hand side of
formula. If missing, variables are looked in the global environment.

index Character, the entropy index to be calculated (see Details).

relative Logical, if TRUE then beta diversity is standardized by its maximum (see De-
tails).

scales Numeric, of length 1, the order of the generalized diversity index to be used.

global Logical, indicates the calculation of beta diversity values, see Details.

nsimul Number of permutation to use if matr is not of class 'permat'. If nsimul =
0, only the FUN argument is evaluated. It is thus possible to reuse the statistic
values without using a null model.

x An object to print.

... Other arguments passed to oecosimu, i.e. method, thin or burnin.

Details

Multiplicative diversity partitioning is based on Whittaker’s (1972) ideas, that has recently been
generalised to one parametric diversity families (i.e. Rényi and Tsallis) by Jost (2006, 2007). Jost
recommends to use the numbers equivalents (Hill numbers), instead of pure diversities, and proofs,
that this satisfies the multiplicative partitioning requirements.

The current implementation of multipart calculates Hill numbers based on the functions renyi
and tsallis (provided as index argument). If values for more than one scales are desired, it
should be done in separate runs, because it adds extra dimensionality to the implementation, which
has not been resolved efficiently.

Alpha diversities are then the averages of these Hill numbers for each hierarchy levels, the global
gamma diversity is the alpha value calculated for the highest hierarchy level. When global =
TRUE, beta is calculated relative to the global gamma value:

\[ \beta_i = \gamma / \alpha_i \]

when global = FALSE, beta is calculated relative to local gamma values (local gamma means
the diversity calculated for a particular cluster based on the pooled abundance vector):

\[ \beta_{ij} = \alpha_{(i+1)j} / \text{mean}(\alpha_{ij}) \]
where \( j \) is a particular cluster at hierarchy level \( i \). Then beta diversity value for level \( i \) is the mean of the beta values of the clusters at that level, \( \beta_i = \text{mean}(\beta_{ij}) \).

If \( \text{relative} = \text{TRUE} \), the respective beta diversity values are standardized by their maximum possible values (\( \text{mean}(\beta_{ij})/\beta_{\text{max},ij} \)) given as \( \beta_{\text{max},ij} = n_j \) (the number of lower level units in a given cluster \( j \)).

The expected diversity components are calculated \( \text{nsimul} \) times by individual based randomisation of the community data matrix. This is done by the "r2dtable" method in oecosimu by default.

Value

An object of class 'multipart' with same structure as 'oecosimu' objects.

Author(s)

Péter Sólymos, <solymos@ualberta.ca>

References


See Also

See adipart for additive diversity partitioning, hiersimu for hierarchical null model testing and oecosimu for permutation settings and calculating \( p \)-values.

Examples

data(mite)
data(mite.xy)
data(mite.env)
## Function to get equal area partitions of the mite data
cutter <- function (x, cut = seq(0, 10, by = 2.5)) {
  out <- rep(1, length(x))
  for (i in 2:(length(cut) - 1))
    out[which(x > cut[i] & x <= cut[i + 1])] <- i
  return(as.factor(out))
}
## The hierarchy of sample aggregation
levsm <- data.frame(  
  l1=as.factor(1:nrow(mite)),
  l2=cutter(mite.xy$y, cut = seq(0, 10, by = 2.5)),
  l3=cutter(mite.xy$y, cut = seq(0, 10, by = 5)),
  l4=cutter(mite.xy$y, cut = seq(0, 10, by = 10)))
## Multiplicative diversity partitioning
multipart(mite ~ ., levsm, index="renyi", scales=1, nsimul=25)
multipart(mite ~ ., levsm, index="renyi", scales=1, nsimul=25, relative=TRUE)
multipart(mite ~ ., levsm, index="renyi", scales=1, nsimul=25, global=TRUE)
Description

Patches or local communities are regarded as nested if they all could be subsets of the same community. In general, species poor communities should be subsets of species rich communities, and rare species should only occur in species rich communities.

Usage

```r
nestedchecker(comm)
nestedn0(comm)
nesteddisc(comm)
nestedtemp(comm, ...)
nestednoidf(comm, order = TRUE, weighted = FALSE)
## S3 method for class 'nestedtemp'
plot(x, kind = c("temperature", "incidence"),
     col=rev(heat.colors(100)), names = FALSE, ...)
```

Arguments

- `comm` Community data.
- `x` Result object for a plot.
- `col` Colour scheme for matrix temperatures.
- `kind` The kind of plot produced.
- `names` Label columns and rows in the plot using names in `comm`. If it is a logical vector of length 2, row and column labels are returned accordingly.
- `order` Order rows and columns by frequencies.
- `weighted` Use species abundances as weights of interactions.
- `...` Other arguments to functions.

Details

The nestedness functions evaluate alternative indices of nestedness. The functions are intended to be used together with Null model communities and used as an argument in `oecosimu` to analyse the non-randomness of results.

Function `nestedchecker` gives the number of checkerboard units, or 2x2 submatrices where both species occur once but on different sites (Stone & Roberts 1990).

Function `nestedn0` implements nestedness measure N0 which is the number of absences from the sites which are richer than the most pauperate site species occurs (Patterson & Atmar 1986).

Function `nesteddisc` implements discrepancy index which is the number of ones that should be shifted to fill a row with ones in a table arranged by species frequencies (Brualdi & Sanderson...
The original definition arranges species (columns) by their frequencies, but did not have any method of handling tied frequencies.

The `nesteddisc` function tries to order tied columns to minimize the discrepancy statistic but this is rather slow, and with a large number of tied columns there is no guarantee that the best ordering was found. In that case a warning of tied columns will be issued.

Function `nestedtemp` finds the matrix temperature which is defined as the sum of “surprises” in arranged matrix. In arranged unsurprising matrix all species within proportion given by matrix fill are in the upper left corner of the matrix, and the surprise of the absence or presences is the diagonal distance from the fill line (Atmar & Patterson 1993). Function tries to pack species and sites to a low temperature (Rodríguez-Gironés & Santamaria 2006), but this is an iterative procedure, and the temperatures usually vary among runs. Function `nestedtemp` also has a `plot` method which can display either incidences or temperatures of the surprises. Matrix temperature was rather vaguely described (Atmar & Patterson 1993), but Rodríguez-Gironés & Santamaria (2006) are more explicit and their description is used here. However, the results probably differ from other implementations, and users should be cautious in interpreting the results. The details of calculations are explained in the `vignette` Design decisions and implementation that you can read using functions `vignette` or `vegandocs`. Function `nestedness` in the `bipartite` package is a direct port of the BINMATNEST programme of Rodríguez-Gironés & Santamaria (2006).

Function `nestednodf` implements a nestedness metric based on overlap and decreasing fill (Almeida-Neto et al., 2008). Two basic properties are required for a matrix to have the maximum degree of nestedness according to this metric: (1) complete overlap of 1’s from right to left columns and from down to up rows, and (2) decreasing marginal totals between all pairs of columns and all pairs of rows. The nestedness statistic is evaluated separately for columns (`N columns`) for rows (`N rows`) and combined for the whole matrix (`NODF`). If you set `order = FALSE`, the statistic is evaluated with the current matrix ordering allowing tests of other meaningful hypothesis of matrix structure than default ordering by row and column totals (breaking ties by total abundances when `weighted = TRUE`) (see Almeida-Neto et al. 2008). With `weighted = TRUE`, the function finds the weighted version of the index (Almeida-Neto & Ulrich, 2011). However, this requires quantitative null models for adequate testing.

**Value**

The result returned by a nestedness function contains an item called `statistic`, but the other components differ among functions. The functions are constructed so that they can be handled by `oecosimu`.

**Author(s)**

Jari Oksanen and Gustavo Carvalho (nestednodf).

**References**


See Also

In general, the functions should be used with `oecosimu` which generates Null model communities to assess the non-randomness of nestedness patterns.

Examples

```r
data(sipoo)
## Matrix temperature
out <- nestedtemp(sipoo)
out
plot(out)
plot(out, kind="incid")
## Use oecosimu to assess the non-randomness of checker board units
nestedchecker(sipoo)
oecosimu(sipoo, nestedchecker, "quasiswap")
## Another Null model and standardized checkerboard score
oecosimu(sipoo, nestedchecker, "r00", statistic = "C.score")
```

---

**oecosimu**  
*Null Models for Biological Communities*

**Description**

Null models generate random communities with different criteria to study the significance of nestedness or other community patterns. The function only simulates binary (presence/absence) models with constraint for total number of presences, and optionally for numbers of species and/or species frequencies.
Usage

```r
oecosimu(comm, nestfun, method, nsimul = 99, burnin = 0, thin = 1,
          statistic = "statistic", alternative = c("two.sided", "less", "greater"),
          ...)
commsimulator(x, method, thin=1)
## S3 method for class 'oecosimu'
as.ts(x, ...)
## S3 method for class 'oecosimu'
as.mcmc(x)
## S3 method for class 'oecosimu'
density(x, ...)
## S3 method for class 'oecosimu'
densityplot(x, data, xlab = "Simulated", ...)
```

Arguments

- **comm**: Community data.
- **x**: Community data for `commsimulator`, or an `oecosimu` result object for `as.ts`, `as.mcmc`, `density` and `densityplot`.
- **nestfun**: Function to analyse nestedness. Some functions are provided in `vegan`, but any function can be used if it accepts the community as the first argument, and returns either a plain number or the result in list item with the name defined in argument `statistic`. See Examples for defining your own functions.
- **method**: Null model method. See details.
- **nsimul**: Number of simulated null communities.
- **burnin**: Number of null communities discarded before proper analysis in sequential methods "swap" and "tswap".
- **thin**: Number of discarded null communities between two evaluations of nestedness statistic in sequential methods "swap" and "tswap".
- **statistic**: The name of the statistic returned by `nestfun`.
- **alternative**: a character string specifying the alternative hypothesis, must be one of "two.sided" (default), "greater" or "less". You can specify just the initial letter.
- **data**: Ignored argument of the generic function.
- **xlab**: Label of the x-axis.
- **...**: Other arguments to functions.

Details

Function `oecosimu` is a wrapper that evaluates a nestedness statistic using function given by `nestfun`, and then simulates a series of null models using `commsimulator` or other functions (depending on method argument), and evaluates the statistic on these null models. The `vegan` packages contains some nestedness functions that are described separately (`nestedchecker`, `nesteddisc`, `nestedn0`, `nestedtemp`), but many other functions can be used as long as they are meaningful with binary or quantitative community models. An applicable function must return either the statistic as a plain number, or as a list element "statistic" (like `chisq.test`), or
in an item whose name is given in the argument statistic. The statistic can be a single number
(like typical for a nestedness index), or it can be a vector. The vector indices can be used to analyse
site (row) or species (column) properties, see treedive for an example.

Function commsimulator implements binary (presence/absence) null models for community
composition. The implemented models are r00 which maintains the number of presences but fills
these anywhere so that neither species (column) nor site (row) totals are preserved. Methods r0, r1
and r2 maintain the site (row) frequencies. Method r0 fills presences anywhere on the row with no
respect to species (column) frequencies, r1 uses column marginal frequencies as probabilities, and
r2 uses squared column sums. Methods r1 and r2 try to simulate original species frequencies, but
they are not strictly constrained. All these methods are reviewed by Wright et al. (1998). Method
c0 maintains species frequencies, but does not honour site (row) frequencies (Jonsson 2001).

The other methods maintain both row and column frequencies. Methods swap and tswap implement
sequential methods, where the matrix is changed only little in one step, but the changed matrix
is used as an input if the next step. Methods swap and tswap inspect random 2x2 submatrices
and if they are checkerboard units, the order of columns is swapped. This changes the matrix struc-
ture, but does not influence marginal sums (Gotelli & Entsminger 2003). Method swap inspects
submatrices so long that a swap can be done. Miklós & Podani (2004) suggest that this may lead
into biased sequences, since some columns or rows may be more easily swapped, and they suggest
trying a fixed number of times and doing zero to many swaps at one step. This method is imple-
mented by method tswap or trial swap. Function commsimulator makes only one trial swap in
time (which probably does nothing), but oecosimu estimates how many submatrices are expected
before finding a swappable checkerboard, and uses that ratio to thin the results, so that on average
one swap will be found per step of tswap. However, the checkerboard frequency probably changes
during swaps, but this is not taken into account in estimating the thin. One swap still changes the
matrix only little, and it may be useful to thin the results so that the statistic is only evaluated after
burnin steps (and thinned).

Methods quasiswap and backtracking are not sequential, but each call produces a matrix
that is independent of previous matrices, and has the same marginal totals as the original data.
The recommended method is quasiswap which is much faster because it is implemented in C.
Method backtracking is provided for comparison, but it is so slow that it may be dropped
from future releases of vegan (or also implemented in C). Method quasiswap (Miklós & Podani
2004) implements a method where matrix is first filled honouring row and column totals, but with
integers that may be larger than one. Then the method inspects random 2x2 matrices and performs
a quasiswap on them. Quasiswap is similar to ordinary swap, but it also can reduce numbers above
one to ones maintaining marginal totals. Method backtracking implements a filling method
with constraints both for row and column frequencies (Gotelli & Entsminger 2001). The matrix is
first filled randomly using row and column frequencies as probabilities. Typically row and column
sums are reached before all incidences are filled in. After that begins “backtracking”, where some
of the points are removed, and then filling is started again, and this backtracking is done so may
times that all incidences will be filled into matrix. The quasiswap method is not sequential, but
it produces a random incidence matrix with given marginal totals.

Function as.ts transforms the simulated results of sequential methods into a time series or a ts
object. This allows using analytic tools for time series in studying the sequences (see examples).
Function as.mcmc transforms the simulated results of sequential methods into an mcmc object of
the coda package. The coda package provides functions for the analysis of stationarity, adequacy of
sample size, autocorrelation, need of burn-in and much more for sequential methods. Please consult
the documentation of coda package.
Function `density` provides an interface to the standard `density` function for the simulated values. Function `densityplot` is an interface to the `densityplot` function of the `lattice` package. The `density` can be used meaningfully only for single statistics and must be plotted separately. The `densityplot` function can handle multiple statistics, and it plots the results directly. In addition to the density, the `densityplot` also shows the observed value of the statistic (provided it is within the graph limits). The `densityplot` function is defined as a generic function in the `lattice` package and you must either load the `lattice` library before calling `densityplot`, or use the longer form `densityplot.oecosimu` when you first time call the function.

As a result of `method = "r2dtable"` in `oecosimu`, quantitative community null models are used to evaluate the statistic. This setting uses the `r2dtable` function to generate random matrices with fixed row and column totals (hypergeometric distribution). This null model is used in diversity partitioning function (see `adipart`).

The `method` argument can be a function with first argument taking the community matrix, and optionally with `burnin` and `thin` argument. The function must return a matrix-like object with same dimensions. But be careful, blindly applying permuted matrices for null model testing can be dangerous.

Value

Function `oecosimu` returns the result of `nestfun` with one added component called `oecosimu`. The `oecosimu` component contains the simulated values of the statistic (item `simulated`), the name of the method, two-sided $P$ value and z-value of the statistic based on simulation. The `commsimulator` returns a null model matrix or a swap of the input matrix.

Note

Functions `commsimulator` and `oecosimu` do not have default `nestfun` nor default `method`, because there is no clear natural choice. If you use these methods, you must be able to choose your own strategy. The choice of nestedness index is difficult because the functions seem to imply very different concepts of structure and randomness. The choice of swapping method is also problematic. Method `r00` has some heuristic value of being really random. However, it produces null models which are different from observed communities in most respects, and a “significant” result may simply mean that not all species are equally common (`r0` is similar with this respect). It is also difficult to find justification for `r2`. The methods maintaining both row and column totals only study the community relations, but they can be very slow. Moreover, they regard marginal totals as constraints instead of results of occurrence patterns. You should evaluate timings in small trials (one cycle) before launching an extensive simulation. One swap is fast, but it changes data only little, and you may need long `burnin` and strong `thinning` in large matrices. You should plot the simulated values to see that they are more or less stationary and there is no trend. Method `quasiswap` is implemented in C and it is much faster than `backtrack`. Method `backtrack` may be removed from later releases of `vegan` because it is slow, but it is still included for comparison.

If you wonder about the name of `oecosimu`, look at journal names in the References (and more in `nestedtemp`).

Author(s)

Jari Oksanen
### References


### See Also

- `r2dtable` generates table with given marginals but with entries above one. Functions `permatfull` and `permatswap` generate Null models for count data. Function `rndtaxa` (*labdsv* package) randomizes a community table. See also `nestedtemp` (that also discusses other nestedness functions) and `treedive` for another application.

### Examples

```r
## Use the first eigenvalue of correspondence analysis as an index
## of structure: a model for making your own functions.
data(sipoo)
out <- oecosimu(sipoo, decorana, "swap", burnin=100, thin=10, statistic="evals")
out
## Inspect the swap sequence as a time series object
plot(as.ts(out))
laq.plot(as.ts(out))
acf(as.ts(out))
## Density plot: needs lattice
require(lattice)
densityplot(out, as.table = TRUE)
## Use quantitative null models to compare
## mean Bray-Curtis dissimilarities
data(dune)
meandist <- function(x) mean(vegdist(x, "bray"))
mbcl <- oecosimu(dune, meandist, "r2dtable")
mbcl
## Define a custom function that shuffles
## cells in each rows
f <- function(x) {
  apply(x, 2, function(z) sample(z, length(z)))
}
mbc2 <- oecosimu(as.matrix(dune), meandist, f)
mbc2
```
Description

Functions to add convex hulls, arrows, line segments, regular grids of points, ‘spider’ graphs, ellipses or cluster dendrogram to ordination diagrams. The ordination diagrams can be produced by vegan plot.cca, plot.decorana or ordiplot.

Usage

```r
ordihull(ord, groups, display = "sites", draw = c("lines","polygon", "none"),
          show.groups, label = FALSE, ...)
ordiellipse(ord, groups, display="sites", kind = c("sd","se"), conf,
            draw = c("lines","polygon", "none"), w = weights(ord, display),
            col = NULL, show.groups, label = FALSE, ...)
ordispider(ord, groups, display="sites", w = weights(ord, display),
           show.groups, label = FALSE, ...)
ordiarrows(ord, groups, levels, replicates, display = "sites",
           show.groups, startmark, label = FALSE, ...)
ordisegments(ord, groups, levels, replicates, display = "sites",
             show.groups, label = FALSE, ...)
ordigrid(ord, levels, replicates, display = "sites", lty = c(1,1),
            col = c(1,1), lwd = c(1,1), ...)
ordicluster(ord, cluster, prune = 0, display = "sites",
             w = weights(ord, display), ...)
```

Arguments

- **ord**: An ordination object or an `ordiplot` object.
- **groups**: Factor giving the groups for which the graphical item is drawn.
- **levels, replicates**: Alternatively, regular groups can be defined with arguments `levels` and `replicates`, where `levels` gives the number of groups, and `replicates` the number of successive items at the same group.
- **display**: Item to displayed.
- **draw**: Use either `lines` or `polygon` to draw the line. Graphical parameters are passed to both. The main difference is that polygons may be filled and non-transparent. With `none` nothing is drawn, but the function returns the `invisible` plotting data.
- **show.groups**: Show only given groups. This can be a vector, or `TRUE` if you want to show items for which condition is `TRUE`. This argument makes it possible to use different colours and line types for groups. The default is to show all groups.
- **label**: Label the groups by their names. In `ordiellipse, ordihull and ordispider` the the group name is in the centroid of the object, in `ordiarrows in the`
ordihull

start of the arrow, and in ordisegments at both ends. ordiellipse and ordihull use standard text, and others use ordilabel.

startmark plotting character used to mark the first item. The default is to use no mark, and for instance, startmark = 1 will draw a circle. For other plotting characters, see pch in points.

w Weights used to find the average within group. Weights are used automatically for cca and decorana results, unless undone by the user. w=NULL sets equal weights to all points.

kind Whether standard deviations of points (sd) or standard deviations of their (weighted) averages (se) are used.

conf Confidence limit for ellipses, e.g. 0.95. If given, the corresponding sd or se is multiplied with the corresponding value found from the Chi-squared distribution with 2df.

cluster Result of hierarchic cluster analysis, such as hclust or agnes.

col Colour of ellipses or ellipse fills in ordiellipse or lines in ordigrid. For other functions the effect depends on the underlining functions this argument is passed to.

lty, lwd Line type, line width used for levels and replicates in ordigrid.

prune Number of upper level hierarchies removed from the dendrogram. If prune > 0, dendrogram will be disconnected.

... Parameters passed to graphical functions such as lines, segments, arrows, polygon or to scores to select axes and scaling etc.

Details

Function ordihull draws lines or polygons for the convex hulls found by function chull encircling the items in the groups.

Function ordiellipse draws lines or polygons for dispersion ellipses using either standard deviation of point scores or standard error of the (weighted) average of scores, and the (weighted) correlation defines the direction of the principal axis of the ellipse. An ellipsoid hull can be drawn with function ellipsoidhull of package cluster.

Functions ordihull and ordiellipse return the invisible plotting structure. In ordihull this is a list of coordinates of the hulls (which can be extracted with scores), and in ordiellipse a list of covariance matrices and scales used in drawing the ellipses. These result objects have a summary method that returns the coordinates of the centres of the ellipses or hulls and their surface areas in user units. The centres of the hulls may differ from the location of the label which is the centre of the points instead of the centre of the polygon. With draw = "none" only the result object is returned and nothing is drawn.

Function ordiarrows draws arrows and ordisegments draws line segments between successive items in the groups. Function ordigrid draws line segments both within the groups and for the corresponding items among the groups.

Function ordispider draws a ‘spider’ diagram where each point is connected to the group centroid with segments. Weighted centroids are used in the correspondence analysis methods cca and decorana or if the user gives the weights in the call. If ordispider is called with cca or
**rda** result without **groups** argument, the function connects each ‘WA’ scores to the corresponding ‘LC’ score. If the argument is a (**invisible**) ordihull object, the function will connect the points of the hull to their centroid.

Function **ordicluster** overlays a cluster dendrogram onto ordination. It needs the result from a hierarchic clustering such as **hclust** or **agnes**, or other with a similar structure. Function **ordicluster** connects cluster centroids to each other with line **segments**. Function uses centroids of all points in the clusters, and is therefore similar to average linkage methods.

**Note**

These functions add graphical items to ordination graph: You must draw a graph first.

**Author(s)**

Jari Oksanen

**See Also**

The functions pass parameters to basic graphical functions, and you may wish to change the default values in **arrows**, **lines**, **segments** and **polygon**. You can pass parameters to **scores** as well. Underlying function for **ordihull** is **chull**.

**Examples**

```r
data(dune)
data(dune.env)
mod <- cca(dune ~ Management, dune.env)
attach(dune.env)
## pass non-graphical arguments without warnings
plot(mod, type="n", scaling = 3)
## Catch the invisible result of ordihull...
pl <- ordihull(mod, Management, scaling = 3, label = TRUE)
## ... and find centres and areas of the hulls
summary(pl)
## use ordispider to label and mark the hull
plot(mod, type = "n")
pl <- ordihull(mod, Management, scaling = 3)
ordispider(pl, col="red", lty=3, label = TRUE)
## ordispider to connect WA and LC scores
plot(mod, dis=c("wa","lc"), type="p")
ordispider(mod)
## Other types of plots
plot(mod, type = "p", display="sites")
ordicluster(mod, hclust(vegdist(dune)), prune=3, col = "blue")
plot(mod, type="n", display = "sites")
text(mod, display="sites", labels = as.character(Management))
pl <- ordiellipse(mod, Management, kind="se", conf=0.95, lwd=2, col="blue")
summary(pl)
```
Function `ordilabel` is similar to `text`, but the text is on an opaque label. This can help in crowded ordination plots: you still cannot see all text labels, but at least the uppermost are readable. Argument `priority` helps to make the most important labels visible.

**Usage**

`ordilabel(x, display, labels, choices = c(1, 2), priority, cex = 0.8, fill = "white", border = NULL, col = NULL, xpd = TRUE, ...)`

**Arguments**

- **x**: An ordination object an any object known to `scores`.
- **display**: Kind of scores displayed (passed to `scores`).
- **labels**: Optional text used in plots. If this is not given, the text is found from the ordination object.
- **choices**: Axes shown (passed to `scores`).
- **priority**: Vector of the same length as the number of labels. The items with high priority will be plotted uppermost.
- **cex**: Character expansion for the text (passed to `text`).
- **fill**: Background colour of the labels (the `col` argument of `polygon`).
- **border**: The colour and visibility of the border of the label as defined in `polygon`.
- **col**: Text colour. Default NULL will give the value of `border` or `par("fg")` if `border` is NULL.
- **xpd**: Draw labels also outside the plot region (see `par`).
- **...**: Other arguments (passed to `text`).

**Details**

The function may be useful with crowded ordination plots, in particular together with argument `priority`. You will not see all text labels, but at least some are readable. Other alternatives to crowded plots are `identify.ordiplot`, `orditorp` and `orditkplot`.

**Author(s)**

Jari Oksanen

**See Also**

`scores`, `polygon`, `text`. The function is modelled after `s.label` in `ade4` package.
Examples

data(dune)
ord <- cca(dune)
plot(ord, type = "n")
ordilabel(ord, dis="sites", cex=1.2, font=3, fill="hotpink", col="blue")
## You may prefer separate plots, but here species as well
ordilabel(ord, dis="sp", font=2, priority=colSums(dune))

ordiplot

Alternative plot and identify Functions for Ordination

Description

Ordination plot function especially for congested plots. Function ordiplot always plots only unlabelled points, but identify.ordiplot can be used to add labels to selected site, species or constraint points. Function identify.ordiplot can be used to identify points from plot.cca, plot.decorana, plot.procrustes and plot.rad as well.

Usage

ordiplot(ord, choices = c(1, 2), type="points", display, xlim, ylim, ...)
## S3 method for class 'ordiplot'
identify(x, what, labels, ...)
## S3 method for class 'ordiplot'
points(x, what, select, ...)
## S3 method for class 'ordiplot'
text(x, what, labels, select, ...)

Arguments

ord A result from an ordination.
choices Axes shown.
type The type of graph which may be "points", "text" or "none" for any ordination method.
display Display only "sites" or "species". The default for most methods is to display both, but for cca, rda and capscale it is the same as in plot.cca.
xlim, ylim the x and y limits (min,max) of the plot.
... Other graphical parameters.
x A result object from ordiplot.
what Items identified in the ordination plot. The types depend on the kind of plot used. Most methods know sites and species, functions cca and rda know in addition constraints (for 'LC' scores), centroids and biplot, and plot.procrustes ordination plot has heads and points.
labels Optional text used for labels. Row names will be used if this is missing.
select Items to be displayed. This can either be a logical vector which is TRUE for displayed items or a vector of indices of displayed items.
Function `ordiplot` draws an ordination diagram using black circles for sites and red crosses for species. It returns invisibly an object of class `ordiplot` which can be used by `identify.ordiplot` to label selected sites or species, or constraints in `cca` and `rda`.

The function can handle output from several alternative ordination methods. For `cca`, `rda` and `decorana` it uses their `plot` method with option `type = "points"`. In addition, the `plot` functions of these methods return invisibly an `ordiplot` object which can be used by `identify.ordiplot` to label points. For other ordinations it relies on `scores` to extract the scores.

For full user control of plots, it is best to call `ordiplot` with `type = "none"` and save the result, and then add sites and species using `points.ordiplot` or `text.ordiplot` which both pass all their arguments to the corresponding default graphical functions.

Function `ordiplot` returns invisibly an object of class `ordiplot` with items `sites`, `species` and `constraints` (if these are available in the ordination object). Function `identify.ordiplot` uses this object to label the point.

The purpose of these functions is to provide similar functionality as the `plot`, `plotid` and `specid` methods in library `labdsv`. The functions are somewhat limited in parametrization, but you can call directly the standard `identify` and `plot` functions for a better user control.

Author(s)

Jari Oksanen

See Also

`identify` for basic operations, `plot.cca`, `plot.decorana`, `plot.procrustes` which also produce objects for `identify.ordiplot` and `scores` for extracting scores from non-vegan ordinations.

Examples

```r
# Draw a cute NMDS plot from a non-vegan ordination (isoMDS).
# Function `metaMDS` would be an easier alternative.
data(dune)
dune.dis <- vegdist(wisconsin(dune))
library(MASS)
dune.mds <- isoMDS(dune.dis)
dune.mds <- postMDS(dune.mds, dune.dis)
dune.mds$species <- wascores(dune.mds$points, dune, expand = TRUE)
fig <- ordiplot(dune.mds, type = "none")
points(fig, "sites", pch=21, col="red", bg="yellow")
text(fig, "species", col="blue", cex=0.9)
```

```r
# Default plot of the previous using `identify` to label selected points
## Not run:
fig <- ordiplot(dune.mds)
```
identify(fig, "spec")
## End(Not run)

description

Function ordiplot3d displays three-dimensional ordination graphics using `scatterplot3d`. Function ordirgl displays three-dimensional dynamic ordination graphs which can be rotated and zoomed into using rgl package. Both work with all ordination results form vegan and all ordination results known by scores function.

usage

ordiplot3d(object, display = "sites", choices = 1:3, ax.col = 2, 
arr.len = 0.1, arr.col = 4, envfit, xlab, ylab, zlab, ...)
ordirgl(object, display = "sites", choices = 1:3, type = "p", 
ax.col = "red", arr.col = "yellow", text, envfit, ...)
orglpoints(object, display = "sites", choices = 1:3, ...)
orgltext(object, text, display = "sites", choices = 1:3, justify = "center", 
adj = 0.5, ...)
orglsegments(object, groups, display = "sites", choices = 1:3, ...)
orglspider(object, groups, display = "sites", w = weights(object, display), 
choices = 1:3, ...)

arguments

object
  An ordination result or any object known by scores.
display
  Display "sites" or "species" or other ordination object recognized by scores.
choices
  Selected three axes.
arr.len
  'Length' (width) of arrow head passed to arrows function.
arr.col
  Colour of biplot arrows and centroids of environmental variables.
type
  The type of plots: "p" for points or "t" for text labels.
ax.col
  Axis colour (concerns only the crossed axes through the origin).
text
  Text to override the default with type = "t".
envfit
  Fitted environmental variables from envfit displayed in the graph.
xlab, ylab, zlab
  Axis labels passed to scatterplot3d. If missing, labels are taken from the ordination result. Set to NA to suppress labels.
justify, adj
  Text justification passed to rgl.texts. One of these is used depending on the version of rgl installed.
groups  Factor giving the groups for which the graphical item is drawn.

w  Weights used to find the average within group. Weights are used automatically for cca and decorana results, unless undone by the user. w=NULL sets equal weights to all points.

...  Other parameters passed to graphical functions.

Details

Both function display three-dimensional ordination graphics. Function ordiplot3d plots static scatter diagrams using scatterplot3d. Function ordirgl plots dynamic graphics using OpenGL in rgl. Both functions use most default settings of underlying graphical functions, and you must consult their help pages to change graphics to suit your taste (see scatterplot3d, rgl, rgl.points, rgl.texts). Both functions will display only one selected set of scores, typically either "sites" or "species", but for instance cca also has "lc" scores. In constrained ordination (cca, rda, capscale), biplot arrows and centroids are always displayed similarly as in two-dimensional plotting function plot.cca. Alternatively, it is possible to display fitted environmental vectors or class centroids from envfit in both graphs. These are displayed similarly as the results of constrained ordination, and they can be shown only for non-constrained ordination. The user must remember to specify at least three axes in envfit if the results are used with these functions.

Function ordiplot3d plots only points. However, it returns invisibly an object inheriting from ordiplot so that you can use identify.ordiplot to identify "points" or "arrows". The underlying scatterplot3d function accepts type = "n" so that only the axes, biplot arrows and centroids of environmental variables will be plotted, and the ordination scores can be added with text.ordiplot or points.ordiplot. Further, you can use any functions from the ordihull family with the invisible result of ordiplot3d, but you must remember to specify the display as "points" or "arrows". To change the viewing angle, orientation etc. you must see scatterplot3d.

Function ordirgl makes a dynamic three-dimensional graph that can be rotated with mouse, and zoomed into with mouse buttons or wheel (but Mac users with one-button mouse should see rgl.viewpoint), or try ctrl-button. MacOS X users must start X11 before calling rgl commands. Function ordirgl uses default settings, and you should consult the underlying functions rgl.points, rgl.texts to see how to control the graphics. Function ordirgl always cleans its graphic window before drawing. Functions orglpoints adds points and orgltext adds text to existing ordirgl windows. In addition, function orglsegments combines points within "groups" with line segments similarly as ordisegments. Function orglspider works similarly as ordispider: it connects points to their weighted centroid within "groups", and in constrained ordination it can connect "wa" or weighted averages scores to corresponding "lc" or linear combination scores if "groups" is missing. In addition, basic rgl functions rgl.points, rgl.texts, rgl.lines and many others can be used.

Value

Function ordiplot3d returns invisibly an object of class "ordiplot3d" inheriting from ordiplot. The return object will contain the coordinates projected onto two dimensions for "points", and possibly for the heads of "arrows" and "centroids" of environmental variables. Functions like identify.ordiplot, points.ordiplot, text.ordiplot can use this result, as well as ordihull and other functions documented with the latter. In addition, the result will
contain the object returned by `scatterplot3d`, including function `xyz.converter` which projects three-dimensional coordinates onto the plane used in the current plot. Function `ordirgl` returns nothing.

**Warning**

Function `ordirgl` uses OpenGL package `rgl` which may not be functional in all platforms, and can crash R in some: use `save.image` before trying `ordirgl`. Mac users must start `X11` (and first install `X11` and some other libraries) before being able to use `rgl`. It seems that `rgl.texts` does not always position the text like supposed, and it may be safe to verify text location with corresponding points.

**Note**

The user interface of `rgl` changed in version 0.65, but the `ordirgl` functions do not yet fully use the new capabilities. However, they should work both in old and new versions of `rgl`.

**Author(s)**

Jari Oksanen

**See Also**

`scatterplot3d`, `rgl`, `rgl.points`, `rgl.texts`, `rgl.viewpoint`, `ordiplot`, `identify.ordiplot`, `text.ordiplot`, `points.ordiplot`, `ordihull`, `plot.cca`, `envfit`.

**Examples**

```r
## Examples are not run, because they need non-standard packages
## 'scatterplot3d' and 'rgl' (and the latter needs user interaction).

### Default 'ordiplot3d'
## Not run:
data(dune)
data(dune.env)
ord <- cca(dune ~ A1 + Moisture, dune.env)
ordiplot3d(ord)

### A boxed 'pin' version
ordiplot3d(ord, type = "h")

### More user control
pl <- ordiplot3d(ord, angle=15, type="n")
points(pl, "points", pch=16, col="red", cex = 0.7)

### identify(pl, "arrows", col="blue") would put labels in better positions
text(pl, "arrows", col="blue", pos=3)
text(pl, "centroids", col="blue", pos=1, cex = 1.2)

### ordirgl
ordirgl(ord, size=2)
ordirgl(ord, display = "species", type = "t")
rgl.quit()
```

## End(Not run)
ordipointlabel  Ordination Plots with Points and Optimized Locations for Text

Description

The function `ordipointlabel` produces ordination plots with points and text label to the points. The points are in the exact location given by the ordination, but the function tries to optimize the location of the text labels to minimize overplotting text. The function may be useful with moderately crowded ordination plots.

Usage

```r
ordipointlabel(x, display = c("sites", "species"), choices = c(1, 2),
    col = c(1, 2), pch = c("o", "+"), font = c(1, 1),
    cex = c(0.8, 0.8), add = FALSE, ...)
```

Arguments

- `x` A result object from ordination.
- `display` Scores displayed in the plot.
- `choices` Axes shown.
- `col, pch, font, cex` Colours, point types, font style and character expansion for each kind of scores displayed in the plot. These should be vectors of the same length as the number of items in `display`.
- `add` Add to an existing plot.
- `...` Other arguments passed to `points` and `text`.

Details

The function uses simulated annealing (`optim`, `method = "SANN"`) to optimize the location of the text labels to the points. There are eight possible locations: up, down, sides and corners. There is a weak preference to text right above the point, and a weak avoidance of corner positions. The exact locations and the goodness of solution varies between runs, and there is no guarantee of finding the global optimum. The optimization can take a long time in difficult cases with a high number of potential overlaps. Several sets of scores can be displayed in one plot.

The function is modelled after `pointLabel` in `maptools` package (which has chained dependencies of S4 packages).

Value

The function returns invisibly an object of class `ordipointlabel` with items `xy` for coordinates of points, `labels` for coordinates of labels, items `pch`, `cex` and `font` for graphical parameters of each point or label. In addition, it returns the result of `optim` as an attribute "optim". The unit of overlap is the area of character "m", and with variable `cex` it is the smallest alternative. The result object inherits from `orditkplot` result, and can be replotted with its `plot` command. It may be...
possible to further edit the result object with `orditkplot`, but for good results it is necessary that the points span the whole horizontal axis without empty margins.

**Note**

The function is designed for ordination graphics, and the optimization works properly with plots of isometric aspect ratio.

**Author(s)**

Jari Oksanen

**References**

See `pointLabel` for references.

**See Also**

`pointLabel` for the model implementation, and `optim` for the optimization.

**Examples**

```r
data(dune)
ord <- cca(dune)
ordipointlabel(ord)
```

---

### ordiresids

*Plots of Residuals and Fitted Values for Constrained Ordination*

**Description**

The function provides `plot.lm` style diagnostic plots for the results of constrained ordination from `cca`, `rda` and `capscale`. Normally you do not need these plots, because ordination is descriptive and does not make assumptions on the distribution of the residuals. However, if you permute residuals in significance tests (`anova.cca`), you may be interested in inspecting that the residuals really are exchangeable and independent of fitted values.

**Usage**

```r
ordiresids(x, kind = c("residuals", "scale", "qqmath"),
           residuals = "working", type = c("p", "smooth", "g"),
           formula, ...)```
Arguments

- **x**: Ordination result from `cca`, `rda` or `capscale`.
- **kind**: The type of plot: "residuals" plot residuals against fitted values, "scale" the square root of absolute residuals against fitted values, and "qqmath" the residuals against expected distribution (defaults `qnorm`), unless defined differently in the `formula` argument.
- **residuals**: The kind of residuals and fitted values. The argument is passed on to `fitted.cca` with alternatives "working" and "response".
- **type**: The type of plot. The argument is passed on to `lattice` functions.
- **formula**: Formula to override the default plot. The formula can contain items `Fitted`, `Residuals`, `Species` and `Sites` (provided that names of species and sites are available in the ordination result).
- **...**: Other arguments passed to `lattice` functions.

Details

The default plots are similar as in `plot.lm`, but they use `Lattice` functions `xyplot` and `qqmath`. The alternatives have default formulae but these can be replaced by the user. The elements available in formula or in the `groups` argument are `Fitted`, `Residuals`, `Species` and `Sites`.

Value

The function return a `Lattice` object that can displayed as plot.

Author(s)

Jari Oksanen

See Also

`plot.lm`, `Lattice`, `xyplot`, `qqmath`.

Examples

```r
data(varespec)
data(varechem)
mod <- cca(varespec ~ A1 + P + K, varechem)
ordiresids(mod)
```
Choose a Model by Permutation Tests in Constrained Ordination

Description

Automatic stepwise model building for constrained ordination methods (cca, rda, capscale). The function ordistep is modelled after step and can do forward, backward and stepwise model selection using permutation tests. Function ordiR2step performs forward model choice solely on adjusted $R^2$ and P-value, for ordination objects created by rda or capscale.

Usage

ordistep(object, scope, direction = c("both", "backward", "forward"), 
Pin = 0.05, Pout = 0.1, pstep = 100, perm.max = 1000, steps = 50, 
trace = TRUE, ...)
ordiR2step(object, scope, direction = c("both", "forward"), 
Pin = 0.05, pstep = 100, perm.max = 1000, 
trace = TRUE, ...)

Arguments

object In ordistep, an ordination object inheriting from cca or rda. In ordiR2step, the object must inherit from rda, that is, it must have been computed using rda or capscale.
scope Defines the range of models examined in the stepwise search. This should be either a single formula, or a list containing components upper and lower, both formulae. See step for details. In ordiR2step, this defines the upper scope; it can also be an ordination object from with the model is extracted.
direction The mode of stepwise search, can be one of "both", "backward", or "forward", with a default of "both". If the scope argument is missing, the default for direction is "backward".
Pin, Pout Limits of permutation $P$-values for adding (Pin) a term to the model, or dropping (Pout) from the model. Term is added if $P \leq \text{Pin}$, and removed if $P > \text{Pout}$.
pstep Number of permutations in one step. See add1.cca.
perm.max Maximum number of permutation in anova.cca.
steps Maximum number of iteration steps of dropping and adding terms.
trace If positive, information is printed during the model building. Larger values may give more information.
... Any additional arguments to add1.cca and drop1.cca.
Details

The basic functions for model choice in constrained ordination are `add1.cca` and `drop1.cca`. With these functions, ordination models can be chosen with standard R function `step` which bases the term choice on AIC. AIC-like statistics for ordination are provided by functions `deviance.cca` and `extractAIC.cca` (with similar functions for `rda`). Actually, constrained ordination methods do not have AIC, and therefore the `step` may not be trusted. This function provides an alternative using permutation P-values.

Function `ordistep` defines the model, scope of models considered, and direction of the procedure similarly as `step`. The function alternates with drop and add steps and stops when the model was not changed during one step. The – and + signs in the summary table indicate which stage is performed. The number of permutations is selected adaptively with respect to the defined decision limit. It is often sensible to have `Pout > Pin` in stepwise models to avoid cyclic adds and drops of single terms.

Function `ordiR2step` builds model so that it maximizes adjusted $R^2$ (function `RsquareAdj`) at every step, and stopping when the adjusted $R^2$ starts to decrease, or the adjusted $R^2$ of the scope is exceeded, or the selected permutation P-value is exceeded (Blanchet et al. 2008). The direction has choices "forward" and "both", but it is very exceptional that a term is dropped with the adjusted $R^2$ criterion. Function uses adjusted $R^2$ as the criterion, and it cannot be used if the criterion cannot be calculated. Therefore it is unavailable for `cca`.

Functions `ordistep` (based on P values) and `ordiR2step` (based on adjusted $R^2$ and hence on eigenvalues) can select variables in different order.

Value

Functions return the selected model with one additional component, `anova`, which contains brief information of steps taken. You can suppress voluminous output during model building by setting `trace = FALSE`, and find the summary of model history in the `anova` item.

Author(s)

Jari Oksanen

References


See Also

The function handles constrained ordination methods `cca`, `rda` and `capscale`. The underlying functions are `add1.cca` and `drop1.cca`, and the function is modelled after standard `step` (which also can be used directly but uses AIC for model choice, see `extractAIC.cca`). Function `ordiR2step` builds upon `RsquareAdj`.

Examples

```r
## See add1.cca for another example
```
### Dune data

data(dune)
data(dune.env)
mod0 <- rda(dune ~ 1, dune.env)  # Model with intercept only
mod1 <- rda(dune ~ ., dune.env)  # Model with all explanatory variables

## Example without scope. Default direction is "backward"
ordistep(mod1, perm.max = 200)

## Example of ordiR2step, forward
ordiR2step(mod0, scope = formula(mod1), direction="forward", perm.max = 200)

### Mite data

data(mite)
data(mite.env)
mite.hel = decostand(mite, "hel")
mod0 <- rda(mite.hel ~ 1, mite.env)  # Model with intercept only
mod1 <- rda(mite.hel ~ ., mite.env)  # Model with all explanatory variables

## Example of ordiR2step with default direction = "both"
## (This never goes "backward" but evaluates included terms.)
step.res <- ordiR2step(mod0, mod1, perm.max = 200)
step.res$anova  # Summary table

## Example of ordiR2step with direction = "forward"
step.res <- ordiR2step(mod0, scope = formula(mod1), direction="forward")
step.res <- ordiR2step(mod0, scope = formula(mod1), direction="forward", trace=0)
step.res$anova  # Summary table

---

**ordisurf**  
*Fit and Plot Smooth Surfaces of Variables on Ordination.*

**Description**

Function `ordisurf` fits a smooth surface for given variable and plots the result on ordination diagram.

**Usage**

## Default S3 method:
ordisurf(x, y, choices=c(1, 2), knots=10, family="gaussian", col="red", thinplate = TRUE, add = FALSE, display = "sites", w = weights(x), main, nlevels = 10, levels, labcex = 0.6, bubble = FALSE, cex = 1, select = FALSE, method = "GCV.Cp", gamma = 1, plot = TRUE, ...)

---

---

---
Arguments

x For `ordisurf` an ordination configuration, either a matrix or a result known by `scores`. For `plot.ordisurf` and `object` of class "`ordisurf" as returned by `ordisurf`.

y Variable to be plotted.

choices Ordination axes.

knots Number of initial knots in `gam` (one more than degrees of freedom). If `knots = 0` or `knots = 1` the function will fit a linear trend surface, and if `knots = 2` the function will fit a quadratic trend surface instead of a smooth surface.

family Error distribution in `gam`.

col Colour of contours.

thinplate Use thinplate splines in `gam`.

add Add contours on an existing diagram or draw a new plot.

display Type of scores known by `scores`: typically "sites" for ordinary site scores or "lc" for linear combination scores.

w Prior weights on the data. Concerns mainly `cca` and `decorana` results which have nonconstant weights.

main The main title for the plot, or as default the name of plotted variable in a new plot.

nlevels, levels Either a vector of levels for which contours are drawn, or suggested number of contours in `nlevels` if `levels` are not supplied.

labcex Label size in contours. Setting this zero will suppress labels.

bubble Use “bubble plot” for points, or vary the point diameter by the value of the plotted variable. If `bubble` is numeric, its value is used for the maximum symbol size (as in `cex`), or if `bubble = TRUE`, the value of `cex` gives the maximum. The minimum size will always be `cex = 0.4`. The option only has an effect if `add = FALSE`.

cex Character expansion of plotting symbols.

select Logical; specify `gam` argument "select". If this is `TRUE` then `gam` can add an extra penalty to each term so that it can be penalized to zero. This means that the smoothing parameter estimation that is part of fitting can completely remove terms from the model. If the corresponding smoothing parameter is estimated as zero then the extra penalty has no effect.
method character; the smoothing parameter estimation method. Options allowed are: "GCV.Cp" uses GCV for models with unknown scale parameter and Mallows’ Cp/UBRE/AIC for models with known scale; "GACV.Cp" as for "GCV.Cp" but uses GACV (Generalised Approximate CV) instead of GCV; "REML" and "ML" use restricted maximum likelihood or maximum likelihood estimation for both known and unknown scale; and "P-REML" and "P-ML" use REML or ML estimation but use a Pearson estimate of the scale.

gamma Multiplier to inflate model degrees of freedom in GCV or UBRE/AIC score by. This effectively places an extra penalty on complex models. An oft used value if gamma = 1.4.

plot logical; should any plotting be done by ordisurf? Useful if all you want is the fitted response surface model.

formula, data Alternative definition of the fitted model as x ~ y, or left-hand side is the ordination x and right-hand side the single fitted continuous variable y. The variable y must be in the working environment or in the data frame or environment given by data. All other arguments of are passed to the default method.

object An ordisurf result object.

newdata Coordinates in two-dimensional ordination for new points.

what character; what type of plot to produce. "contour" produces a contour plot of the response surface, see contour for details. "persp" produces a perspective plot of the same, see persp for details. "gam" plots the fitted GAM model, an object that inherits from class "gam" returned by ordisurf, see plot.gam.

... Other parameters passed to gam, or to the graphical functions. See Note below for exceptions.

Details

Function ordisurf fits a smooth surface using thinplate splines in gam, and uses predict.gam to find fitted values in a regular grid. The smooth surface can be fitted with an extra penalty that allows the entire smoother to be penalized back to 0 degrees of freedom, effectively removing the term from the model. The addition of this extra penalty is invoked by setting argument select to TRUE. The function plots the fitted contours with convex hull of data points either over an existing ordination diagram or draws a new plot. If select == TRUE and the smooth is effectively penalised out of the model, no contours will be plotted.

gam determines the degree of smoothness for the fitted response surface during model fitting. Argument method controls how gam performs this smoothness selection. See gam for details of the available options. Using "REML" or "ML" yields p-values for smooths with the best coverage properties if such things matter to you.

The function uses scores to extract ordination scores, and x can be any result object known by that function.

User can supply a vector of prior weights w. If the ordination object has weights, these will be used. In practise this means that the row totals are used as weights with cca or decorana results. If you do not like this, but want to give equal weights to all sites, you should set w = NULL. The
behaviour is consistent with \texttt{envfit}. For complete accordance with constrained \texttt{cca}, you should set \texttt{display = "lc"} (and possibly \texttt{scaling = 2}).

Function \texttt{calibrate} returns the fitted values of the response variable. The \texttt{newdata} must be coordinates of points for which the fitted values are desired. The function is based on \texttt{predict.gam} and will pass extra arguments to that function.

\textbf{Value}

Function is usually called for its side effect of drawing the contour plot. The function returns the result object of class "\texttt{ordisurf}" that inherits from \texttt{gam} used internally to fit the surface, but adds an item \texttt{grid} that contains the data for the grid surface. The item \texttt{grid} has elements \texttt{x} and \texttt{y} which are vectors of axis coordinates, and element \texttt{z} that is a matrix of fitted values for \texttt{contour}. The values outside the convex hull of observed points are \texttt{NA} in \texttt{z}. The \texttt{gam} component of the result can be used for further analysis like predicting new values (see \texttt{predict.gam}).

\textbf{Note}

The default is to use thinplate splines. These make sense in ordination as they have equal smoothing in all directions and are rotation invariant.

Graphical arguments supplied to \texttt{plot.ordisurf} are passed on to the underlying plotting functions, \texttt{contour}, \texttt{persp}, and \texttt{plot.gam}. The exception to this is that arguments \texttt{col} and \texttt{cex} can not currently be passed to \texttt{plot.gam} because of a bug in the way that function evaluates arguments when arranging the plot.

A work-around is to call \texttt{plot.gam} directly on the result of a call to \texttt{ordisurf}. See the Examples for an illustration of this.

\textbf{Author(s)}

Dave Roberts, Jari Oksanen and Gavin L. Simpson

\textbf{See Also}

For basic routines \texttt{gam}, and \texttt{scores}. Function \texttt{envfit} provides a more traditional and compact alternative.

\textbf{Examples}

```r
data(varespec)
data(varechem)
library(MASS)
vare.dist <- vegdist(varespec)
vare.mds <- isoMDS(vare.dist)
with(varechem, ordisurf(vare.mds, Baresoil, bubble = 5))

## as above but with extra penalties on smooth terms:
with(varechem, ordisurf(vare.mds, Baresoil, bubble = 5, col = "blue",
                        add = TRUE, select = TRUE))

## Cover of Cladina arbuscula
fit <- with(varespec, ordisurf(vare.mds, Cla.arb, family=quasipoisson))
```
## Get fitted values
calibrate(fit)

## Plot method
plot(fit, what = "contour")

## Plotting the "gam" object
plot(fit, what = "gam") ## 'col' and 'cex' not passed on
## or via plot.gam directly
plot.gam(fit, cex = 2, pch = 1, col = "blue")
## 'col' effects all objects drawn...

---

orditkplot

 Ordination Plot with Movable Labels

### Description

Function `orditkplot` produces an editable ordination plot with points and labels. The labels can be moved with mouse, and the edited plot can be saved as an encapsulated postscript file or exported via R `plot` function to other graphical formats, or saved in the R session for further processing.

### Usage

`orditkplot(x, display = "species", choices = 1:2, width, xlim, ylim,
    tcex = 0.8, tcol, pch = 1, pcol, pbg, pcex = 0.7, labels, ...)`

### Arguments

- **x**: An ordination result or any other object that `scores` can handle, or for the `plot` function the object dumped from the interactive `orditkplot` session.
- **display**: Type of `scores` displayed. For ordination scores this typically is either "species" or "sites", and for `orditkplot` result it is either "points" or "labels".
- **choices**: Axes displayed.
- **width**: Width of the plot in inches; defaults to the current width of the graphical device.
- **xlim, ylim**: x and y limits for plots: points outside these limits will be completely removed.
- **tcex**: Character expansion for text labels.
- **tcol**: Colour of text labels.
pch, pcol, pbg

Point type and outline and fill colours. Defaults pcol="black" and pbg="transparent". Argument pbg has an effect only in filled plotting characters pch = 21 to 25.

pceEx
Expansion factor for point size.

labels
Labels used instead of row names.

... Other arguments passed to the function. These can be graphical parameters (see par) used in the plot, or extra arguments to scores. These arguments are ignored in plot, but honoured in text and points.

Details

Function orditkplot uses tcltk package to draw Tcl/Tk based ordination graphics with points and labels. The function opens an editable canvas with fixed points, but the labels can be dragged with mouse to better positions or edited. In addition, it is possible to zoom to a part of the graph.

The function knows the following mouse operations:

- **Left mouse button** can be used to move labels to better positions. A line will connect a label to the corresponding point.
- **Double clicking left mouse button** opens a window where the label can be edited. After editing the label, hit the Return key.
- **Right mouse button** (or alternatively, Shift-Mouse button with one-button mouse) can be used for zooming to a part of the graph. Keeping the mouse button down and dragging will draw a box of the zoomed area, and after releasing the button, a new plot window will be created (this is still preliminary: all arguments are not passed to the new plot).

In addition there are buttons for the following tasks: **Copy to EPS** copies the current plot to an encapsulated postscript (eps) file using standard Tcl/Tk utilities. The faithfulness of this copy is system dependent. **Button Export plot** uses plot.orditkplot function to redraw the plot into graphical file formats. Depending on the system, the following graphical formats may be available: eps, pdf, png, jpeg or bmp. The file type is deduced from the file suffix or the selection of the file type in the dialogue box. Alternatively, the same dialogue can be used to save the plot to an editable xfig file. **Button Dump to R** writes the edited coordinates of labels and points to the R session for further processing, and the plot.orditkplot function can be used to display the results. For faithful replication of the plot, the graph must have similar dimensions as the orditkplot canvas had originally. The plot function cannot be configured, but it uses the same settings as the original Tcl/Tk plot. However, points and text functions are fully configurable, and unaware of the original Tcl/Tk plot settings (probably you must set cex at least to get a decent plot). Finally, **button Dismiss** closes the window.

The produced plot will have equal aspect ratio. The width of the horizontal axis is fixed, but vertical axes will be scaled to needed height, and you can use scrollbar to move vertically if the whole canvas does not fit the window. If you use dumped labels in ordinary R plots, your plot must have the same dimensions as the orditkplot canvas to have identical location of the labels.

The function only displays one set of scores. However, you can use ordipointlabel to produce a result object that has different points and text types for several sets of scores and this can further edited with orditkplot. For a good starting solution you need to scale the ordipointlabel result so that the points span over the whole horizontal axis.
The plot is a Tcl/Tk canvas, but the function tries to replicate standard graphical device of the platform, and it honours several graphical parameters (see `par`). Many of the graphical parameters can be given on the command line, and they will be passed to the function without influencing other graphical devices in R. At the moment, the following graphical parameters are honoured: `pch bg`, `cex`, `cex.axis`, `cex.lab`, `col` (for labels), `col.axis`, `col.lab`, `family` (for font faces), `fg`, `font`, `font.axis`, `font.lab`, `lheight`, `lwd` (for the box), `mar`, `mex`, `mgp`, `ps`, `tcl`. These can be set with `par`, and they also will influence other plots similarly.

The Tk canvas text cannot be rotated, and therefore vertical axis is not labelled, and `las` parameter will not be honoured in the Tcl/Tk plot, but it will be honoured in the exported R plots and in `plot.orditkplot`.

**Value**

Function returns nothing useful directly, but you can save the edited graph to a file or dump the edited positions to an R session for further processing and plotting.

**Note**

You need `tcltk` package and R must have been configured with `capabilities` for `tcltk` when building the binary. Depending on your OS, you may need to start X11 and set the display before loading `tcltk` and starting the function (for instance, with `Sys.setenv("DISPLAY"=":0")`). See `tcltk-package`.

**Author(s)**

Jari Oksanen

**See Also**

Function `ordipointlabel` is an automatic procedure with similar goals of avoiding overplotting. See `ordiplot`, `plot.cca`, `ordirgl` and `orditorp` for alternative ordination plots, and `scores` for extracting ordination scores.

**Examples**

```r
## The example needs user interaction and is not executed directly.
## It should work when pasted to the window.
## Not run:
data(varespec)
ord <- cca(varespec)
## Do something with the graph and end by clicking "Dismiss"
orditkplot(ord, mar = c(4,4,1,1)+.1, font=3)
## Use ordipointlabel to produce a plot that has both species and site
## scores in different colors and plotting symbols
pl <- ordipointlabel(ord)
orditkplot(pl)
## End(Not run)
```
orditorp  

Add Text or Points to Ordination Plots

Description

The function adds text or points to ordination plots. Text will be used if this can be done without overwriting other text labels, and points will be used otherwise. The function can help in reducing clutter in ordination graphics, but manual editing may still be necessary.

Usage

orditorp(x, display, labels, choices = c(1, 2), priority, cex = 0.7, pcesx, col = par("col"), pcol, pch = par("pch"), air = 1, ...)

Arguments

x  
A result object from ordination or an ordiplot result.
display  
Items to be displayed in the plot. Only one alternative is allowed. Typically this is "sites" or "species".
labels  
Optional text used for labels. Row names will be used if this is missing.
choices  
Axes shown.
priority  
Text will be used for items with higher priority if labels overlap. This should be vector of the same length as the number of items plotted.
cex, pcesx  
Text and point sizes, see plot.default..
col, pcol  
Text and point colours, see plot.default.
pch  
Plotting character, see points.
air  
Amount of empty space between text labels. Values <1 allow overlapping text.
...  
Other arguments to scores (and its various methods), text and points.

Details

Function orditorp will add either text or points to an existing plot. The items with high priority will be added first and text will be used if this can be done without overwriting previous labels, and points will be used otherwise. If priority is missing, labels will be added from the outskirts to the centre. Function orditorp can be used with most ordination results, or plotting results from ordiplot or ordination plot functions (plot.cca, plot.decorana, plot.metaMDS).

Arguments can be passed to the relevant scores method for the ordination object (x) being drawn. See the relevant scores help page for arguments that can be used.

Value

The function returns invisibly a logical vector where TRUE means that item was labelled with text and FALSE means that it was marked with a point. The returned vector can be used as the select argument in ordination text and points functions.
Examples

```r
## A cluttered ordination plot :
data(BCI)
mod <- cca(BCI)
plot(mod, dis="sp", type="t")
# Now with orditorp and abbreviated species names
cnam <- make.cepnames(names(BCI))
plot(mod, dis="sp", type="n")
stems <- colSums(BCI)
orditorp(mod, "sp", label = cnam, priority = stems, pch="+", pcol="grey")
```

Description

Functions `ordicloud`, `ordisplom` and `ordixyplot` provide an interface to plot ordination results using Trellis functions `cloud`, `splom` and `xyplot` in package `lattice`.

Usage

```r
ordixyplot(x, data = NULL, formula, display = "sites", choices = 1:3,
 panel = "panel.ordi", aspect = "iso", envfit,
 type = c("p", "biplot"), ...)
ordisplom(x, data = NULL, formula = NULL, display = "sites", choices = 1:3,
 panel = "panel.ordi", type = "p", ...)
ordicloud(x, data = NULL, formula, display = "sites", choices = 1:3,
 panel = "panel.ordi3d", prepanel = "prepanel.ordi3d", ...)
```

Arguments

- **x**: An ordination result that `scores` knows: any ordination result in `vegan` and many others.
- **data**: Optional data to amend ordination results. The ordination results are found from `x`, but you may give here data for other variables needed in plots. Typically these are environmental data.
- **formula**: Formula to define the plots. A default formula will be used if this is omitted. The ordination axes must be called by the same names as in the ordination results (and these names vary among methods). In `ordisplom`, special character . refers to the ordination result.
- **display**: The kind of scores: an argument passed to `scores`.
- **choices**: The axes selected: an argument passed to `scores`. 
**ordixyplot**

- **panel, prepanel**
  - The names of the panel and prepanel functions.

- **aspect**
  - The aspect of the plot (passed to the **lattice** function).

- **envfit**
  - Result of **envfit** function displayed in **ordixyplot**. Please note that this needs same choices as **ordixyplot**.

- **type**
  - The type of plot. This knows the same alternatives as **panel.xyplot**. In addition **ordixyplot** has alternatives "biplot" and "arrows". The first displays fitted vectors and factor centroids of **envfit**, or in constrained ordination, the biplot arrows and factor centroids if **envfit** is not given. The second (type = "arrows") is a trellis variant of **ordiarrows** and draws arrows by groups. The line parameters are controlled by **trellis.par.set** for **superpose.line**, and the user can set length, angle and ends parameters of **panel.arrows**.

- **...**
  - Arguments passed to **scores** methods or **lattice** functions.

**Details**

The functions provide an interface to the corresponding **lattice** functions. All graphical parameters are passed to the **lattice** function so that these graphs are extremely configurable. See **Lattice** and **xyplot.splom** and **cloud** for details, usage and possibilities.

The argument **x** must always be an ordination result. The scores are extracted with **vegan** function **scores** so that these functions work with all **vegan** ordinations and many others.

The **formula** is used to define the models. All functions have simple default formulae which are used if **formula** is missing. If **formula** is omitted in **ordisplom** it produces a pairs plot of ordination axes and variables in **data**. If **formula** is given, ordination results must be referred to as . and other variables by their names. In other functions, the formula must use the names of ordination scores and names of **data**.

The ordination scores are found from **x**, and **data** is optional. The **data** should contain other variables than ordination scores to be used in plots. Typically, they are environmental variables (typically factors) to define panels or plot symbols.

The proper work is done by the panel function. The layout can be changed by defining own panel functions. See **panel.xyplot**, **panel.splom** and **panel.cloud** for details and survey of possibilities.

Ordination graphics should always be isometric: same scale should be used in all axes. This is controlled (and can be changed) with argument **aspect** in **ordixyplot**. In **ordicloud** the isometric scaling is defined in **panel** and **prepanel** functions. You must replace these functions if you want to have non-isometric scaling of graphs. You cannot select isometric scaling in **ordisplom**.

**Value**

The function return **Lattice** objects of class "**trellis**".

**Author(s)**

Jari Oksanen
See Also

Lattice, xyplot, splom, cloud, panel.splom, panel.cloud

Examples

data(dune)
data(dune.env)
ord <- cca(dune)
## Pairs plots
ordisplom(ord)
ordisplom(ord, data=dune.env, choices=1:2)
ordisplom(ord, data=dune.env, form = ~ . | Management, groups=Manure)
## Scatter plot
ordixyplot(ord, data=dune.env, form = CA1 ~ CA2 | Management,
    groups=Manure)
## Choose a different scaling
ordixyplot(ord, scaling = 3)
## ... Slices of third axis
ordixyplot(ord, form = CA1 ~ CA2 | equal.count(CA3, 4), type = c("g","p"))
## Display environmental variables
ordixyplot(ord, envfit = envfit(ord ~ Management + A1, dune.env, choices=1:3))
## 3D Scatter plots
ordicloud(ord, form = CA2 ~ CA3*CA1, groups = Manure, data = dune.env)
ordicloud(ord, form = CA2 ~ CA3*CA1 | Management, groups = Manure,
data = dune.env, auto.key = TRUE, type = c("p","h"))

Description

This function computed classical PCNM by the principal coordinate analysis of a truncated distance
matrix. These are commonly used to transform (spatial) distances to rectangular data that suitable
for constrained ordination or regression.

Usage

pcnm(dis, threshold, w, dist.ret = FALSE)

Arguments

dis A distance matrix.
threshold A threshold value or truncation distance. If missing, minimum distance giving
connected network will be used. This is found as the longest distance in the
minimum spanning tree of dis.
w Prior weights for rows.
dist.ret Return the distances used to calculate the PCNMs.
Details

Principal Coordinates of Neighbourhood Matrix (PCNM) map distances between rows onto rectangular matrix on rows using a truncation threshold for long distances (Borcard & Legendre 2002). If original distances were Euclidean distances in two dimensions (like normal spatial distances), they could be mapped onto two dimensions if there is no truncation of distances. Because of truncation, there will be a higher number of principal coordinates. The selection of truncation distance has a huge influence on the PCNM vectors. The default is to use the longest distance to keep data connected. The distances above truncation threshold are given an arbitrary value of 4 times threshold. For regular data, the first PCNM vectors show a wide scale variation and later PCNM vectors show smaller scale variation (Borcard & Legendre 2002), but for irregular data the interpretation is not as clear.

The PCNM functions are used to express distances in rectangular form that is similar to normal explanatory variables used in, e.g., constrained ordination (rda, cca and capscale) or univariate regression (lm) together with environmental variables (row weights should be supplied with cca; see Examples). This is regarded as a more powerful method than forcing rectangular environmental data into distances and using them in partial mantel analysis (mantel.partial) together with geographic distances (Legendre et al. 2008, but see Tuomisto & Ruokolainen 2008).

The function is based on pcnm function in Dray’s unreleased spacemakeR package. The differences are that the current function uses rspantree as an internal support function. The current function also can use prior weights for rows by using weighted metric scaling of wcmdscale. The use of row weights allows finding orthonormal PCNMs also for correspondence analysis (e.g., cca).

Value

A list of the following elements:

- **values**: Eigenvalues obtained by the principal coordinates analysis.
- **vectors**: Eigenvectors obtained by the principal coordinates analysis. They are scaled to unit norm. The vectors can be extracted with scores function. The default is to return all PCNM vectors, but argument choices selects the given vectors.
- **threshold**: Truncation distance.
- **dist**: The distance matrix where values above threshold are replaced with arbitrary value of four times the threshold. String "pcnm" is added to the method attribute, and new attribute threshold is added to the distances. This is returned only when dist.ret = TRUE.

Author(s)

Jari Oksanen, based on the code of Stephane Dray.

References


See Also

*spantree*.

Examples

```r
## Example from Borcard & Legendre (2002)
data(mite.xy)
pcnm1 <- pcnm(dist(mite.xy))
op <- par(mfrow=c(1,3))
## Map of PCNMs in the sample plot
ordisurf(mite.xy, scores(pcnm1, choi=1), bubble = 4, main = "PCNM 1")
ordisurf(mite.xy, scores(pcnm1, choi=2), bubble = 4, main = "PCNM 2")
ordisurf(mite.xy, scores(pcnm1, choi=3), bubble = 4, main = "PCNM 3")
par(op)
## Plot first PCNMs against each other
ordisplom(pcnm1, choices=1:4)
## Weighted PCNM for CCA
data(mite)
rs <- rowSums(mite)/sum(mite)
pcnmw <- pcnm(dist(mite.xy), w = rs)
ord <- cca(mite ~ scores(pcnmw))
## Multiscale ordination: residual variance should have no distance
trend
msoplot(mso(ord, mite.xy))
```

permat

*Matrix Permutation Algorithms for Presence-Absence and Count Data*

Description

Individual (for count data) or incidence (for presence-absence data) based null models can be generated for community level simulations. Options for preserving characteristics of the original matrix (rows/columns sums, matrix fill) and restricted permutations (based on strata) are discussed in the Details section.

Usage

```r
permatfull(m, fixedmar = "both", shuffle = "both", strata = NULL, mtype = "count", times = 99)
permatswap(m, method = "quasiswap", fixedmar="both", shuffle = "both", strata = NULL, mtype = "count", times = 99, burnin = 0, thin = 1)
## S3 method for class 'permat'
print(x, digits = 3, ...)
## S3 method for class 'permat'
summary(object, ...)```
## S3 method for class 'summary.permat'
print(x, digits = 2, ...)
## S3 method for class 'permat'
plot(x, type = "bray", ylab, xlab, col, lty,
    lowess = TRUE, plot = TRUE, text = TRUE, ...)
## S3 method for class 'permat'
lines(x, type = "bray", ...)
## S3 method for class 'permat'
as.ts(x, type = "bray", ...)
## S3 method for class 'permat'
as.mcmc(x)

### Arguments

- **m**
  A community data matrix with plots (samples) as rows and species (taxa) as columns.

- **fixedmar**
  Character, stating which of the row/column sums should be preserved ("none", "rows", "columns", "both").

- **strata**
  Numeric vector or factor with length same as nrow(m) for grouping rows within strata for restricted permutations. Unique values or levels are used.

- **mtype**
  Matrix data type, either "count" for count data, or "prab" for presence-absence type incidence data.

- **times**
  Number of permuted matrices.

- **method**
  Character for method used for the swap algorithm ("swap", "tswap", "quasiswap", "backtrack") as described for function `commsimulator`. If mtype="count" the "quasiswap", "swap", "swsh" and "abuswap" methods are available (see details).

- **shuffle**
  Character, indicating whether individuals ("ind"), samples ("samp") or both ("both") should be shuffled, see details.

- **burnin**
  Number of null communities discarded before proper analysis in sequential ("swap", "tswap") methods.

- **thin**
  Number of discarded permuted matrices between two evaluations in sequential ("swap", "tswap") methods.

- **x, object**
  Object of class "permat".

- **digits**
  Number of digits used for rounding.

- **ylab, xlab, col, lty**
  Graphical parameters for the `plot` method.

- **type**
  Character, type of plot to be displayed: "bray" for Bray-Curtis dissimilarities, "chisq" for Chi-squared values.

- **lowess, plot, text**
  Logical arguments for the `plot` method, whether a locally weighted regression curve should be drawn, the plot should be drawn, and statistic values should be printed on the plot.

- **...**
  Other arguments passed to methods.
Details

The function `permatfull` is useful when matrix fill is allowed to vary, and matrix type is `count`. The `fixedmar` argument is used to set constraints for permutation. If none of the margins are fixed, cells are randomised within the matrix. If rows or columns are fixed, cells within rows or columns are randomised, respectively. If both margins are fixed, the `r2dtable` function is used that is based on Patefield’s (1981) algorithm. For presence absence data, matrix fill should be necessarily fixed, and `permatfull` is a wrapper for the function `commsimulator`. The `r00`, `r0`, `c0`, `quasiswap` algorithms of `commsimulator` are used for "none", "rows", "columns", "both" values of the `fixedmar` argument, respectively.

The `shuffle` argument only have effect if the `mtype = "count"` and `permatfull` function is used with "none", "rows", "columns" values of `fixedmar`. All other cases for count data are individual based randomisations. The "samp" and "both" options result fixed matrix fill. The "both" option means that individuals are shuffled among non zero cells ensuring that there are no cell with zeros as a result, then cell (zero and new valued cells) are shuffled.

The function `permatswap` is useful when with matrix fill (i.e. the proportion of empty cells) and row/columns sums should be kept constant. `permatswap` uses different kinds of swap algorithms, and row and columns sums are fixed in all cases. For presence-absence data, the `swap` and `tswap` methods of `commsimulator` can be used. For count data, a special swap algorithm ("swapcount") is implemented that results in permuted matrices with fixed marginals and matrix fill at the same time.

The 'quasiswapcount' algorithm (method="quasiswap" and `mtype="count"`) uses the same trick as Carsten Dormann's `swap.web` function in the package `bipartite`. First, a random matrix is generated by the `r2dtable` function retaining row and column sums. Then the original matrix fill is reconstructed by sequential steps to increase or decrease matrix fill in the random matrix. These steps are based on swapping 2x2 submatrices (see 'swapcount' algorithm for details) to maintain row and column totals. This algorithm generates independent matrices in each step, so `burnin` and `thin` arguments are not considered. This is the default method, because this is not sequential (as `swapcount` is) so independence of subsequent matrices does not have to be checked.

The `swapcount` algorithm (method="swap" and `mtype="count"`) tries to find 2x2 submatrices (identified by 2 random row and 2 random column indices), that can be swapped in order to leave column and row totals and fill unchanged. First, the algorithm finds the largest value in the submatrix that can be swapped (d) and whether in diagonal or antidiagonal way. Submatrices that contain values larger than zero in either diagonal or antidiagonal position can be swapped. Swap means that the values in diagonal or antidiagonal positions are decreased by d, while remaining cells are increased by d. A swap is made only if fill doesn’t change. This algorithm is sequential, subsequent matrices are not independent, because swaps modify little if the matrix is large. In these cases many burnin steps and thinning is needed to get independent random matrices. Although this algorithm is implemented in C, large burnin and thin values can slow it down considerably. WARNING: according to simulations, this algorithm seems to be biased and non random, thus its use should be avoided!

The algorithm "swsh" in the function `permatswap` is a hybrid algorithm. First, it makes binary quasiswaps to keep row and column incidences constant, then non-zero values are modified according to the `shuffle` argument (only "samp" and "both" are available in this case, because it is applied only on non-zero values).

The algorithm "abuswap" produces two kinds of null models (based on `fixedmar="columns"` or `fixedmar="rows"`) as described in Hardy (2008; randomization scheme 2x and 3x, respec-
permat

These preserve column and row occurrences, and column or row sums at the same time.

Constraints on row/column sums, matrix fill, total sum and sums within strata can be checked by the summary method. plot method is for visually testing the randomness of the permuted matrices, especially for the sequential swap algorithms. If there are any tendency in the graph, higher burnin and thin values can help for sequential methods. New lines can be added to existing plot with the lines method.

Unrestricted and restricted permutations: if strata is NULL, functions perform unrestricted permutations. Otherwise, it is used for restricted permutations. Each strata should contain at least 2 rows in order to perform randomization (in case of low row numbers, swap algorithms can be rather slow). If the design is not well balanced (i.e. same number of observations within each stratum), permuted matrices may be biased because same constraints are forced on submatrices of different dimensions. This often means, that the number of potential permutations will decrease with their dimensions. So the more constraints we put, the less randomness can be expected.

The plot method is useful for graphically testing for trend and independence of permuted matrices. This is especially important when using sequential algorithms ("swap", "tswap", "abuswap").

The as.ts method can be used to extract Bray-Curtis dissimilarities or Chi-squared values as time series. This can further used in testing independence (see Examples). The method as.mcmc is useful for accessing diagnostic tools available in the coda package.

Value

Functions permatfull and permatswap return an object of class "permat" containing the the function call (call), the original data matrix used for permutations (orig) and a list of permuted matrices with length times (perm).

The summary method returns various statistics as a list (including mean Bray-Curtis dissimilarities calculated pairwise among original and permuted matrices, Chi-square statistics, and check results of the constraints; see Examples). Note that when strata is used in the original call, summary calculation may take longer.

The plot creates a plot as a side effect.

The as.ts method returns an object of class "ts".

Author(s)

Péter Sólymos, <solymos@ualberta.ca> and Jari Oksanen

References

Original references for presence-absence algorithms are given on help page of commsimulator.


See Also

For other functions to permute matrices: `commsimulator`, `r2dtable`, `sample`, `swap.web`.

For the use of these permutation algorithms: `oecosimu`, `adipart`, `hiersimu`.

For time-series diagnostics: `Box.test`, `lag.plot`, `tsdiag`, `ar`, `arima`.

Examples

```r
## A simple artificial community data matrix.
m <- matrix(c(
  1, 3, 2, 0, 3, 1,
  0, 2, 1, 0, 2, 1,
  0, 0, 1, 2, 0, 3,
  0, 0, 0, 1, 4, 3
), 4, 6, byrow=TRUE)
## Using the quasiswap algorithm to create a
## list of permuted matrices, where
## row/columns sums and matrix fill are preserved:
x1 <- permatswap(m, "quasiswap")
summary(x1)
## Unrestricted permutation retaining
## row/columns sums but not matrix fill:
x2 <- permatfull(m)
summary(x2)
## Unrestricted permutation of presence-absence type
## not retaining row/columns sums:
x3 <- permatfull(m, "none", mtype="prab")
x3$orig ## note: original matrix is binarized!
summary(x3)
## Restricted permutation,
## check sums within strata:
x4 <- permatfull(m, strata=c(1,1,2,2))
summary(x4)
## Not sequential algorithm
data(BCI)
a <- permatswap(BCI, "quasiswap")
## Sequential algorithm
b <- permatswap(BCI, "abuswap", fixedmar="col",
  burnin=0, thin=100, times=50)
## Extract Bray-Curtis dissimilarities
bc <- as.ts(b)
## Lag plot
lag.plot(bc)
## First order autoregressive model
mar <- arima(bc, c(1,0,0))
```
## Ljung-Box test of residuals
Box.test(mar$residuals)
## Graphical diagnostics
tsdia(mar)

---

permCheck | Utility functions for permutation schemes

### Description

permCheck provides checking of permutation schemes for validity. numPerms calculates the maximum number of permutations possible under the current permutation scheme. allPerms enumerates all possible permutations for the given scheme. getNumObs is a utility function to return the number of observations for a range of R and ordination objects. permuplot produces a graphical representation of the selected permutation design.

### Usage

```r
permCheck(object, control = permControl(), make.all = TRUE)
```

```r
## S3 method for class 'permCheck'
summary(object, ...)
```

```r
numPerms(object, control = permControl())
```

```r
allPerms(n, control = permControl(), max = 9999,
observed = FALSE)
```

```r
## S3 method for class 'allPerms'
summary(object, ...)
```

```r
getNumObs(object, ...)
```

```r
## Default S3 method:
getNumObs(object, ...)
```

```r
## S3 method for class 'numeric'
getNumObs(object, ...)
```

```r
## S3 method for class 'integer'
getNumObs(object, ...)
```

```r
permuplot(n, control = permControl(), col = par("col"),
hcol = "red", shade = "lightgrey", xlim = NULL, ylim = NULL,
inset = 0.1, main = NULL, sub = NULL, ann = par("ann"),
cex = par("cex"), ...)
```
Arguments

object an R object. Specifically, for `getNumObs` any object handled by `scores`, data frames, matrices, and numeric and integer vectors. See Details for a complete description, especially for `numPerms`. For `summary.permCheck` an object of class "permCheck". For `summary.allPerms` an object of class "allPerms".

control a list of control values describing properties of the permutation design, as returned by a call to `permControl`

make.all logical; should `permCheck` generate all possible permutations? Useful if want to check permutation design but not produce the matrix of all permutations.

n the number of observations or an 'object' from which the number of observations can be determined via `getNumObs`.

max the maximum number of permutations, below which complete enumeration will be attempted. See Details.

observed logical, should the observed ordering of samples be returned as part of the complete enumeration? Default is FALSE to facilitate usage in higher level functions.

col, xlim, ylim, main, sub, ann, cex
Graphical parameters.

hcol Colour to use for highlighting observations and the border colour of the polygons drawn when `type = "strata"`

shade The polygon shading colour (passed to argument `col` of function `polygon`) when `type = "strata"`

inset Proportion of range of x and y coordinates to add to the plot x and y limits. Used to create a bit of extra space around the margin of each plot.

... arguments to other methods. For `permuplot` graphical parameters can be passed to plotting functions, though note that not all parameters will be accepted gracefully at the moment.

Details

`permCheck`, `allPerms`, `numPerms` and `permuplot` are utility functions for working with the new permutation schemes available in `permuted.index2`.

`permCheck` is used to check the current permutation schemes against the object to which it will be applied. It calculates the maximum number of possible permutations for the number of observations in `object` and the permutation scheme described by `control`. The returned object contains component `control`, an object of class "permControl" suitably modified if `permCheck` identifies a problem.

The main problem is requesting more permutations than possible with the number of observations and the permutation design. In such cases, `nperm` is reduced to equal the number of possible permutations, and complete enumeration of all permutations is turned on (`control$complete` is set to TRUE).

Alternatively, if the number of possible permutations is low, and less than `control$minperm`, it is better to enumerate all possible permutations, and as such complete enumeration of all permutations is turned on (`control$complete` is set to TRUE).
permCheck

Function `numPerms` returns the number of permutations for the passed `object` and the selected permutation scheme. `object` can be one of a data frame, matrix, an object for which a `scores` method exists, or a numeric or integer vector. In the case of a numeric or integer vector, a vector of length 1 can be used and it will be expanded to a vector of length `object` (i.e., `1:object`) before computing the number of permutations. As such, `object` can be the number of observations not just the object containing the observations.

Function `allPerms` enumerates all possible permutations for the number of observations and the selected permutation scheme. It has `print` and `summary` methods. `allPerms` returns a matrix containing all possible permutations, possibly containing the observed ordering (if argument `observed` is `TRUE`). The rows of this matrix are the various permutations and the columns reflect the number of samples.

With free permutation designs, and restricted permutation schemes with large numbers of observations, there are a potentially huge number of possible permutations of the samples. It would be inefficient, not to mention incredibly time consuming, to enumerate them all. Storing all possible permutations would also become problematic in such cases. To control this and guard against trying to evaluate too large a number of permutations, if the number of possible permutations is larger than `max`, `allPerms` exits with an error.

Function `getNumObs` is a simple generic function to return the number of observations in a range of `R` objects. The default method will work for any object for which a `scores` method exists. This includes matrices and data frames, as well as specific methods for numeric or integer vectors.

`permuplot` is a graphical utility function, which produces a graphical representation of a permutation design. It takes the number of observations and an object returned by `permControl` as arguments and produces a plot on the currently active device. If strata are present in the design, the plotting region is split into sufficient plotting regions (one for each stratum), and the design in each stratum plotted.

Free permutation designs are represented by plotting the observation number at random x and y coordinates. Series designs (time series or line transects) are represented by plotting the observation numbers comprising the series in a circle and the start of the permuted series is highlighted using colour `hcol`. Grid designs are drawn on a regular grid and the top left observation in the original grid is highlighted using colour `hcol`. Note the ordering used is R’s standard ordering for matrices - columns are filled first.

Value

For `permCheck` a list containing the maximum number of permutations possible and an object of class "permControl".

For `allPerms`, and object of class "allPerms", a matrix whose rows are the set of all possible permutations for the supplies number of observations and permutation scheme selected. The matrix has two additional attributes `control` and `observed`. Attribute `control` contains the argument `control` (possibly updated via `permCheck`). Attribute `observed` contains argument `observed`.

For `numPerms`, the (numeric) number of possible permutations.

For `getNumObs`, the (numeric) number of observations in `object`.

For `permuplot`, a plot on the currently active device.
Note

In general, mirroring "series" or "grid" designs doubles or quadruples, respectively, the number of permutations without mirroring (within levels of strata if present). This is not true in two special cases:

1. In "grid" designs where the number of columns is equal to 2, and
2. In "series" designs where the number of observations in a series is equal to 2.

For example, with 2 observations there are 2 permutations for "series" designs:

1. 1-2, and
2. 2-1.

If these two permutations were mirrored, we would have:

1. 2-1, and
2. 1-2.

It is immediately clear that this is the same set of permutations without mirroring (if one reorders the rows). A similar situation arises in "grid" designs where the number of columns per grid is equal to 2. Note that the number of rows per grid is not an issue here.

Author(s)

Gavin Simpson

See Also

permuted.index2 and permControl.

Examples

```r
## use example data from ?pyrifos
eample(pyrifos)

## Demonstrate the maximum number of permutations for the pyrifos data
## under a series of permutation schemes

## no restrictions - lots of perms
(check1 <- permCheck(pyrifos, control = permControl(type = "free")))
summary(check1)

## no strata but data are series with no mirroring, so 132 permutations
permCheck(pyrifos, control = permControl(type = "series",
mirror = FALSE))

## no strata but data are series with mirroring, so 264 permutations
permCheck(pyrifos, control = permControl(type = "series",
mirror = TRUE))

## unrestricted within strata
```
permCheck(pyrifos, control = permControl(strata = ditch, type = "free"))

## time series within strata, no mirroring
permCheck(pyrifos, control = permControl(strata = ditch, type = "series", mirror = FALSE))

## time series within strata, with mirroring
permCheck(pyrifos, control = permControl(strata = ditch, type = "series", mirror = TRUE))

## time series within strata, no mirroring, same permutation within strata
permCheck(pyrifos, control = permControl(strata = ditch, type = "series", constant = TRUE))

## time series within strata, with mirroring, same permutation within strata
permCheck(pyrifos, control = permControl(strata = ditch, type = "series", mirror = TRUE, constant = TRUE))

## permute strata
permCheck(pyrifos, permControl(strata = ditch, type = "free", permute.strata = TRUE))

## this should also also for arbitrary vectors
vec1 <- permCheck(1:100)
vec2 <- permCheck(1:100, permControl())
all.equal(vec1, vec2)
vec3 <- permCheck(1:100, permControl(type = "series"))
all.equal(100, vec3$n)
vec4 <- permCheck(1:100, permControl(type = "series", mirror = TRUE))
all.equal(vec4$n, 200)

## enumerate all possible permutations
fac <- g1(2, 6)
ctrl <- permControl(type = "grid", mirror = FALSE, strata = fac, constant = TRUE, nrow = 3, ncol = 2)
numPerms(1:12, control = ctrl)
(tmp <- allPerms(12, control = ctrl, observed = TRUE))
(tmp2 <- allPerms(12, control = ctrl))

## turn on mirroring
ctrl$mirror <- TRUE
numPerms(1:12, control = ctrl)
(tmp3 <- allPerms(12, control = ctrl, observed = TRUE))
(tmp4 <- allPerms(12, control = ctrl))

## prints out details of the permutation scheme as well as the matrix of permutations
summary(tmp)
summary(tmp2)

## different numbers of observations per level of strata
fac <- factor(rep(1:3, times = c(3, 2, 2)))

## free permutations in levels of strata
numPerms(7, permControl(type = "free", strata = fac))
allPerms(7, permControl(type = "free", strata = fac))
## series permutations in levels of strata
numPerms(7, permControl(type = "series", strata = fac))
allPerms(7, permControl(type = "series", strata = fac))

## allPerms can work with a vector
vec <- c(3,4,5)
allPerms(vec)

## Tests for permuplot
n <- 25
## standard permutation designs
permuplot(n, permControl(type = "free"))
permuplot(n, permControl(type = "series"))
permuplot(n, permControl(type = "grid", nrow = 5, ncol = 5))

## restricted perms with mirroring
permuplot(n, permControl(type = "series", mirror = TRUE))
permuplot(n, permControl(type = "grid", nrow = 5, ncol = 5, mirror = TRUE))

## perms within strata
fac <- gl(6, 20)
control <- permControl(type = "free", strata = fac)
permuplot(120, control = control, cex = 0.8)
control <- permControl(type = "series", strata = fac)
permuplot(120, control = control, cex = 0.8)
fac <- gl(6, 25)
control <- permControl(type = "grid", strata = fac,
                       nrow = 5, ncol = 5)
permuplot(150, control = control, cex = 0.8)

## perms within strata with mirroring
fac <- gl(6, 20)
control <- permControl(type = "series", strata = fac,
                       mirror = TRUE)
permuplot(120, control = control, cex = 0.8)
fac <- gl(6, 25)
control <- permControl(type = "grid", strata = fac,
                       nrow = 5, ncol = 5, mirror = TRUE)
permuplot(150, control = control, cex = 0.8)

## same perms within strata
fac <- gl(6, 20)
control <- permControl(type = "free", strata = fac,
                       constant = TRUE)
permuplot(120, control = control, cex = 0.8)
control <- permControl(type = "series", strata = fac,
                       constant = TRUE)
permuplot(120, control = control, cex = 0.8)
fac <- gl(6, 25)
control <- permControl(type = "grid", strata = fac,
                       nrow = 5, ncol = 5, constant = TRUE)
permutations

permuplot(150, control = control, cex = 0.8)

## same perms within strata with mirroring
fac <- gl(6, 20)
control <- permControl(type = "series", strata = fac,
mirror = TRUE, constant = TRUE)
permuplot(120, control = control, cex = 0.8)
fac <- gl(6, 25)
control <- permControl(type = "grid", strata = fac,
  nrow = 5, ncol = 5, mirror = TRUE,
  constant = TRUE)
permuplot(150, control = control, cex = 0.8)

permutations  Permutation tests in Vegan

Description

Unless stated otherwise, vegan currently provides for two types of permutation test:

1. Free permutation of DATA, also known as randomisation, and
2. Free permutation of DATA within the levels of a factor variable.

We use DATA to mean either the observed data themselves or some function of the data, for example the residuals of an ordination model in the presence of covariables.

The second type of permutation test above is available if the function providing the test accepts an argument strata or passes additional arguments (via ...) to permuted.index.

The Null hypothesis for these two types of permutation test assumes free exchangeability of DATA (within the levels of strata if specified). Dependence between observations, such as that which arises due to spatial or temporal autocorrelation, or more-complicated experimental designs, such as split-plot designs, violates this fundamental assumption of the test and requires restricted permutation test designs. The next major version of Vegan will include infrastructure to handle these more complicated permutation designs.

Again, unless otherwise stated in the help pages for specific functions, permutation tests in Vegan all follow the same format/structure:

1. An appropriate test statistic is chosen. Which statistic is chosen should be described on the help pages for individual functions.
2. The value of the test statistic is evaluate for the observed data and analysis/model and recorded. Denote this value \( x_0 \).
3. The DATA are randomly permuted according to one of the above two schemes, and the value of the test statistic for this permutation is evaluated and recorded.
4. Step 3 is repeated a total of \( n \) times, where \( n \) is the number of permutations requested. Denote these values as \( x_i \), where \( i = 1, ..., n \).
5. The values of the test statistic for the \( n \) permutations of the \( DATA \) are added to the value of the test statistic for the observed data. These \( n + 1 \) values represent the Null or randomisation distribution of the test statistic. The observed value for the test statistic is included in the Null distribution because under the Null hypothesis being tested, the observed value is just a typical value of the test statistic, inherently no different from the values obtained via permutation of \( DATA \).

6. The number of times that a value of the test statistic in the Null distribution is equal to or greater than the value of the test statistic for the observed data is recorded. Note the point mentioned in step 5 above; the Null distribution includes the observed value of the test statistic. Denote this count as \( N \).

7. The permutation p-value is computed as

\[
p = \frac{N}{n + 1}
\]

The above description illustrates why the default number of permutations specified in Vegan functions takes values of 199 or 999 for example. Once the observed value of the test statistic is added to this number of random permutations of \( DATA \), pretty p-values are achievable because \( n + 1 \) becomes 200 or 1000, for example.

The minimum achievable p-value is

\[
p_{\text{min}} = \frac{1}{n + 1}
\]

A more common definition, in ecological circles, for \( N \) would be the number of \( x_i \) greater than or equal to \( x_0 \). The permutation p-value would then be defined as

\[
p = \frac{N + 1}{n + 1}
\]

The + 1 in the numerator of the above equation represents the observed statistic \( x_0 \). The minimum p-value would then be defined as

\[
p_{\text{min}} = \frac{0 + 1}{n + 1}
\]

However this definition discriminates between the observed statistic and the other \( x_i \). Under the Null hypothesis there is no such distinction, hence we prefer the definition used in the numbered steps above.

One cannot simply increase the number of permutations (\( n \)) to achieve a potentially lower p-value unless the number of observations available permits such a number of permutations. This is unlikely to be a problem for all but the smallest data sets when free permutation (randomisation) is valid, but in designs where \( \text{strata} \) is specified and there are a low number of observations within each level of \( \text{strata} \), there may not be as many actual permutations of the data as you might want.

It is currently the responsibility of the user to determine the total number of possible permutations for their \( DATA \). No checks are made within Vegan functions to ensure a sensible number of permutations is chosen.

Limits on the total number of permutations of \( DATA \) are more severe in temporally or spatially ordered data or experimental designs with low replication. For example, a time series of \( n = 100 \) observations has just 100 possible permutations including the observed ordering.
In situations where only a low number of permutations is possible due to the nature of DATA or the experimental design, enumeration of all permutations becomes important and achievable computationally. Currently, Vegan does not include functions to perform complete enumeration of the set of possible permutations. The next major release of Vegan will include such functionality, however.

Author(s)
Gavin Simpson

See Also
permutest, permuted.index

permutated.index2  Unrestricted and restricted permutations

Description
Unrestricted and restricted permutation designs for time series, line transects, spatial grids and blocking factors.

Usage
permuted.index2(n, control = permControl())

permControl(strata = NULL, nperm = 199, complete = FALSE,
  type = c("free", "series", "grid"),
  permute.strata = FALSE,
  maxperm = 9999, minperm = 99,
  mirror = FALSE, constant = FALSE,
  ncol = NULL, nrow = NULL,
  all.perms = NULL)

permute(i, n, control)

Arguments

n  numeric; the length of the returned vector of permuted values. Usually the number of observations under consideration.

control  a list of control values describing properties of the permutation design, as returned by a call to permControl.

strata  An integer vector or factor specifying the strata for permutation. If supplied, observations are permuted only within the specified strata.

nperm  the number of permutations.

complete  logical; should complete enumeration of all permutations be performed?
type  

permuted.index2

permute.strata

maxperm

minperm

mirror

constant

ncol, nrow

all.perms

i

Details

permuted.index2 can generate permutations for a wide range of restricted permutation schemes. A small selection of the available combinations of options is provided in the Examples section below.

Argument mirror determines whether grid or series permutations can be mirrored. Consider the sequence 1,2,3,4. The relationship between consecutive observations is preserved if we reverse the sequence to 4,3,2,1. If there is no inherent direction in your experimental design, mirrored permutations can be considered part of the Null model, and as such increase the number of possible permutations. The default is to not use mirroring so you must explicitly turn this on using mirror = TRUE in permControl.

To permute strata rather than the observations within the levels of strata, use permute.strata = TRUE. However, note that the number of observations within each level of strata must be equal!

For some experiments, such as BACI designs, one might wish to use the same permutation within each level of strata. This is controlled by argument constant. If constant = TRUE then the same permutation will be generated for each level of strata. The default is constant = FALSE.

permute is a higher level utility function for use in a loop within a function implementing a permutation test. The main purpose of permute is to return the correct permutation in each iteration of the loop, either a random permutation from the current design or the next permutation from control$all.perms if it is not NULL and control$complete is TRUE.

Value

For permuted.index2 a vector of length n containing a permutation of the observations 1, ..., n using the permutation scheme described by argument control.

For permControl a list with components for each of the possible arguments.
Note

permuted.index2 is currently used in one Vegan function: `permtest.betadisper`. Over time, the other functions that currently use the older `permuted.index` will be updated to use `permuted.index2`.

Author(s)

Gavin Simpson

See Also

`permCheck`, a utility function for checking permutation scheme described by `permControl`.

Examples

```r
set.seed(1234)

## unrestricted permutations
permuted.index2(20)

## observations represent a time series of line transect
permuted.index2(20, control = permControl(type = "series"))

## observations represent a time series of line transect
## but with mirroring allowed
permuted.index2(20, control = permControl(type = "series", mirror = TRUE))

## observations represent a spatial grid
perms <- permuted.index2(20, permControl(type = "grid",
                                        ncol = 4, nrow = 5))

## view the permutation as a grid
matrix(matrix(1:20, nrow = 5, ncol = 4)[perms], ncol = 4, nrow = 5)

## random permutations in presence of strata
block <- gl(4, 5)
permuted.index2(20, permControl(strata = block, type = "free"))

## as above but with same random permutation within strata
permuted.index2(20, permControl(strata = block, type = "free",
                                 constant = TRUE))

## time series within each level of block
permuted.index2(20, permControl(strata = block, type = "series"))

## as above, but with same permutation for each level
permuted.index2(20, permControl(strata = block, type = "series",
                                 constant = TRUE))

## spatial grids within each level of block
permuted.index2(100, permControl(strata = block, type = "grid",
                                   ncol = 5, nrow = 5))

## as above, but with same permutation for each level
permuted.index2(100, permControl(strata = block, type = "grid",
                                   ncol = 5, nrow = 5, constant = TRUE))
```
## permuting levels of block instead of observations

```r
permuted.index2(20, permControl(strata = block, type = "free",
permute.strata = TRUE))
```

## Simple function using permute() to assess significance
## of a t.test

```r
pt.test <- function(x, group, control) {
  ## function to calculate t
  t.statistic <- function(x, y) {
    m <- length(x)
    n <- length(y)
    ## means and variances, but for speed
    xbar <- .Internal(mean(x))
    ybar <- .Internal(mean(y))
    xvar <- .Internal(cov(x, NULL, 1, FALSE))
    yvar <- .Internal(cov(y, NULL, 1, FALSE))
    pooled <- sqrt(((m-1)*xvar + (n-1)*yvar) / (m+n-2))
    (xbar - ybar) / (pooled * sqrt(1/m + 1/n))
  }
  ## check the control object
  control <- permCheck(x, control)$control
  ## number of observations
  nobs <- getNumObs(x)
  ## group names
  lev <- names(table(group))
  ## vector to hold results, +1 because of observed t
  t.permu <- numeric(length = control$nperm) + 1
  ## calculate observed t
  t.permu[1] <- t.statistic(x[group == lev[1]], x[group == lev[2]])
  ## generate randomisation distribution of t
  for(i in seq_along(t.permu)) {
    ## return a permutation
    want <- permute(i, nobs, control)
    ## calculate permuted t
    t.permu[i+1] <- t.statistic(x[want][group == lev[1]],
                               x[want][group == lev[2]])
  }
  ## pval from permutation test
  pval <- sum(abs(t.permu) >= abs(t.permu[1])) / (control$nperm + 1)
  ## return value
  return(list(t.stat = t.permu[1], pval = pval))
}
```

## generate some data with slightly different means

```r
set.seed(1234)
g1 <- rnorm(20, mean = 9)
g2 <- rnorm(20, mean = 10)
dat <- c(g1, g2)
## grouping variable
gp <- gl(2, 20, labels = paste(\"Group\", 1:2))
## create the permutation design
control <- permControl(type = "free", nperm = 999)
```
## perform permutation t test
perm.val <- pt.test(dat, grp, control)
perm.val

## compare perm.val with the p-value from t.test()
t.test(dat ~ grp, var.equal = TRUE)

---

**permutest.betadisper**

*Permutation test of multivariate homogeneity of groups dispersions (variances)*

---

### Description

Implements a permutation-based test of multivariate homogeneity of group dispersions (variances) for the results of a call to `betadisper`.

### Usage

```r
## S3 method for class 'betadisper'
permutest(x, pairwise = FALSE, 
    control = permControl(nperm = 999), ...)
```

### Arguments

- **x**
  - an object of class "betadisper", the result of a call to `betadisper`.
- **pairwise**
  - logical; perform pairwise comparisons of group means?
- **control**
  - a list of control values for the permutations to replace the default values returned by the function `permControl`
- **...**
  - Arguments passed to other methods.

### Details

To test if one or more groups is more variable than the others, ANOVA of the distances to group centroids can be performed and parametric theory used to interpret the significance of F. An alternative is to use a permutation test. `permutest.betadisper` permutes model residuals to generate a permutation distribution of F under the Null hypothesis of no difference in dispersion between groups.

Pairwise comparisons of group mean dispersions can be performed by setting argument `pairwise` to `TRUE`. A classical t test is performed on the pairwise group dispersions. This is combined with a permutation test based on the t statistic calculated on pairwise group dispersions. An alternative to the classical comparison of group dispersions, is to calculate Tukey’s Honest Significant Differences between groups, via `TukeyHSD.betadisper`.

---
Value

`permutest.betadisper` returns a list of class "permutest.betadisper" with the following components:

- `tab` the ANOVA table which is an object inheriting from class "data.frame".
- `pairwise` a list with components observed and permuted containing the observed and permuted p-values for pairwise comparisons of group mean distances (dispersions or variances).
- `groups` character; the levels of the grouping factor.
- `control` a list, the result of a call to `permControl`.

Author(s)

Gavin L. Simpson

References


See Also

For the main fitting function see `betadisper`. For an alternative approach to determining which groups are more variable, see `TukeyHSD.betadisper`.

Examples

data(varespec)

## Bray-Curtis distances between samples
dis <- vegdist(varespec)

## First 16 sites grazed, remaining 8 sites ungrazed
groups <- factor(c(rep(1,16), rep(2,8)), labels = c("grazed","ungrazed"))

## Calculate multivariate dispersions
mod <- betadisper(dis, groups)
mod

## Perform test
anova(mod)

## Permutation test for F
permutest(mod, pairwise = TRUE)

## Tukey's Honest Significant Differences
(mod.HSD <- TukeyHSD(mod))
plot(mod.HSD)
plot.cca

Plot or Extract Results of Constrained Correspondence Analysis or Redundancy Analysis

Description

Functions to plot or extract results of constrained correspondence analysis (cca), redundancy analysis (rda) or constrained analysis of principal coordinates (capscale).

Usage

## S3 method for class 'cca'
plot(x, choices = c(1, 2), display = c("sp", "wa", "cn"),
     scaling = 2, type, xlim, ylim, const, ...)

## S3 method for class 'cca'
text(x, display = "sites", labels, choices = c(1, 2), scaling = 2,
    arrow.mul, head.arrow = 0.05, select, const, ...)

## S3 method for class 'cca'
points(x, display = "sites", choices = c(1, 2), scaling = 2,
    arrow.mul, head.arrow = 0.05, select, const, ...)

## S3 method for class 'cca'
scores(x, choices=c(1,2), display=c("sp","wa","cn"), scaling=2, ...)

## S3 method for class 'rda'
scores(x, choices=c(1,2), display=c("sp","wa","cn"), scaling=2, ...)

## S3 method for class 'cca'
scores(x, choices=c(1,2), display=c("sp","wa","cn"), scaling=2, const, ...)

## S3 method for class 'summary.cca'
scores(x, choices=c(1,2), display=c("sp","wa","cn"), scaling=2, const, ...)

Arguments

x, object A cca result object.

choices Axes shown.

display Scores shown. These must include some of the alternatives species or sp for species scores, sites or wa for site scores, lc for linear constraints or "LC scores", or bp for biplot arrows or cn for centroids of factor constraints instead of an arrow.

scaling Scaling for species and site scores. Either species (2) or site (1) scores are scaled by eigenvalues, and the other set of scores is left unscaled, or with 3 both are scaled symmetrically by square root of eigenvalues. Corresponding negative
values can be used in `cca` to additionally multiply results with $\sqrt{\frac{1}{1 - \lambda}}$.
This scaling is known as Hill scaling (although it has nothing to do with Hill’s rescaling of `decorana`).
With corresponding negative values in `rda`, species scores are divided by standard deviation of each species and multiplied with an equalizing constant. Unscaled raw scores stored in the result can be accessed with `scaling = 0`.

`type` Type of plot: partial match to `text` for text labels, `points` for points, and `none` for setting frames only. If omitted, `text` is selected for smaller data sets, and `points` for larger.

`xlim`, `ylim` the x and y limits (min,max) of the plot.

`labels` Optional text to be used instead of row names.

`arrow.mul` Factor to expand arrows in the graph. Arrows will be scaled automatically to fit the graph if this is missing.

`head.arrow` Default length of arrow heads.

`select` Items to be displayed. This can either be a logical vector which is TRUE for displayed items or a vector of indices of displayed items.

`const` General scaling constant to `rda` scores. The default is to use a constant that gives biplot scores, that is, scores that approximate original data (see vignette ‘decision-vegan.pdf’ with `vegandocs` for details and discussion). If `const` is a vector of two items, the first is used for species, and the second item for site scores.

`axes` Number of axes in summaries.

`digits` Number of digits in output.

`n`, `head`, `tail` Number of rows printed from the head and tail of species and site scores. Default NA prints all.

`...` Parameters passed to other functions.

### Details

Same `plot` function will be used for `cca` and `rda`. This produces a quick, standard plot with current scaling.

The `plot` function sets colours (`col`), plotting characters (`pch`) and character sizes (`cex`) to certain standard values. For a fuller control of produced plot, it is best to call `plot` with `type="none"` first, and then add each plotting item separately using `text.cca` or `points.cca` functions.

These use the default settings of standard `text` and `points` functions and accept all their parameters, allowing a full user control of produced plots.

Environmental variables receive a special treatment. With `display="bp"`, arrows will be drawn. These are labelled with `text` and unlabelled with `points`. The basic `plot` function uses a simple (but not very clever) heuristics for adjusting arrow lengths to plots, but the user can give the expansion factor in `mul.arrow`. With `display="cn"` the centroids of levels of `factor` variables are displayed (these are available only if there were factors and a formula interface was used in `cca` or `rda`). With this option continuous variables still are presented as arrows and ordered factors as arrows and centroids.
If you want to have still a better control of plots, it is better to produce them using primitive plot commands. Function scores helps in extracting the needed components with the selected scaling.

Function summary lists all scores and the output can be very long. You can suppress scores by setting axes = 0 or display = NA or display = NULL. You can display some first or last (or both) rows of scores by using head or tail or explicit print command for the summary.

Palmer (1993) suggested using linear constraints (“LC scores”) in ordination diagrams, because these gave better results in simulations and site scores (“WA scores”) are a step from constrained to unconstrained analysis. However, McCune (1997) showed that noisy environmental variables (and all environmental measurements are noisy) destroy “LC scores” whereas “WA scores” were little affected. Therefore the plot function uses site scores (“WA scores”) as the default. This is consistent with the usage in statistics and other functions in R (lda, cancor).

Value

The plot function returns invisibly a plotting structure which can be used by function identify.ordiplot to identify the points or other functions in the ordiplot family.

Note

Package ade4 has function cca which returns constrained correspondence analysis of the same class as the vegan function. If you have results of ade4 in your working environment, vegan functions may try to handle them and fail with cryptic error messages. However, there is a simple utility function ade2vegancca which tries to translate ade4 cca results to vegan cca results so that some vegan functions may work partially with ade4 objects (with a warning).

Author(s)

Jari Oksanen

See Also

cca, rda and capscale for getting something to plot, ordiplot for an alternative plotting routine and more support functions, and text, points and arrows for the basic routines.

Examples

data(dune)
data(dune.env)
mod <- cca(dune ~ A1 + Moisture + Management, dune.env)
plot(mod, type="n")
text(mod, dis="cn")
points(mod, pch=21, col="red", bg="yellow", cex=1.2)
text(mod, "species", col="blue", cex=0.8)
## Limited output of 'summary'
head(summary(mod), tail=2)
## Read description of scaling in RDA in vegan:
## Not run: vegandocs("decision")
Principal Response Curves for Treatments with Repeated Observations

Description

Principal Response Curves (PRC) are a special case of Redundancy Analysis (rda) for multivariate responses in repeated observation design. They were originally suggested for ecological communities. They should be easier to interpret than traditional constrained ordination. They can also be used to study how the effects of a factor A depend on the levels of a factor B, that is \( A + A:B \), in a multivariate response experiment.

Usage

```r
prc(response, treatment, time, ...)
## S3 method for class 'prc'
summary(object, axis = 1, scaling = 3, digits = 4, ...)
## S3 method for class 'prc'
plot(x, species = TRUE, select, scaling = 3, axis = 1, type = "l",
     xlab, ylab, ylim, lty = 1:5, col = 1:6, pch, legpos, cex = 0.8,
     ...)
```

Arguments

- `response`: Multivariate response data. Typically these are community (species) data. If the data are counts, they probably should be log transformed prior to the analysis.
- `treatment`: A factor for treatments.
- `time`: An unordered factor defining the observations times in the repeated design.
- `object, x`: An prc result object.
- `axis`: Axis shown (only one axis can be selected).
- `scaling`: Scaling of species scores, identical to the `scaling` in scores.rda.
- `digits`: Number of significant digits displayed.
- `species`: Display species scores.
- `select`: Vector to select displayed species. This can be a vector of indices or a logical vector which is TRUE for the selected species
- `type`: Type of plot: "l" for lines, "p" for points or "b" for both.
- `xlab, ylab`: Text to replace default axis labels.
- `ylim`: Limits for the vertical axis.
- `lty, col, pch`: Line type, colour and plotting characters (defaults supplied).
- `legpos`: The position of the legend. A guess is made if this is not supplied, and NA will suppress legend.
- `cex`: Character expansion for symbols and species labels.
- `...`: Other parameters passed to functions.
Details

PRC is a special case of rda with a single factor for treatment and a single factor for time points in repeated observations. In vegan, the corresponding rda model is defined as rda(response ~ treatment * time + Condition(time)). Since the time appears twice in the model formula, its main effects will be aliased, and only the main effect of treatment and interaction terms are available, and will be used in PRC. Instead of usual multivariate ordination diagrams, PRC uses canonical (regression) coefficients and species scores for a single axis. All that the current functions do is to provide a special summary and plot methods that display the rda results in the PRC fashion. The current version only works with default contrasts (contr.treatment) in which the coefficients are contrasts against the first level, and the levels must be arranged so that the first level is the control (or a baseline). If necessary, you must change the baseline level with function relevel.

Function summary prints the species scores and the coefficients. Function plot plots coefficients against time using matplot, and has similar defaults. The graph (and PRC) is meaningful only if the first treatment level is the control, as the results are contrasts to the first level when unordered factors are used. The plot also displays species scores on the right vertical axis using function linestack. Typically the number of species is so high that not all can be displayed with the default settings, but users can reduce character size or padding (air) in linestack, or select only a subset of the species. A legend will be displayed unless suppressed with legpos = NA, and the functions tries to guess where to put the legend if legpos is not supplied.

Value

The function is a special case of rda and returns its result object (see cca.object). However, a special summary and plot methods display returns differently than in rda.

Warning

The first level of treatment must be the control: use function relevel to guarantee the correct reference level. The current version will ignore user setting of contrasts and always use treatment contrasts (contr.treatment). The time must be an unordered factor.

Author(s)

Jari Oksanen and Cajo ter Braak

References


See Also

rda, anova.cca.
predict.cca

**Prediction Tools for {Constrained} Ordination (CCA, RDA, DCA, CA, PCA)**

### Description

Function `predict` can be used to find site and species scores or estimates of the response data with new data sets. Function `calibrate` estimates values of constraints with new data set. Functions `fitted` and `residuals` return estimates of response data.

### Usage

```r
## S3 method for class 'cca'
fitted(object, model = c("CCA", "CA"),
       type = c("response", "working"), ...)
## S3 method for class 'capscale'
fitted(object, model = c("CCA", "CA", "Imaginary"),
       type = c("response", "working"), ...)
## S3 method for class 'cca'
residuals(object, ...)
## S3 method for class 'cca'
predict(object, newdata, type = c("response", "wa", "sp", "lc", "working"),
        rank = "full", model = c("CCA", "CA"), scaling = FALSE, ...)
## S3 method for class 'cca'
calibrate(object, newdata, rank = "full", ...)
## S3 method for class 'cccc'
coef(object, ...)
## S3 method for class 'decorana'
predict(object, newdata, type = c("response", "sites", "species"),
        rank = 4, ...)
```
Arguments

object  
A result object from `cca`, `rda`, `capscale` or `decorana`.

model  
Show constrained ("CCA") or unconstrained ("CA") results. For `capscale` this can also be "Imaginary" for imaginary components with negative eigenvalues.

newdata  
New data frame to be used in prediction or in calibration. Usually this a new community data frame, but for `predict.cca` type = "lc" and for constrained component with type "response" and "working" it must be an environment data frame. If the original model had row or column names, then new data must contain rows or columns with the same names (row names for species scores, column names for "wa" scores and constraint names of "lc" scores). In other cases the rows or columns must match directly.

type  
The type of prediction, fitted values or residuals: "response" scales results so that the same ordination gives the same results, and "working" gives the values used internally, that is after Chi-square standardization in `cca` and scaling and centring in `rda`. In `capscale` the "response" gives the dissimilarities, and "working" the scaled scores that produce the dissimilarities as Euclidean distances. Alternative "wa" gives the site scores as weighted averages of the community data, "lc" the site scores as linear combinations of environmental data, and "sp" the species scores. In `predict.decorana` the alternatives are scores for "sites" or "species".

rank  
The rank or the number of axes used in the approximation. The default is to use all axes (full rank) of the "model" or all available four axes in `predict.decorana`.

scaling  
Scaling or predicted scores with the same meaning as in `cca`, `rda` and `capscale`.

...  
Other parameters to the functions.

Details

Function `fitted` gives the approximation of the original data matrix or dissimilarities from the ordination result either in the scale of the response or as scaled internally by the function. Function `residuals` gives the approximation of the original data from the unconstrained ordination. With argument `type = "response"` the `fitted.cca` and `residuals.cca` function both give the same marginal totals as the original data matrix, and their entries do not add up to the original data. Functions `fitted.capscale` and `residuals.capscale` give the dissimilarities with `type = "response"`, but these are not additive, but the "working" scores are additive. All variants of `fitted` and `residuals` are defined so that for `model <- cca(y ~ x), cca(fitted(mod))` is equal to constrained ordination, and `cca(residuals(mod))` is equal to unconstrained part of the ordination.

Function `predict` can find the estimate of the original data matrix or dissimilarities (`type = "response"`) with any rank. With `rank = "full"` it is identical to `fitted`. In addition, the function can find the species scores or site scores from the community data matrix for `cca` or `rda`. The function can be used with new data, and it can be used to add new species or site scores to existing ordinations. The function returns (weighted) orthonormal scores by default, and you must specify explicit `scaling` to add those scores to ordination diagrams. With `type = "wa"` the function finds the site scores from species scores. In that case, the new data can contain new sites, but species must match in the original and new data. With `type = "sp"` the function...
predict.cca

finds species scores from site constraints (linear combination scores). In that case the new data can contain new species, but sites must match in the original and new data. With `type = "lc"` the function finds the linear combination scores for sites from environmental data. In that case the new data frame must contain all constraining and conditioning environmental variables of the model formula. With `type = "response"` or `type = "working"` the new data must contain environmental variables if constrained component is desired, and community data matrix if residual or unconstrained component is desired. With these types, the function uses `newdata` to find new "lc" (constrained) or "wa" scores (unconstrained) and then finding the response or working data from these new row scores and species scores.

If a completely new data frame is created, extreme care is needed defining variables similarly as in the original model, in particular with (ordered) factors. If ordination was performed with the formula interface, the `newdata` also can be a data frame or matrix, but extreme care is needed that the columns match in the original and `newdata`.

Function `calibrate.cca` finds estimates of constraints from community ordination or "wa" scores from `cca`, `rda` and `capscale`. This is often known as calibration, bioindication or environmental reconstruction. Basically, the method is similar to projecting site scores onto biplot arrows, but it uses regression coefficients. The function can be called with `newdata` so that cross-validation is possible. The `newdata` may contain new sites, but species must match in the original and new data. The function does not work with 'partial' models with `Condition` term, and it cannot be used with `newdata` for `capscale` results. The results may only be interpretable for continuous variables.

Function `coef` will give the regression coefficients from centred environmental variables (constraints and conditions) to linear combination scores. The coefficients are for unstandardized environmental variables. The coefficients will be `NA` for aliased effects.

Function `predict.decorana` is similar to `predict.cca`. However, `type = "species"` is not available in detrended correspondence analysis (DCA), because detrending destroys the mutual reciprocal averaging (except for the first axis when rescaling is not used). Detrended CA does not attempt to approximate the original data matrix, so `type = "response"` has no meaning in detrended analysis (except with `rank = 1`).

Value

The functions return matrices, vectors or dissimilarities as is appropriate.

Author(s)

Jari Oksanen.

References


See Also

`cca`, `rda`, `capscale`, `decorana`, `vif`, `goodness.cca`
Examples

```r
data(dune)
data(dune.env)
mod <- cca(dune ~ A1 + Management + Condition(Moisture), data=dune.env)
# Definition of the concepts 'fitted' and 'residuals'
mod
cca(fitted(mod))
cca(residuals(mod))
# 'passively'.
freq <- specnumber(dune, MARGIN=2)
freq
mod <- cca(dune[, freq>1] ~ A1 + Management + Condition(Moisture), dune.env)
predict(mod, type="sp", newdata=dune[, freq==1], scaling=2)
# 'passively'.
mod <- cca(dune ~ A1 + Moisture, dune.env)
pred <- calibrate(mod)
pred
with(dune.env, plot(A1, pred[,"A1"] - A1, ylab="Prediction Error"))
bline(h=0)
```

---

**procrustes**

Procrustes Rotation of Two Configurations and PROTEST

### Description

Function `procrustes` rotates a configuration to maximum similarity with another configuration. Function `protest` tests the non-randomness ('significance') between two configurations.

### Usage

```r
procrustes(X, Y, scale = TRUE, symmetric = FALSE, scores = "sites", ...)
```

## S3 method for class 'procrustes'

```r
summary(object, digits = getOption("digits"), ...)
```

## S3 method for class 'procrustes'

```r
plot(x, kind=1, choices=c(1,2), to.target = TRUE,
     type = "p", xlab, ylab, main, ar.col = "blue", len=0.05,
     cex = 0.7, ...)
```

## S3 method for class 'procrustes'

```r
points(x, display = c("target", "rotated"), ...)
```

## S3 method for class 'procrustes'

```r
text(x, display = c("target", "rotated"), labels, ...)
```

## S3 method for class 'procrustes'

```r
lines(x, type = c("segments", "arrows"), choices = c(1, 2), ...)
```

## S3 method for class 'procrustes'

```r
residuals(object, ...)
```
## S3 method for class 'procrustes'
fitted(object, truemean = TRUE, ...)
## S3 method for class 'procrustes'
predict(object, newdata, truemean = TRUE, ...)
protest(X, Y, scores = "sites", permutations = 999, strata, ...)

### Arguments

- **X**: Target matrix
- **Y**: Matrix to be rotated.
- **scale**: Allow scaling of axes of Y.
- **symmetric**: Use symmetric Procrustes statistic (the rotation will still be non-symmetric).
- **scores**: Kind of scores used. This is the display argument used with the corresponding scores function: see scores, scores.cca and scores.cca for alternatives.
- **x, object**: An object of class procrustes.
- **digits**: Number of digits in the output.
- **kind**: For plot function, the kind of plot produced: kind = 1 plots shifts in two configurations, kind = 0 draws a corresponding empty plot, and kind = 2 plots an impulse diagram of residuals.
- **choices**: Axes (dimensions) plotted.
- **xlab, ylab**: Axis labels, if defaults unacceptable.
- **main**: Plot title, if default unacceptable.
- **display**: Show only the "target" or "rotated" matrix as points.
- **to.target**: Draw arrows to point to target.
- **type**: The type of plot drawn. In plot, the type can be "points" or "text" to select the marker for the tail of the arrow, or "none" for drawing an empty plot. In lines the type selects either arrows or line segments to connect target and rotated configuration.
- **truemean**: Use the original range of target matrix instead of centring the fitted values. Function plot.procrustes needs truemean = FALSE.
- **newdata**: Matrix of coordinates to be rotated and translated to the target.
- **permutations**: Number of permutation to assess the significance of the symmetric Procrustes statistic.
- **strata**: An integer vector or factor specifying the strata for permutation. If supplied, observations are permuted only within the specified strata.
- **ar.col**: Arrow colour.
- **len**: Width of the arrow head.
- **labels**: Character vector of text labels. Rownames of the result object are used as default.
- **cex**: Character expansion for points or text.
- **...**: Other parameters passed to functions. In procrustes and protest parameters are passed to scores, in graphical functions to underlying graphical functions.
Details

Procrustes rotation rotates a matrix to maximum similarity with a target matrix minimizing sum of squared differences. Procrustes rotation is typically used in comparison of ordination results. It is particularly useful in comparing alternative solutions in multidimensional scaling. If `scale=FALSE`, the function only rotates matrix $Y$. If `scale=TRUE`, it scales linearly configuration $Y$ for maximum similarity. Since $Y$ is scaled to fit $X$, the scaling is non-symmetric. However, with `symmetric=TRUE`, the configurations are scaled to equal dispersions and a symmetric version of the Procrustes statistic is computed.

Instead of matrix, $X$ and $Y$ can be results from an ordination from which `scores` can extract results. Function `procrustes` passes extra arguments to `scores`, `scores.cca` etc. so that you can specify arguments such as `scaling`.

Function `plot` plots a `procrustes` object and returns invisibly an `ordiplot` object so that function `identify.ordiplot` can be used for identifying points. The items in the `ordiplot` object are called `heads` and `points` with `kind=1` (ordination diagram) and `sites` with `kind=2` (residuals). In ordination diagrams, the arrow heads point to the target configuration if `to.target = TRUE`, and to rotated configuration if `to.target = FALSE`. Target and original rotated axes are shown as cross hairs in two-dimensional Procrustes analysis, and with a higher number of dimensions, the rotated axes are projected onto plot with their scaled and centred range. Function `plot` passes parameters to underlying plotting functions. For full control of plots, you can draw the axes using `plot` with `kind = 0`, and then add items with `points` or `lines`. These functions pass all parameters to the underlying functions so that you can select the plotting characters, their size, colours etc., or you can select the width, colour and type of line `segments` or arrows, or you can select the orientation and head width of `arrows`.

Function `residuals` returns the pointwise residuals, and `fitted` the fitted values, either centred to zero mean (if `true.mean=FALSE`) or with the original scale (these hardly make sense if `symmetric = TRUE`). In addition, there are `summary` and `print` methods.

If matrix $X$ has a lower number of columns than matrix $Y$, then matrix $X$ will be filled with zero columns to match dimensions. This means that the function can be used to rotate an ordination configuration to an environmental variable (most practically extracting the result with the `fitted` function). Function `predict` can be used to add new rotated coordinates to the target. The `predict` function will always translate coordinates to the original non-centred matrix. The function cannot be used with `newdata` for symmetric analysis.

Function `protest` calls `procrustes(...)`, `symmetric = TRUE`) repeatedly to estimate the ‘significance’ of the Procrustes statistic. Function `protest` uses a correlation-like statistic derived from the symmetric Procrustes sum of squares $ss$ as $r = \sqrt{(1 - ss)}$, and sometimes called $m_{12}$. Function `protest` has own `print` method, but otherwise uses `procrustes` methods. Thus `plot` with a `protest` object yields a “Procrustean superimposition plot.”

Value

Function `procrustes` returns an object of class `procrustes` with items. Function `protest` inherits from `procrustes`, but amends that with some new items:

- **Yrot** Rotated matrix $Y$.
- **X** Target matrix.
- **ss** Sum of squared differences between $X$ and $Yrot$. 


rotation Orthogonal rotation matrix.
translation Translation of the origin.
scale Scaling factor.
xmean The centroid of the target.
symmetric Type of ss statistic.
call Function call.
t0 This and the following items are only in class protest: Procrustes correlation from non-permuted solution.
t Procrustes correlations from permutations.
signif ‘Significance’ of t
permutations Number of permutations.
strata The name of the stratifying variable.
stratum.values Values of the stratifying variable.

Note
The function protest follows Peres-Neto & Jackson (2001), but the implementation is still after Mardia et al. (1979).

Author(s)
Jari Oksanen

References

See Also
isoMDS, initMDS for obtaining objects for procrustes, and mantel for an alternative to protest without need of dimension reduction.

Examples
data(varespec)
vare.dist <- vegdist(wisconsin(varespec))
library(MASS) ## isoMDS
mds.null <- isoMDS(vare.dist, tol=1e-7)
mds.alt <- isoMDS(vare.dist, initMDS(vare.dist), maxit=200, tol=1e-7)
vare.proc <- procrustes(mds.alt, mds.null)
vare.proc
summary(vare.proc)
plot(vare.proc)
plot(vare.proc, kind=2)
residuals(vare.proc)
Description

The data are log transformed abundances of aquatic invertebrate in twelve ditches studied in eleven times before and after an insecticide treatment.

Usage

data(pyrifos)

Format

A data frame with 132 observations on the log-transformed abundances of 178 species. There are only twelve sites (ditches, mesocosms), but these were studied repeatedly in eleven occasions. The treatment levels, treatment times, or ditch ID’s are not in the data frame, but the data are very regular, and the example below shows how to obtain these external variables.

Details

This data set was obtained from an experiment in outdoor experimental ditches. Twelve mesocosms were allocated at random to treatments; four served as controls, and the remaining eight were treated once with the insecticide chlorpyrifos, with nominal dose levels of 0.1, 0.9, 6, and 44 µg/ L in two mesocosms each. The example data set invertebrates. Sampling was done 11 times, from week -4 pre-treatment through week 24 post-treatment, giving a total of 132 samples (12 mesocosms times 11 sampling dates), see van den Brink & ter Braak (1999) for details. The data set contains only the species data, but the example below shows how to obtain the treatment, time and ditch ID variables.

Source

CANOCO 4 example data, with the permission of Cajo J. F. ter Braak.

References


Examples

data(pyrifos)
ditch <- gl(12, 1, length=132)
week <- gl(11, 12, labels=c(-4, -1, 0.1, 1, 2, 4, 8, 12, 15, 19, 24))
dose <- factor(rep(c(0.1, 0, 0.9, 0, 44, 6, 0.1, 44, 0.9, 0, 6), 11))
Description

Functions construct rank – abundance or dominance / diversity or Whittaker plots and fit broken-stick, pre-emption, log-Normal, Zipf and Zipf-Mandelbrot models of species abundance.

Usage

## S3 method for class 'data.frame'
radfit(df, ...)
## S3 method for class 'radfit.frame'
plot(x, order.by, BIC = FALSE, model, legend = TRUE,
as.table = TRUE, ...)
## Default S3 method:
radfit(x, ...)
## S3 method for class 'radfit'
plot(x, BIC = FALSE, legend = TRUE, ...)
radlattice(x, BIC = FALSE, ...)  
rad.null(x, family=poisson, ...)
rad.preempt(x, family = poisson, ...)
rad.lognormal(x, family = poisson, ...)
rad.zipf(x, family = poisson, ...)
rad.zipfbrot(x, family = poisson, ...)
## S3 method for class 'radline'
plot(x, xlab = "Rank", ylab = "Abundance", type = "b", ...)
## S3 method for class 'radline'
lines(x, ...)
## S3 method for class 'radline'
bpoints(x, ...)
## S3 method for class 'rad'
plot(x, xlab = "Rank", ylab = "Abundance", log = "y", ...)

Arguments

df  
Data frame where sites are rows and species are columns.

x  
A vector giving species abundances in a site, or an object to be plotted.

order.by  
A vector used for ordering sites in plots.

BIC  
Use Bayesian Information Criterion, BIC, instead of Akaike's AIC. The penalty for a parameter is \( k = \log(S) \) where \( S \) is the number of species, whereas AIC uses \( k = 2 \).

model  
Show only the specified model. If missing, AIC is used to select the model. The model names (which can be abbreviated) are Preemption, Lognormal, Veiled.LN, Zipf, Mandelbrot.
Add legend of line colours.

Arrange panels starting from upper left corner (passed to `xyplot`).

Error distribution (passed to `glm`). All alternatives accepting `link = "log"` in `family` can be used, although not all make sense.

Labels for \(x\) and \(y\) axes.

Type of the plot, "b" for plotting both observed points and fitted lines, "p" for only points, "l" for only fitted lines, and "n" for only setting the frame.

Use logarithmic scale for given axis. The default `log = "y"` gives the traditional plot in community ecology where the pre-emption model is a straight line, and with `log = "xy"` Zipf model is a straight line. With `log = "n"` both axes are in the original arithmetic scale.

Other parameters to functions.

Details

Rank – Abundance Dominance (RAD) or Dominance/Diversity plots (Whittaker 1965) display logarithmic species abundances against species rank order. These plots are supposed to be effective in analysing types of abundance distributions in communities. These functions fit some of the most popular models mainly following Wilson (1991). Function `as.rad` constructs observed RAD data. Functions `rad.XXXX` (where `XXXX` is a name) fit the individual models, and function `radfit` fits all models. The argument of the function `radfit` can be either a vector for a single community or a data frame where each row represents a distinct community. All these functions have their own `plot` functions. When the argument is a data frame, `plot` uses `Lattice` graphics, and other `plot` functions use ordinary graphics. The ordinary graphics functions return invisibly an `ordiplot` object for observed points, and function `identify.ordiplot` can be used to label selected species. The most complete control of graphics can be achieved with `rad.XXXX` methods which have `points` and `lines` functions to add observed values and fitted models into existing graphs. Alternatively, `radlattice` uses `Lattice` graphics to display each `radfit` model in a separate panel together with their AIC or BIC values.

Function `rad.null` fits a brokenstick model where the expected abundance of species at rank \(r\) is \(a_r = (J/S) \sum_{x=r}^S (1/x)\) (Pielou 1975), where \(J\) is the total number of individuals (site total) and \(S\) is the total number of species in the community. This gives a Null model where the individuals are randomly distributed among observed species, and there are no fitted parameters. Function `rad.preempt` fits the niche preemption model, a.k.a. geometric series or Motomura model, where the expected abundance \(a\) of species at rank \(r\) is \(a_r = J\alpha(1-\alpha)^{r-1}\). The only estimated parameter is the preemption coefficient \(\alpha\) which gives the decay rate of abundance per rank. The niche preemption model is a straight line in a RAD plot. Function `rad.lognormal` fits a log-Normal model which assumes that the logarithmic abundances are distributed Normally, or \(a_r = \exp(\log \mu + \log \sigma N)\), where \(N\) is a Normal deviate. Function `rad.zipf` fits the Zipf model \(a_r = Jp_1r^\gamma\) where \(p_1\) is the fitted proportion of the most abundant species, and \(\gamma\) is a decay coefficient. The Zipf – Mandelbrot model (`rad.zipfbrot`) adds one parameter: \(a_r = Jc(r+\beta)^\gamma\) after which \(p_1\) of the Zipf model changes into a meaningless scaling constant \(c\). There are grand narratives about ecological mechanisms behind each model (Wilson 1991), but several alternative and contrasting mechanisms can produce similar models and a good fit does not imply a specific mechanism.

Log-Normal and Zipf models are generalized linear models (`glm`) with logarithmic link function. Zipf-Mandelbrot adds one nonlinear parameter to the Zipf model, and is fitted using `nlm` for the
nonlinear parameter and estimating other parameters and log-Likelihood with glm. Pre-emption model is fitted as purely nonlinear model. There are no estimated parameters in the Null model. The default family is poisson which is appropriate only for genuine counts (integers), but other families that accept link = "log" can be used. Family Gamma may be appropriate for abundance data, such as cover. The “best” model is selected by AIC. Therefore “quasi” families such as quasipoisson cannot be used: they do not have AIC nor log-Likelihood needed in non-linear models.

Value

Function rad.XXXX will return an object of class radline, which is constructed to resemble results of glm and has many (but not all) of its components, even when only nlm was used in fitting. At least the following glm methods can be applied to the result: fitted, residuals.glm with alternatives "deviance" (default), "pearson", "response", function coef, AIC, extractAIC, and deviance. Function radfit applied to a vector will return an object of class radfit with item y for the constructed RAD, item family for the error distribution, and item models containing each radline object as an item. In addition, there are special AIC, coef and fitted implementations for radfit results. When applied to a data frame radfit will return an object of class radfit.frame which is a list of radfit objects; function summary can be used to display the results for individual radfit objects. The functions are still preliminary, and the items in the radline objects may change.

Note

The RAD models are usually fitted for proportions instead of original abundances. However, nothing in these models seems to require division of abundances by site totals, and original observations are used in these functions. If you wish to use proportions, you must standardize your data by site totals, e.g. with decostand and use appropriate family such as Gamma.

The lognormal model is fitted in a standard way, but I do think this is not quite correct – at least it is not equivalent to fitting Normal density to log abundances like originally suggested (Preston 1948).

Some models may fail. In particular, estimation of the Zipf-Mandelbrot model is difficult. If the fitting fails, NA is returned.

Wilson (1991) defined preemption model as \( r = J p_1 (1 - \alpha)^{r-1} \), where \( p_1 \) is the fitted proportion of the first species. However, parameter \( p_1 \) is completely defined by \( \alpha \) since the fitted proportions must add to one, and therefore I handle preemption as a one-parameter model.

Veiled log-Normal model was included in earlier releases of this function, but it was removed because it was flawed: an implicit veil line also appears in the ordinary log-Normal. The latest release version with rad.veil was 1.6-10.

Author(s)

Jari Oksanen

References


**See Also**

`fisherfit` and `prestonfit`. An alternative approach is to use `qqnorm` or `qqplot` with any distribution. For controlling graphics: `Lattice`, `xyplot`, `lset`.

**Examples**

```r
data(BCI)
mod <- rad.lognormal(BCI[1,])
mod
plot(mod)
mod <- radfit(BCI[1,])
## Standard plot overlaid for all models
## Pre-emption model is a line
plot(mod)
## log for both axes: Zipf model is a line
plot(mod, log = "xy")
## Lattice graphics separately for each model
radlattice(mod)
# Take a subset of BCI to save time and nerves
mod <- radfit(BCI[2:5,])
mod
plot(mod, pch=".")
```

---

**rankindex**

*Compares Dissimilarity Indices for Gradient Detection*

**Description**

Rank correlations between dissimilarity indices and gradient separation.

**Usage**

```r
rankindex(grad, veg, indices = c("euc", "man", "gow", "bra", "kul"),
           stepacross = FALSE, method = "spearman", ...)
```

**Arguments**

- `grad`: The gradient variable or matrix.
- `veg`: The community data matrix.
- `indices`: Dissimilarity indices compared, partial matches to alternatives in `vegdist`. Alternatively, it can be a (named) list of functions returning objects of class 'dist'.
Use *stepacross* to find a shorter path dissimilarity. The dissimilarities for site pairs with no shared species are set NA using *no.shared* so that indices with no fixed upper limit can also be analysed.

**method**  
Correlation method used.

...  
Other parameters to *stepacross*.

**Details**

A good dissimilarity index for multidimensional scaling should have a high rank-order similarity with gradient separation. The function compares most indices in *vegdist* against gradient separation using rank correlation coefficients in *cor.test*. The gradient separation between each point is assessed as Euclidean distance for continuous variables, and as Gower metric for mixed data using function *daisy* when *grad* has factors.

The *indices* argument can accept any dissimilarity indices besides the ones calculated by the *vegdist* function. For this, the argument value should be a (possibly named) list of functions. Each function must return a valid ‘dist’ object with dissimilarities, similarities are not accepted and should be converted into dissimilarities beforehand.

**Value**

Returns a named vector of rank correlations.

**Note**

There are several problems in using rank correlation coefficients. Typically there are very many ties when \(n(n-1)/2\) gradient separation values are derived from just \(n\) observations. Due to floating point arithmetics, many tied values differ by machine epsilon and are arbitrarily ranked differently by *rank* used in *cor.test*. Two indices which are identical with certain transformation or standardization may differ slightly (magnitude \(10^{-15}\)) and this may lead into third or fourth decimal instability in rank correlations. Small differences in rank correlations should not be taken too seriously. Probably this method should be replaced with a sounder method, but I do not yet know which… You may experiment with *mantel*, *anosim* or even *protest*.

Earlier version of this function used *method* = "kendall", but that is far too slow in large data sets.

The functions returning dissimilarity objects should be self contained, because the ... argument passes additional parameters to *stepacross* and not to the functions supplied via the *indices* argument.

**Author(s)**

Jari Oksanen, with additions from Peter Solymos

**References**

See Also

vegdist, stepacross, no.shared, isoMDS, cor, Machine, and for alternatives anosim, mantel and protest.

Examples

data(varespec)
data(varechem)
## The next scales all environmental variables to unit variance.
## Some would use PCA transformation.
rankindex(scale(varechem), varespec)
rkindex(scale(varechem), wisconsin(varespec))
## Using non vegdist indices as functions
funs <- list(Manhattan=function(x) dist(x, "manhattan"),
             Gower=function(x) cluster:::daisy(x, "gower"),
             Ochiai=function(x) designdist(x, "1-J/sqrt(A*B)"))
rkindex(scale(varechem), varespec, funs)

read.cep  

Reads a CEP (Canoco) data file

Description

read.cep reads a file formatted by relaxed strict CEP format used by Canoco software, among others.

Usage

read.cep(file, maxdata=10000, positive=TRUE, trace=FALSE, force=FALSE)

Arguments

file File name (character variable).
maxdata Maximum number of non-zero entries.
positive Only positive entries, like in community data.
trace Work verbosely.
force Run function, even if R refuses first.

Details

Cornell Ecology Programs (CEP) introduced several data formats designed for punched cards. One of these was the ‘condensed strict’ format which was adopted by popular software DECORANA and TWINSPAN. Later, Cajo ter Braak wrote Canoco based on DECORANA, where he adopted the format, but relaxed it somewhat (that’s why I call it a ‘relaxed strict’ format). Further, he introduced a more ordinary ‘free’ format, and allowed the use of classical Fortran style ‘open’ format with fixed field widths. This function should be able to deal with all these Canoco formats, whereas it cannot read many of the traditional CEP alternatives.

All variants of CEP formats have:
• Two or three title cards, most importantly specifying the format (or word FREE) and the number of items per record (number of species and sites for FREE format).
• Data in one of three accepted formats:
  1. Condensed format: First number on the line is the site identifier, and it is followed by pairs (‘couplets’) of numbers identifying the species and its abundance (an integer and a floating point number).
  2. Open Fortran format, where the first number on the line must be the site number, followed by abundance values in fields of fixed widths. Empty fields are interpreted as zeros.
  3. ‘Free’ format, where the numbers are interpreted as abundance values. These numbers must be separated by blank space, and zeros must be written as zeros.
• Species and site names, given in Fortran format (10A8): Ten names per line, eight columns for each.

With option positive = TRUE the function removes all lines and columns with zero or negative marginal sums. In community data with only positive entries, this removes empty sites and species. If data entries can be negative, this ruins data, and such data sets should be read in with option positive = FALSE.

Value

Returns a data frame, where columns are species and rows are sites. Column and row names are taken from the CEP file, and changed into unique R names by make.names after stripping the blanks.

Note

The function relies on smooth linking of Fortran file IO in R session. This is not guaranteed to work, and therefore the function may not work in your system, but it can crash the R session. Therefore the default is that the function does not run. If you still want to try:

1. Save your session
2. Run read.cep() with switch force=TRUE

If you transfer files between operating systems or platforms, you should always check that your file is formatted to your current platform. For instance, if you transfer files from Windows to Linux, you should change the files to unix format, or your session may crash when Fortran program tries to read the invisible characters that Windows uses at the end of each line.

If you compiled vegan using gfortran, the input is probably corrupted. You either should compile vegan with other FORTRAN compilers or not to use read.cep. The problems still persist in gfortran 4.01.

Author(s)

Jari Oksanen

References

Examples

```r
## Provided that you have the file `dune.spe'
## Not run:
theclassic <- read.cep("dune.spe", force=T)
## End(Not run)
```

### renyi

**Rényi and Hill Diversities and Corresponding Accumulation Curves**

**Description**

Function *renyi* find Rényi diversities with any scale or the corresponding Hill number (Hill 1973). Function *renyiaccum* finds these statistics with accumulating sites.

**Usage**

```r
renyi(x, scales = c(0, 0.25, 0.5, 1, 2, 4, 8, 16, 32, 64, Inf), hill = FALSE)
## S3 method for class 'renyi'
plot(x, ...)
renyiaccum(x, scales = c(0, 0.5, 1, 2, 4, Inf), permutations = 100,
          raw = FALSE, ...)
## S3 method for class 'renyiaccum'
plot(x, what = c("mean", "Qnt 0.025", "Qnt 0.975"), type = "l",
     ...)
## S3 method for class 'renyiaccum'
persp(x, theta = 220, col = heat.colors(100), zlim, ...)
rgl.renyiaccum(x, rgl.height = 0.2, ...)
```

**Arguments**

- **x** Community data matrix or plotting object.
- **scales** Scales of Rényi diversity.
- **hill** Calculate Hill numbers.
- **permutations** Number of random permutations in accumulating sites.
- **raw** if FALSE then return summary statistics of permutations, and if TRUE then returns the individual permutations.
- **what** Items to be plotted.
- **type** Type of plot, where type = "1" means lines.
- **theta** Angle defining the viewing direction (azimuthal) in `persp`.
- **col** Colours used for surface. Single colour will be passed on, and vector colours will be selected by the midpoint of a rectangle in `persp`.
- **zlim** Limits of vertical axis.
- **rgl.height** Scaling of vertical axis.
- **...** Other arguments which are passed to *renyi* and to graphical functions.
Details

Common diversity indices are special cases of Rényi diversity

\[ H_a = \frac{1}{1-a} \log \sum p_i^a \]

where \( a \) is a scale parameter, and Hill (1975) suggested to use so-called “Hill numbers” defined as \( N_a = \exp(H_a) \). Some Hill numbers are the number of species with \( a = 0 \), \( \exp(H') \) or the exponent of Shannon diversity with \( a = 1 \), inverse Simpson with \( a = 2 \) and \( 1/\max(p_i) \) with \( a = \infty \). According to the theory of diversity ordering, one community can be regarded as more diverse than another only if its Rényi diversities are all higher (Tóthmérész 1995).

The plot method for renyi uses lattice graphics, and displays the diversity values against each scale in separate panel for each site together with minimum, maximum and median values in the complete data.

Function renyiaccum is similar to specaccum but finds Rényi or Hill diversities at given scales for random permutations of accumulated sites. Its plot function uses lattice function xyplot to display the accumulation curves for each value of scales in a separate panel. In addition, it has a persp method to plot the diversity surface against scale and number and sites. Dynamic graphics with rgl.renyiaccum use rgl package, and produces similar surface as persp with a mesh showing the empirical confidence levels.

Value

Function renyi returns a data frame of selected indices. Function renyiaccum with argument raw = FALSE returns a three-dimensional array, where the first dimension are the accumulated sites, second dimension are the diversity scales, and third dimension are the summary statistics mean, stdev, min, max, Qnt 0.025 and Qnt 0.975. With argument raw = TRUE the statistics on the third dimension are replaced with individual permutation results.

Author(s)

Roeland Kindt <r.kindt@cgiar.org> and Jari Oksanen

References

http://www.worldagroforestry.org/treesandmarkets/tree_diversity_analysis.asp


See Also

diversity for diversity indices, and specaccum for ordinary species accumulation curves, and xyplot, persp and rgl for controlling graphics.
Examples

data(BCI)
i <- sample(nrow(BCI), 12)
mod <- renyi(BCI[i,])
plot(mod)
mod <- renyiaccum(BCI[i,])
plot(mod, as.table=TRUE, col = c(1, 2, 2))
persp(mod)

---

RsquareAdj  Adjusted R-square

Description

The functions finds the adjusted R-square.

Usage

## Default S3 method:
RsquareAdj(x, n, m, ...)
## S3 method for class 'rda'
RsquareAdj(x, ...)

Arguments

x

Unadjusted R-squared or an object from which the terms for evaluation or adjusted R-squared can be found.

n, m

Number of observations and number of degrees of freedom in the fitted model.

...  Other arguments (ignored).

Details

The default method finds the adjusted R-squared from the unadjusted R-squared, number of observations, and number of degrees of freedom in the fitted model. The specific methods find this information from the fitted result object. There are specific methods for rda, cca, lm and glm. Adjusted, or even unadjusted, R-squared may not be available in some cases, and then the functions will return NA. There is no adjusted R-squared in cca, in partial rda, and R-squared values are available only for gaussian models in glm.

Value

The functions return a list of items r.squared and adj.r.squared.

References

see Also

`varpart` uses `RsquareAdj`.

Examples

data(mite)
data(mite.env)
# rda
m <- rda(decostand(mite, "hell") ~ ., mite.env)
RsquareAdj(m)
# default method
RsquareAdj(0.8, 20, 5)

Description

Function to access either species or site scores for specified axes in some ordination methods.

Usage

```r
## Default S3 method:
scores(x, choices, display=c("sites", "species"), ...)
```

Arguments

- `x`  
  An ordination result.
- `choices`  
  Ordination axes. If missing, default method returns all axes.
- `display`  
  Partial match to access scores for sites or species.
- `...`  
  Other parameters (unused).

Details

Function `scores` is a generic method in `vegan`. Several `vegan` functions have their own `scores` methods with their own defaults and with some new arguments. This help page describes only the default method. For other methods, see, e.g., `scores.cca`, `scores.rda`, `scores.decorana`.

All `vegan` ordination functions should have a `scores` method which should be used to extract the scores instead of directly accessing them. Scaling and transformation of scores should also happen in the `scores` function. If the `scores` function is available, the results can be plotted using `ordiplot`, `ordixyplot` etc., and the ordination results can be compared in `procrustes` analysis.

The `scores.default` function is used to extract scores from non-`vegan` ordination results. Most standard ordination methods of libraries `mva`, `multiv` and `MASS` do not have a specific class, and no specific method can be written for them. However, `scores.default` guesses where some commonly used functions keep their site scores and possible species scores.
If \( x \) is a matrix, `scores.default` returns the chosen columns of that matrix, ignoring whether species or sites were requested (do not regard this as a bug but as a feature, please). Currently the function seems to work at least for `isoMDS`, `prcomp`, `princomp` and some `ade4` objects. It may work in other cases or fail mysteriously.

**Value**

The function returns a matrix of scores.

**Author(s)**

Jari Oksanen

**See Also**

`scores.cca`, `scores.decorana`. These have somewhat different interface – `scores.cca` in particular – but all work with keywords `display="sites"` and return a matrix. However, they may also return a list of matrices, and some other `scores` methods will have quite different arguments.

**Examples**

```r
data(varespec)
vare.pca <- prcomp(varespec)
scores(vare.pca, choices=c(1,2))
```

---

**Description**

Screeplot methods for plotting variances of ordination axes/components and overlaying broken stick distributions. Also, provides alternative screeplot methods for `princomp` and `prcomp`.

**Usage**

```r
## S3 method for class 'cca'
screepplot(x, bstick = FALSE, type = c("barplot", "lines"),
          npcs = min(10, if (is.null(x$CCA)) x$CA$rank else x$CCA$rank),
           ptype = "o", bst.col = "red", bst.lty = "solid",
           xlab = "Component", ylab = "Inertia",
           main = deparse(substitute(x)), legend = bstick,
           ...)

## S3 method for class 'decorana'
screepplot(x, bstick = FALSE, type = c("barplot", "lines"),
           npcs = 4,
           ptype = "o", bst.col = "red", bst.lty = "solid",
           ...)
```
xlab = "Component", ylab = "Inertia",
main = deparse(substitute(x)),
...)

## S3 method for class 'prcomp'
screepplot(x, bstick = FALSE, type = c("barplot", "lines"),
npcs = min(10, length(x$sdev)),
ptype = "o", bst.col = "red", bst.lty = "solid",
xlab = "Component", ylab = "Inertia",
main = deparse(substitute(x)), legend = bstick,
...)

## S3 method for class 'princomp'
screepplot(x, bstick = FALSE, type = c("barplot", "lines"),
npcs = min(10, length(x$sdev)),
ptype = "o", bst.col = "red", bst.lty = "solid",
xlab = "Component", ylab = "Inertia",
main = deparse(substitute(x)), legend = bstick,
...)

bstick(n, ...)

## Default S3 method:
bstick(n, tot.var = 1, ...)

## S3 method for class 'cca'
bstick(n, ...)

## S3 method for class 'prcomp'
bstick(n, ...)

## S3 method for class 'princomp'
bstick(n, ...)

## S3 method for class 'decorana'
bstick(n, ...)

Arguments

x an object from which the component variances can be determined.
bstick logical; should the broken stick distribution be drawn?
npcs the number of components to be plotted.
type the type of plot.
ptype if type == "lines" or bstick = TRUE, a character indicating the type of plotting used for the lines; actually any of the types as in plot.default.
bst.col, bst.lty the colour and line type used to draw the broken stick distribution.
xlab, ylab, main
  graphics parameters.

legend
  logical; draw a legend?

n
  an object from which the variances can be extracted or the number of variances
  (components) in the case of bstick.default.

tot.var
  the total variance to be split.

... arguments passed to other methods.

Details

The functions provide screeplots for most ordination methods in vegan and enhanced versions with
broken stick for prcomp and princomp.

Function bstick gives the brokenstick values which are ordered random proportions, defined as
\[ p_i = \left( \frac{\text{tot}}{n} \right) \sum_{x=1}^{n} \left( \frac{1}{x} \right) \] (Legendre & Legendre 1998), where \( \text{tot} \) is the total and \( n \) is the number
of brokenstick components (cf. radfit). Broken stick has been recommended as a stopping rule
in principal component analysis (Jackson 1993): principal components should be retained as long
as observed eigenvalues are higher than corresponding random broken stick components.

The bstick function is generic. The default needs the number of components and the total,
and specific methods extract this information from ordination results. There also is a bstick
method for cca. However, the broken stick model is not strictly valid for correspondence analysis
(CA), because eigenvalues of CA are defined to be \( \leq 1 \), whereas brokenstick components have
no such restrictions. The brokenstick components are not available for decorana where the sum
of eigenvalues (total inertia) is unknown, and the eigenvalues of single axes are not additive in
detrended analysis.

Value

Function screeplot draws a plot on the currently active device, and returns invisibly the xy.coords
of the points or bars for the eigenvalues.

Function bstick returns a numeric vector of broken stick components.

Note

Function screeplot is generic from R version 2.5.0. In these versions you can use plain screeplot
command without suffices cca, prcomp etc.

Author(s)

Gavin L. Simpson

References

and statistical approaches. Ecology 74, 2204–2214.

## simulate.rda

**Simulate Responses with Gaussian Error or Permuted Residuals for Constrained Ordination**

### Description

Function simulates a response data frame so that it adds Gaussian error to the fitted responses of Redundancy Analysis (*rda*), Constrained Correspondence Analysis (*cca*) or distance-based RDA (*capscale*). The function is a special case of generic *simulate*, and works similarly as *simulate.lm*.

### Usage

```r
## S3 method for class 'rda'
simulate(object, nsim = 1, seed = NULL, indx = NULL, rank = "full", ...)
```

### Arguments

- `object` an object representing a fitted *rda* model.
- `nsim` number of response vectors to simulate. (Not yet used, and values above 1 will give an error).
- `seed` an object specifying if and how the random number generator should be initialized ('seeded'). See *simulate* for details.
- `indx` Index of residuals added to the fitted values, such as produced by *permuted.index*, *permuted.index2* or *sample*. The index can have duplicate entries so that bootstrapping is allowed. If null, parametric simulation is used and Gaussian error is added to the fitted values.
- `rank` The rank of the constrained component: passed to *predict.rda* or *predict.cca*.
- `...` additional optional arguments (ignored).

### Examples

```r
data(varespec)
vare.pca <- rda(varespec, scale = TRUE)
bstick(vare.pca)
screepplot(vare.pca, bstick = TRUE, type = "lines")
```
Details

The implementation follows "lm" method of `simulate`, and adds Gaussian (Normal) error to the fitted values (fitted.rda) using function `rnorm`. The standard deviations are estimated independently for each species (column) from the residuals after fitting the constraints. Alternatively, the function can take a permutation index that is used to add permuted residuals (unconstrained component) to the fitted values. Raw data are used in rda. Internal Chi-square transformed data in cca within the function, but the returned data frame is similar to the original input data. The simulation is performed on internal metric scaling data in capscale, but the function returns the Euclidean distances calculated from the simulated data. The simulation uses only the real components, and the imaginary dimensions are ignored.

Value

Returns a data frame with similar additional arguments on random number seed as `simulate`.

Author(s)

Jari Oksanen

See Also

`simulate` for the generic case and for lm objects. Functions fitted.rda and fitted.cca return fitted values without the error component.

Examples

data(dune)
data(dune.env)
mod <- rda(dune ~ Moisture + Management, dune.env)
## One simulation
update(mod, simulate(mod) ~ .)
## An impression of confidence regions of site scores
plot(mod, display="sites")
for (i in 1:5) lines(procrustes(mod, update(mod, simulate(mod) ~ .)), col="blue")

sipoo

*Birds in the Archipelago of Sipoo (Sibbo)*

Description

Land birds on islands covered by coniferous forest in the Sipoo archipelago, southern Finland (land-bridge/ oceanic distinction unclear from source).

Usage

data(sipoo)
Format

A data frame with 18 sites and 50 species (Simberloff & Martin, 1991, Appendix 3). The species are referred by 4+4 letter abbreviation of their Latin names (but using five letters in two species names to make these unique). The example gives the areas of the studies islands in hectares.

Source

http://www.aics-research.com/nested/

References


Examples

data(sipoo)
## Areas of the islands in hectares
sipoo.area <- c(1.1, 2.1, 2.2, 3.1, 3.5, 5.8, 6, 6.1, 6.5, 11.4, 13, 14.5, 16.1, 17.5, 28.7, 40.5, 104.5, 233)

---

**spantree**  
*Minimum Spanning Tree*

Description

Function **spantree** finds a minimum spanning tree connecting all points, but disregarding dissimilarities that are at or above the threshold or NA.

Usage

spantree(d, toolong = 0)  
## S3 method for class 'spantree'  
cophenetic(x)  
spandepth(x)  
## S3 method for class 'spantree'  
plot(x, ord, cex = 0.7, type = "p", labels, dlim,  
     FUN = sammon, ...)  
## S3 method for class 'spantree'  
lines(x, ord, display="sites", ...)

Arguments

d  
Dissimilarity data inheriting from class *dist* or a an object, such as a matrix, that can be converted to a dissimilarity matrix. Functions *vegdist* and *dist* are some functions producing suitable dissimilarity data.
toolong  Shortest dissimilarity regarded as NA. The function uses a fuzz factor, so that dissimilarities close to the limit will be made NA, too. If toolong = 0 (or negative), no dissimilarity is regarded as too long.
x    A spantree result object.
ord  An ordination configuration, or an ordination result known by scores.
cex  Character expansion factor.
type Observations are plotted as points with type="p" or type="b", or as text label with type="t". The tree (lines) will always be plotted.
labels Text used with type="t" or node names if this is missing.
dlim  A ceiling value used to highest cophenetic dissimilarity.
FUN  Ordination function to find the configuration from cophenetic dissimilarities.
display Type of scores used for ord.
... Other parameters passed to functions.

Details

Function spantree finds a minimum spanning tree for dissimilarities (there may be several minimum spanning trees, but the function finds only one). Dissimilarities at or above the threshold toolong and NAs are disregarded, and the spanning tree is found through other dissimilarities. If the data are disconnected, the function will return a disconnected tree (or a forest), and the corresponding link is NA. Connected subtrees can be identified using distconnected.

Function cophenetic finds distances between all points along the tree segments. Function spandepth returns the depth of each node. The nodes of a tree are either leaves (with one link) or internal nodes (more than one link). The leaves are recursively removed from the tree, and the depth is the layer at which the leaf was removed. In disconnected spantree object (in a forest) each tree is analysed separately and disconnected nodes not in any tree have depth zero.

Function plot displays the tree over a supplied ordination configuration, and lines adds a spanning tree to an ordination graph. If configuration is not supplied for plot, the function ordinates the cophenetic dissimilarities of the spanning tree and overlays the tree on this result. The default ordination function is sammon (package MASS), because Sammon scaling emphasizes structure in the neighbourhood of nodes and may be able to beautifully represent the tree (you may need to set dlim, and sometimes the results will remain twisted). These ordination methods do not work with disconnected trees, but you must supply the ordination configuration. Function lines will overlay the tree in an existing plot.

Function spantree uses Prim’s method implemented as priority-first search for dense graphs (Sedgewick 1990). Function cophenetic uses function stepacross with option path = "extended". The spantree is very fast, but cophenetic is slow in very large data sets.

Value

Function spantree returns an object of class spantree which is a list with two vectors, each of length $n - 1$. The number of links in a tree is one less the number of observations, and the first item is omitted. The items are

kid  The child node of the parent, starting from parent number two. If there is no link from the parent, value will be NA and tree is disconnected at the node.
Corresponding distance. If `kid = NA`, then `dist = 0`.

Names of nodes as found from the input dissimilarities.

The function call.

Note

In principle, minimum spanning tree is equivalent to single linkage clustering that can be performed using `hclust` or `agnes`. However, these functions combine clusters to each other and the information of the actually connected points (the “single link”) cannot be recovered from the result. The graphical output of a single linkage clustering plotted with `ordicluster` will look very different from an equivalent spanning tree plotted with `lines.spantree`.

Author(s)

Jari Oksanen

References


See Also

`vegdist` or `dist` for getting dissimilarities, and `hclust` or `agnes` for single linkage clustering.

Examples

data(dune)
dis <- vegdist(dune)
tr <- spantree(dis)
## Add tree to a metric scaling
plot(tr, cmdscale(dis), type = "t")
## Find a configuration to display the tree neatly
plot(tr, type = "t")
## Depths of nodes
depths <- spandepth(tr)
plot(tr, type = "t", label = depths)

---

**Description**

Function `specaccum` finds species accumulation curves or the number of species for a certain number of sampled sites or individuals.
Usage

```r
specaccum(comm, method = "exact", permutations = 100,
conditioned = TRUE, gamma = "jack1", ...)
## S3 method for class 'specaccum'
plot(x, add = FALSE, ci = 2, ci.type = c("bar", "line", "polygon"),
col = par("fg"), ci.col = col, ci.lty = 1, xlab,
ylab = x$method, ylim, xvar = c("sites", "individuals"), ...)
## S3 method for class 'specaccum'
boxplot(x, add = FALSE, ...)
```

Arguments

- **comm**: Community data set.
- **method**: Species accumulation method (partial match). Method "collector" adds sites in the order they happen to be in the data, "random" adds sites in random order, "exact" finds the expected (mean) species richness, "coleman" finds the expected richness following Coleman et al. 1982, and "rarefaction" finds the mean when accumulating individuals instead of sites.
- **permutations**: Number of permutations with method = "random".
- **conditioned**: Estimation of standard deviation is conditional on the empirical dataset for the exact SAC.
- **gamma**: Method for estimating the total extrapolated number of species in the survey area by function `specpool`.
- **x**: A `specaccum` result object.
- **add**: Add to an existing graph.
- **ci**: Multiplier used to get confidence intervals from standard deviation (standard error of the estimate). Value `ci = 0` suppresses drawing confidence intervals.
- **ci.type**: Type of confidence intervals in the graph: "bar" draws vertical bars, "line" draws lines, and "polygon" draws a shaded area.
- **col**: Colour for drawing lines.
- **ci.col**: Colour for drawing lines or filling the "polygon".
- **ci.lty**: Line type for confidence intervals or border of the "polygon".
- **xlab, ylab**: Labels for x (defaults `xvar`) and y axis.
- **ylim**: the y limits of the plot.
- **xvar**: Variable used for the horizontal axis: "individuals" can be used only with method = "rarefaction".
- **...**: Other parameters to functions.

Details

Species accumulation curves (SAC) are used to compare diversity properties of community data sets using different accumulator functions. The classic method is "random" which finds the mean SAC and its standard deviation from random permutations of the data, or subsampling without replacement (Gotelli & Colwell 2001). The "exact" method finds the expected SAC using the
method that was independently developed by Ugland et al. (2003), Colwell et al. (2004) and Kindt et al. (2006). The unconditional standard deviation for the exact SAC represents a moment-based estimation that is not conditioned on the empirical data set (sd for all samples > 0), unlike the conditional standard deviation that was developed by Jari Oksanen (not published, sd=0 for all samples). The unconditional standard deviation is based on an estimation of the total extrapolated number of species in the survey area (a.k.a. gamma diversity), as estimated by function `specpool`. Method "coleman" finds the expected SAC and its standard deviation following Coleman et al. (1982). All these methods are based on sampling sites without replacement. In contrast, the method = "rarefaction" finds the expected species richness and its standard deviation by sampling individuals instead of sites. It achieves this by applying function `rarefy` with number of individuals corresponding to average number of individuals per site.

The function has a plot method. In addition, method = "random" has summary and boxplot methods.

Value

The function returns an object of class "specaccum" with items:

- **call** Function call.
- **method** Accumulator method.
- **sites** Number of sites. For method = "rarefaction" this is the number of sites corresponding to a certain number of individuals and generally not an integer, and the average number of individuals is also returned in item `individuals`.
- **richness** The number of species corresponding to number of sites. With method = "collector" this is the observed richness, for other methods the average or expected richness.
- **sd** The standard deviation of SAC (or its standard error). This is NULL in method = "collector", and it is estimated from permutations in method = "random", and from analytic equations in other methods.
- **perm** Permutation results with method = "random" and NULL in other cases. Each column in `perm` holds one permutation.

Note

The SAC with method = "exact" was developed by Roeland Kindt, and its standard deviation by Jari Oksanen (both are unpublished). The method = "coleman" underestimates the SAC because it does not handle properly sampling without replacement. Further, its standard deviation does not take into account species correlations, and is generally too low.

Author(s)

Roeland Kindt <r.kindt@cgiar.org> and Jari Oksanen.

References


**See Also**

*rarefy* and *rrarefy* are related individual based models. Other accumulation models are *poolaccum* for extrapolated richness, and *renyiaccum* and *tsallisaccum* for diversity indices. Underlying graphical functions are *boxplot*, *matlines*, *segments* and *polygon*.

**Examples**

```r
data(BCI)
s1 <- specaccum(BCI)
s2 <- specaccum(BCI, "random")
s2
summary(s2)
plot(s1, ci.type="poly", col="blue", lwd=2, ci.lty=0, ci.col="lightblue")
boxplot(s2, col="yellow", add=TRUE, pch="+")
```

---

**specpool**

*Extrapolated Species Richness in a Species Pool*

**Description**

The functions estimate the extrapolated species richness in a species pool, or the number of unobserved species. Function *specpool* is based on incidences in sample sites, and gives a single estimate for a collection of sample sites (matrix). Function *estimateR* is based on abundances (counts) on single sample site.

**Usage**

```r
specpool(x, pool)
estimateR(x, ...)
specpool2vect(X, index = c("jack1","jack2", "chao", "boot","Species"))
poolaccum(x, permutations = 100, minsize = 3)
estaccumR(x, permutations = 100)
## S3 method for class 'poolaccum'
summary(object, display, alpha = 0.05, ...)
## S3 method for class 'poolaccum'
plot(x, alpha = 0.05, type = c("l","g"), ...)```
Arguments

- **x**: Data frame or matrix with species data or the analysis result for `plot` function.
- **pool**: A vector giving a classification for pooling the sites in the species data. If missing, all sites are pooled together.
- **X, object**: A `specpool` result object.
- **index**: The selected index of extrapolated richness.
- **permutations**: Number of permutations of sampling order of sites.
- **minsize**: Smallest number of sampling units reported.
- **display**: Indices to be displayed.
- **alpha**: Level of quantiles shown. This proportion will be left outside symmetric limits.
- **type**: Type of graph produced in `xyplot`.
- **...**: Other parameters (not used).

Details

Many species will always remain unseen or undetected in a collection of sample plots. The function uses some popular ways of estimating the number of these unseen species and adding them to the observed species richness (Palmer 1990, Colwell & Coddington 1994).

The incidence-based estimates in `specpool` use the frequencies of species in a collection of sites. In the following, $S_P$ is the extrapolated richness in a pool, $S_0$ is the observed number of species in the collection, $a_1$ and $a_2$ are the number of species occurring only in one or only in two sites in the collection, $p_i$ is the frequency of species $i$, and $N$ is the number of sites in the collection. The variants of extrapolated richness in `specpool` are:

- **Chao**
  \[ S_P = S_0 + \frac{a_1^2}{2(a_1 + 1)} \]
- **First order jackknife**
  \[ S_P = S_0 + a_1 \frac{N-1}{N} \]
- **Second order jackknife**
  \[ S_P = S_0 + a_2 \frac{N-3}{N} - a_2 \frac{(N-2)^2}{N(N-1)} \]
- **Bootstrap**
  \[ S_P = S_0 + \sum_{i=1}^{S_0} (1 - p_i) \]

The abundance-based estimates in `estimateR` use counts (frequencies) of species in a single site. If called for a matrix or data frame, the function will give separate estimates for each site. The two variants of extrapolated richness in `estimateR` are Chao (unbiased variant) and ACE. In the Chao estimate $a_i$ refers to number of species with abundance $i$ instead of incidence:

- **Chao**
  \[ S_P = S_0 + \frac{a_1(a_1-1)}{2(a_2+1)} \]
- **ACE**
  \[ S_P = S_{abund} + \frac{S_{rare}}{C_{ace}} + \frac{a_1}{C_{ace}} \gamma_{ace} \]

where

\[ C_{ace} = 1 - \frac{a_1}{N_{rare}} \]

\[ \gamma_{ace} = \max \left[ \frac{S_{rare} \sum_{i=1}^{a_{10}} i(i-1)a_i}{C_{ace} N_{rare}(N_{rare}-1)} - 1, 0 \right] \]

Here $a_i$ refers to number of species with abundance $i$ and $S_{rare}$ is the number of rare species, $S_{abund}$ is the number of abundant species, with an arbitrary threshold of abundance 10 for rare species, and $N_{rare}$ is the number of individuals in rare species.
Functions estimate the standard errors of the estimates. These only concern the number of added species, and assume that there is no variance in the observed richness. The equations of standard errors are too complicated to be reproduced in this help page, but they can be studied in the R source code of the function. The standard error are based on the following sources: Chao (1987) for the Chao estimate and Smith and van Belle (1984) for the first-order Jackknife and the bootstrap (second-order jackknife is still missing). The variance estimator of $S_{ace}$ was developed by Bob O’Hara (unpublished).

Functions poolaccum and estaccumR are similar to specaccum, but estimate extrapolated richness indices of specpool or estimateR in addition to number of species for random ordering of sampling units. Function specpool uses presence data and estaccumR count data. The functions share summary and plot methods. The summary returns quantile envelopes of permutations corresponding the given level of alpha and standard deviation of permutations for each sample size. The plot function shows the mean and envelope of permutations with given alpha for models. The selection of models can be restricted and order changes using the display argument in summary or plot. For configuration of plot command, see xypplot.

Value

Function specpool returns a data frame with entries for observed richness and each of the indices for each class in pool vector. The utility function specpool2vect maps the pooled values into a vector giving the value of selected index for each original site. Function estimateR returns the estimates and their standard errors for each site. Functions poolaccum and estimateR return matrices of permutation results for each richness estimator, the vector of sample sizes and a table of means of permutations for each estimator.

Note

The functions are based on assumption that there is a species pool: The community is closed so that there is a fixed pool size $S_P$. Such cases may exist, although I have not seen them yet. All indices are biased for open communities.

See http://viceroy.eeb.uconn.edu/EstimateS for a more complete (and positive) discussion and alternative software for some platforms.

Author(s)

Bob O’Hara (estimateR) and Jari Oksanen.

References


See Also

veiledspec, diversity, beals, specaccum.

Examples

data(dune)
data(dune.env)
attach(dune.env)
pool <- specpool(dune, Management)
pool
op <- par(mfrow=c(1,2))
boxplot(specnumber(dune) ~ Management, col="hotpink", border="cyan3", notch=TRUE)
boxplot(specnumber(dune)/specpool2vect(pool) ~ Management, col="hotpink", border="cyan3", notch=TRUE)
par(op)
data(BCI)
## Accumulation model
pool <- poolaccum(BCI)
summary(pool, display = "chao")
plot(pool)
## Quantitative model
estimateR(BCI[1:5,])

stepacross

Stepacross as Flexible Shortest Paths or Extended Dissimilarities

Description

Function stepacross tries to replace dissimilarities with shortest paths stepping across intermediate sites while regarding dissimilarities above a threshold as missing data (NA). With path = "shortest" this is the flexible shortest path (Williamson 1978, Bradfield & Kenkel 1987), and with path = "extended" an approximation known as extended dissimilarities (De’ath 1999). The use of stepacross should improve the ordination with high beta diversity, when there are many sites with no species in common.

Usage

stepacross(dis, path = "shortest", toolong = 1, trace = TRUE, ...)

Arguments

dis Dissimilarity data inheriting from class dist or a an object, such as a matrix, that can be converted to a dissimilarity matrix. Functions vegdist and dist are some functions producing suitable dissimilarity data.

path The method of stepping across (partial match) Alternative "shortest" finds the shortest paths, and "extended" their approximation known as extended dissimilarities.
toolong Shortest dissimilarity regarded as NA. The function uses a fuzz factor, so that dissimilarities close to the limit will be made NA, too.

trace Trace the calculations.

... Other parameters (ignored).

Details

Williamson (1978) suggested using flexible shortest paths to estimate dissimilarities between sites which have nothing in common, or no shared species. With path = "shortest" function stepacross replaces dissimilarities that are toolong or longer with NA, and tries to find shortest paths between all sites using remaining dissimilarities. Several dissimilarity indices are semi-metric which means that they do not obey the triangle inequality $d_{ij} \leq d_{ik} + d_{kj}$, and shortest path algorithm can replace these dissimilarities as well, even when they are shorter than toolong.

De'ath (1999) suggested a simplified method known as extended dissimilarities, which are calculated with path = "extended". In this method, dissimilarities that are toolong or longer are first made NA, and then the function tries to replace these NA dissimilarities with a path through single stepping stone points. If not all NA could be replaced with one pass, the function will make new passes with updated dissimilarities as long as all NA are replaced with extended dissimilarities. This mean that in the second and further passes, the remaining NA dissimilarities are allowed to have more than one stepping stone site, but previously replaced dissimilarities are not updated. Further, the function does not consider dissimilarities shorter than toolong, although some of these could be replaced with a shorter path in semi-metric indices, and used as a part of other paths. In optimal cases, the extended dissimilarities are equal to shortest paths, but they may be longer.

As an alternative to defining too long dissimilarities with parameter toolong, the input dissimilarities can contain NAs. If toolong is zero or negative, the function does not make any dissimilarities into NA. If there are no NAs in the input and toolong = 0, path = "shortest" will find shorter paths for semi-metric indices, and path = "extended" will do nothing. Function no.shared can be used to set dissimilarities to NA.

If the data are disconnected or there is no path between all points, the result will contain NAs and a warning is issued. Several methods cannot handle NA dissimilarities, and this warning should be taken seriously. Function distconnected can be used to find connected groups and remove rare outlier observations or groups of observations.

Alternative path = "shortest" uses Dijkstra’s method for finding flexible shortest paths, implemented as priority-first search for dense graphs (Sedgewick 1990). Alternative path = "extended" follows De’ath (1999), but implementation is simpler than in his code.

Value

Function returns an object of class dist with extended dissimilarities (see functions vegdist and dist). The value of path is appended to the method attribute.

Note

The function changes the original dissimilarities, and not all like this. It may be best to use the function only when you really must: extremely high beta diversity where a large proportion of dissimilarities are at their upper limit (no species in common).
Semi-metric indices vary in their degree of violating the triangle inequality. Morisita and Horn–Morisita indices of `vegdist` may be very strongly semi-metric, and shortest paths can change these indices very much. Mountford index violates basic rules of dissimilarities: non-identical sites have zero dissimilarity if species composition of the poorer site is a subset of the richer. With Mountford index, you can find three sites \( i, j, k \) so that \( d_{ik} = 0 \) and \( d_{jk} = 0 \), but \( d_{ij} > 0 \). The results of `stepacross` on Mountford index can be very weird. If `stepacross` is needed, it is best to try to use it with more metric indices only.

**Author(s)**

Jari Oksanen

**References**


**See Also**

Function `distconnected` can find connected groups in disconnected data, and function `no.shared` can be used to set dissimilarities as `NA`. See `swan` for an alternative approach. Function `stepacross` is an essential component in `isomap` and `cophenetic.spantree`.

**Examples**

```r
# There are no data sets with high beta diversity in vegan, but this
# should give an idea.
data(dune)
dis <- vegdist(dune)
edis <- stepacross(dis)
plot(edis, dis, xlab = "Shortest path", ylab = "Original")
## Manhattan distance have no fixed upper limit.
dis <- vegdist(dune, "manhattan")
is.na(dis) <- no.shared(dune)
dis <- stepacross(dis, toolong=0)
```

**Indices of Taxonomic Diversity and Distinctness**

Function finds indices of taxonomic diversity and distinctness, which are averaged taxonomic distances among species or individuals in the community (Clarke & Warwick 1998, 2001)
**Usage**

```r
taxondive(comm, dis, match.force = FALSE)
taxa2dist(x, varstep = FALSE, check = TRUE, labels)
```

**Arguments**

- `comm` Community data.
- `dis` Taxonomic distances among taxa in `comm`. This should be a `dist` object or a symmetric square matrix.
- `match.force` Force matching of column names in `comm` and labels in `dis`. If `FALSE`, matching only happens when dimensions differ, and in that case the species must be in identical order in both.
- `x` Classification table with a row for each species or other basic taxon, and columns for identifiers of its classification at higher levels.
- `varstep` Vary step lengths between successive levels relative to proportional loss of the number of distinct classes.
- `check` If `TRUE`, remove all redundant levels which are different for all rows or constant for all rows and regard each row as a different basal taxon (species). If `FALSE` all levels are retained and basal taxa (species) also must be coded as variables (columns). You will get a warning if species are not coded, but you can ignore this if that was your intention.
- `labels` The `labels` attribute of taxonomic distances. Row names will be used if this is not given. Species will be matched by these labels in `comm` and `dis` in `taxondive` if these have different dimensions.

**Details**

Clarke & Warwick (1998, 2001) suggested several alternative indices of taxonomic diversity or distinctness. Two basic indices are called taxonomic diversity ($\Delta$) and distinctness ($\Delta^*$):

\[
\Delta = \frac{(\sum_{i<j} \omega_{ij} x_i x_j)}{(n(n-1)/2)}
\]

\[
\Delta^* = \frac{(\sum_{i<j} \omega_{ij} x_i x_j)}{(\sum_{i<j} x_i x_j)}
\]

The equations give the index value for a single site, and summation goes over species $i$ and $j$. Here $\omega$ are taxonomic distances among taxa, and $x$ are species abundances, and $n$ is the total abundance for a site. With presence/absence data both indices reduce to the same index $\Delta^+$, and for this index Clarke & Warwick (1998) also have an estimate of its standard deviation. Clarke & Warwick (2001) presented two new indices: $s\Delta^+$ is the product of species richness and $\Delta^+$, and index of variation in taxonomic distinctness ($\Lambda^+$) defined as

\[
\Lambda^+ = \frac{(\sum_{i<j} \omega_{ij}^2)}{(n(n-1)/2)} - (\Delta^+)^2
\]

The `dis` argument must be species dissimilarities. These must be similar to dissimilarities produced by `dist`. It is customary to have integer steps of taxonomic hierarchies, but other kind of dissimilarities can be used, such as those from phylogenetic trees or genetic differences. Further, the `dis` need not be taxonomic, but other species classifications can be used.
Function `taxa2dist` can produce a suitable `dist` object from a classification table. Each species (or basic taxon) corresponds to a row of the classification table, and columns give the classification at different levels. With `varstep = FALSE` the successive levels will be separated by equal steps, and with `varstep = TRUE` the step length is relative to the proportional decrease in the number of classes (Clarke & Warwick 1999). With `check = TRUE`, the function removes classes which are distinct for all species or which combine all species into one class, and assumes that each row presents a distinct basic taxon. The function scales the distances so that longest path length between taxa is 100 (not necessarily when `check = FALSE`).

Function `plot.taxondive` plots $\Delta^+$ against Number of species, together with expectation and its approximate 2*sd limits. Function `summary.taxondive` finds the $z$ values and their significances from Normal distribution for $\Delta^+$. 

Value

Function returns an object of class `taxondive` with following items:

- **Species** Number of species for each site.
- **D, Dstar, Dplus, SDplus, Lambda** $\Delta, \Delta^*, \Delta^+, s\Delta^+$ and $\Lambda^+$ for each site.
- **sd.Dplus** Standard deviation of $\Delta^+$.
- **ED, EDstar, EDplus** Expected values of corresponding statistics.

Function `taxa2dist` returns an object of class "dist", with an attribute "steps" for the step lengths between successive levels.

Note

The function is still preliminary and may change. The scaling of taxonomic dissimilarities influences the results. If you multiply taxonomic distances (or step lengths) by a constant, the values of all Deltas will be multiplied with the same constant, and the value of $\Lambda^+$ by the square of the constant.

Author(s)

Jari Oksanen

References


See Also

diversity.
Examples

```r
## Preliminary: needs better data and some support functions
data(dune)
data(dune.taxon)
# Taxonomic distances from a classification table with variable step lengths.
taxdis <- taxa2dist(dune.taxon, varstep=TRUE)
plot(hclust(taxdis), hang = -1)
# Indices
mod <- taxondive(dune, taxdis)
mod
summary(mod)
plot(mod)
```

---

### Description

Functional diversity is defined as the total branch length in a trait dendrogram connecting all species, but excluding the unnecessary root segments of the tree (Petchey and Gaston 2006).

### Usage

- `treedive(comm, tree, match.force = FALSE)`
- `treeheight(tree)`
- `treedist(x, tree, relative = TRUE, ...)`

### Arguments

- `comm, x` Community data frame or matrix.
- `tree` A dendrogram which for `treedive` must be for species (columns).
- `match.force` Force matching of column names in `comm` and labels in `tree`. If `FALSE`, matching only happens when dimensions differ, and in that case the species must be in identical order in both.
- `relative` Use distances relative to the height of combined tree.
- `...` Other arguments passed to functions (ignored).

### Details

Function `treeheight` finds the sum of lengths of connecting segments in a dendrogram produced by `hclust`, or other dendrogram that can be coerced to a correct type using `as.hclust`. When applied to a clustering of species traits, this is a measure of functional diversity (Petchey and Gaston 2002, 2006).

Function `treedive` finds the `treeheight` for each site (row) of a community matrix. The function uses a subset of dendrogram for those species that occur in each site, and excludes the tree root if that is not needed to connect the species (Petchey and Gaston 2006). The subset of the
dendrogram is found by first calculating cophenetic distances from the input dendrogram, then reconstructing the dendrogram for the subset of the cophenetic distance matrix for species occurring in each site. Diversity is 0 for one species, and NA for empty communities.

Function `treedist` finds the dissimilarities among trees. Pairwise dissimilarity of two trees is found by combining species in a common tree and seeing how much of the tree height is shared and how much is unique. With `relative = FALSE` the dissimilarity is defined as $2(A \cup B) - A - B$, where $A$ and $B$ are heights of component trees and $A \cup B$ is the height of the combined tree. With `relative = TRUE` the dissimilarity is $(2(A \cup B) - A - B)/(A \cup B)$. Although the latter formula is similar to Jaccard dissimilarity (see `vegdist`, `designdist`), it is not in the range 0...1, since combined tree can add a new root. When two zero-height trees are combined into a tree of above zero height, the relative index attains its maximum value 2. The dissimilarity is zero from a combined zero-height tree.

The functions need a dendrogram of species traits as an input. If species traits contain `factor` or `ordered` factor variables, it is recommended to use Gower distances for mixed data (function `daisy` in package `cluster`), and usually the recommended clustering method is UPGMA (method = "average" in function `hclust`) (Podani and Schmera 2006).

It is possible to analyse the non-randomness of functional diversity using `oecosimu`. This needs specifying an adequate Null model, and the results will change with this choice.

Value

A vector of diversity values or a single tree height, or a dissimilarity structure that inherits from `dist` and can be used similarly.

Author(s)

Jari Oksanen

References


See Also

taxondive is something very similar from another world.

Examples

```r
## There is no data set on species properties yet, and therefore
## the example uses taxonomy
data(dune)
```
data(dune.taxon)
d <- taxa2dist(dune.taxon, varstep=TRUE)
c1 <- hclust(d, "aver")
treedive(dune, c1)
## Significance test using Null model communities.
## The current choice fixes only site totals.
oecosimu(dune, treedive, "r0", tree = c1)
## Clustering of tree distances
dtree <- treedist(dune, c1)
plot(hclust(dtree, "aver"))

---

**tsallis**  
*Tsallis Diversity and Corresponding Accumulation Curves*

**Description**

Function *tsallis* find Tsallis diversities with any scale or the corresponding evenness measures. Function *tsallisaccum* finds these statistics with accumulating sites.

**Usage**

```r
tsallis(x, scales = seq(0, 2, 0.2), norm = FALSE, hill = FALSE)
tsallisaccum(x, scales = seq(0, 2, 0.2), permutations = 100, raw = FALSE, ...)
## S3 method for class 'tsallisaccum'
persp(x, theta = 220, phi = 15, col = heat.colors(100), zlim, ...)
```

**Arguments**

- **x**  
  Community data matrix or plotting object.

- **scales**  
  Scales of Tsallis diversity.

- **norm**  
  Logical, if TRUE diversity values are normalized by their maximum (diversity value at equiprobability conditions).

- **hill**  
  Calculate Hill numbers.

- **permutations**  
  Number of random permutations in accumulating sites.

- **raw**  
  If FALSE then return summary statistics of permutations, and if TRUE then returns the individual permutations.

- **theta, phi**  
  angles defining the viewing direction. theta gives the azimuthal direction and phi the colatitude.

- **col**  
  Colours used for surface.

- **zlim**  
  Limits of vertical axis.

- **...**  
  Other arguments which are passed to *tsallis* and to graphical functions.
Details

The Tsallis diversity (also equivalent to Patil and Taillie diversity) is a one-parametric generalised entropy function, defined as:

\[ H_q = \frac{1}{q-1} \left( 1 - \sum_{i=1}^{S} p_i^q \right) \]

where \( q \) is a scale parameter, \( S \) the number of species in the sample (Tsallis 1988, Tothmeresz 1995). This diversity is concave for all \( q > 0 \), but non-additive (Keylock 2005). For \( q = 0 \) it gives the number of species minus one, as \( q \) tends to 1 this gives Shannon diversity, for \( q = 2 \) this gives the Simpson index (see function `diversity`).

If `norm = TRUE`, `tsallis` gives values normalized by the maximum:

\[ H_q(max) = \frac{S^{1-q} - 1}{1 - q} \]

where \( S \) is the number of species. As \( q \) tends to 1, maximum is defined as \( ln(S) \).

If `hill = TRUE`, `tsallis` gives Hill numbers (numbers equivalents, see Jost 2007):

\[ D_q = (1 - (q - 1)H)^{1/(1-q)} \]

Details on plotting methods and accumulating values can be found on the help pages of the functions `renyi` and `renyiaccum`.

Value

Function `tsallis` returns a data frame of selected indices. Function `tsallisaccum` with argument `raw = FALSE` returns a three-dimensional array, where the first dimension are the accumulated sites, second dimension are the diversity scales, and third dimension are the summary statistics `mean`, `stdev`, `min`, `max`, `Qnt 0.025` and `Qnt 0.975`. With argument `raw = TRUE` the statistics on the third dimension are replaced with individual permutation results.

Author(s)

Péter Sólymos, <solymos@ualberta.ca>, based on the code of Roeland Kindt and Jari Oksanen written for `renyi`.

References

See Also

Plotting methods and accumulation routines are based on functions renyi and renyiaccum. An object of class ‘tsallisaccum’ can be used with function rgl.renyiaccum as well. See also settings for persp.

Examples

data(BCI)
i <- sample(nrow(BCI), 12)
x1 <- tsallis(BCI[i,])
x1
diversity(BCI[i,],"simpson") == x1["2"]
plot(x1)
x2 <- tsallis(BCI[i,],norm=TRUE)
x2
plot(x2)
mod1 <- tsallisaccum(BCI[i,])
plot(mod1, as.table=TRUE, col = c(1, 2, 2))
persp(mod1)
mod2 <- tsallisaccum(BCI[i,], norm=TRUE)
persp(mod2,theta=100,phi=30)

varespec

Vegetation and environment in lichen pastures

Description

The varespec data frame has 24 rows and 44 columns. Columns are estimated cover values of 44 species. The variable names are formed from the scientific names, and are self explanatory for anybody familiar with the vegetation type. The varechem data frame has 24 rows and 14 columns, giving the soil characteristics of the very same sites as in the varespec data frame. The chemical measurements have obvious names. Baresoil gives the estimated cover of bare soil, Humdepth the thickness of the humus layer.

Usage

data(varechem)
data(varespec)

References


Examples

data(varespec)
data(varechem)
varpart

Partition the Variation of Community Matrix by 2, 3, or 4 Explanatory Matrices

Description

The function partitions the variation of response table Y with respect to two, three, or four explanatory tables, using redundancy analysis ordination (RDA). If Y contains a single vector, partitioning is by partial regression. Collinear variables in the explanatory tables do NOT have to be removed prior to partitioning.

Usage

varpart(Y, X, ..., data, transfo, scale = FALSE)
showvarparts(parts, labels, ...)
## S3 method for class 'varpart234'
plot(x, cutoff = 0, digits = 1, ...)

Arguments

Y
Data frame or matrix containing the response data table. In community ecology, that table is often a site-by-species table.

X
Two to four explanatory models, variables or tables. These can be defined in three alternative ways: (1) one-sided model formulae beginning with ~ and then defining the model, (2) name of a single numeric variable, or (3) name of data frame or matrix with numeric variables. The model formulae can have factors, interaction terms and transformations of variables. The names of the variables in the model formula are found in data frame given in data argument, and if not found there, in the user environment. Single numeric variables, data frames or matrices are found in the user environment. All entries till the next argument (data or transfo) are interpreted as explanatory models, and the names of these arguments cannot be abbreviated nor omitted.

data
The data frame with the variables used in the formulae in X.

transfo
Transformation for Y (community data) using decostand. All alternatives in decostand can be used, and those preserving Euclidean metric include "hellinger", "chi.square", "total", "norm".

scale
Should the columns of Y be standardized to unit variance?

parts
Number of explanatory tables (circles) displayed.

labels
Labels used for displayed fractions. Default is to use the same letters as in the printed output.

x
The varpart result.

cutoff
The values below cutoff will not be displayed.

digits
The number of significant digits; the number of decimal places is at least one higher.

...
Other parameters passed to functions.
Details

The functions partition the variation in $Y$ into components accounted for by two to four explanatory tables and their combined effects. If $Y$ is a multicolour data frame or matrix, the partitioning is based on redundancy analysis (RDA, see rda), and if $Y$ is a single variable, the partitioning is based on linear regression. A simplified, fast version of RDA is used (function simpleRDA2). The actual calculations are done in functions varpart2 to varpart4, but these are not intended to be called directly by the user.

The function primarily uses adjusted R squares to assess the partitions explained by the explanatory tables and their combinations, because this is the only unbiased method (Peres-Neto et al., 2006). The raw R squares for basic fractions are also displayed, but these are biased estimates of variation explained by the explanatory table.

The identifiable fractions are designated by lower case alphabets. The meaning of the symbols can be found in the separate document "partitioning.pdf" (which can be read using vegandocs), or can be displayed graphically using function showvarparts.

A fraction is testable if it can be directly expressed as an RDA model. In these cases the printed output also displays the corresponding RDA model using notation where explanatory tables after $|$ are conditions (partialled out; see rda for details). Although single fractions can be testable, this does not mean that all fractions simultaneously can be tested, since there number of testable fractions is higher than the number of estimated models.

An abridged explanation of the alphabetic symbols for the individual fractions follows, but computational details should be checked in "partitioning.pdf" (readable with vegandocs) or in the source code.

With two explanatory tables, the fractions explained uniquely by each of the two tables are $a$ and $c$, and their joint effect is $b$ following Borcard et al. (1992).

With three explanatory tables, the fractions explained uniquely by each of the three tables are $a$ to $c$, joint fractions between two tables are $d$ to $f$, and the joint fraction between all three tables is $g$.

With four explanatory tables, the fractions explained uniquely by each of the four tables are $a$ to $d$, joint fractions between two tables are $e$ to $j$, joint fractions between three variables are $k$ to $n$, and the joint fraction between all four tables is $o$.

There is a plot function that displays the Venn diagram and labels each intersection (individual fraction) with the adjusted R squared if this is higher than cutoff. A helper function showvarpart displays the fraction labels.

Value

Function varpart returns an object of class "varpart" with items scale and transfo (can be missing) which hold information on standardizations, tables which contains names of explanatory tables, and call with the function call. The function varpart calls function varpart2, varpart3 or varpart4 which return an object of class "varpart234" and saves its result in the item part. The items in this object are:

- SS.Y: Sum of squares of matrix Y.
- n: Number of observations (rows).
- nsets: Number of explanatory tables.
bigwarning  Warnings on collinearity.
fract  Basic fractions from all estimated constrained models.
indfract  Individual fractions or all possible subsections in the Venn diagram (see showvarparts).
contr1  Fractions that can be found after conditioning on single explanatory table in models with three or four explanatory tables.
contr2  Fractions that can be found after conditioning on two explanatory tables in models with four explanatory tables.

Fraction Data Frames

Items fract, indfract, contr1 and contr2 are all data frames with items:

- Df Degrees of freedom of numerator of the \( F \)-statistic for the fraction.
- R.square Raw R-squared. This is calculated only for fract and this is NA in other items.
- Adj.R.square Adjusted R-squared.
- Testable If the fraction can be expressed as a (partial) RDA model, it is directly Testable, and this field is TRUE. In that case the fraction label also gives the specification of the testable RDA model.

Note

You can use command vegandocs to display document "partitioning.pdf" which presents Venn diagrams showing the fraction names in partitioning the variation of Y with respect to 2, 3, and 4 tables of explanatory variables, as well as the equations used in variation partitioning.

The functions frequently give negative estimates of variation. Adjusted R-squares can be negative for any fraction; unadjusted R squares of testable fractions always will be non-negative. Non-testable fractions cannot be found directly, but by subtracting different models, and these subtraction results can be negative. The fractions are orthogonal, or linearly independent, but more complicated or nonlinear dependencies can cause negative non-testable fractions.

The current function will only use RDA in multivariate partitioning. It is much more complicated to estimate the adjusted R-squares for CCA, and unbiased analysis of CCA is not currently implemented.

Author(s)

Pierre Legendre, Departement de Sciences Biologiques, Universite de Montreal, Canada. Adapted to vegan by Jari Oksanen.

References

(a) References on variation partitioning

(b) Reference on transformations for species data

(c) Reference on adjustment of the bimultivariate redundancy statistic


See Also

For analysing testable fractions, see rda and anova.cca. For data transformation, see decostand. Function inertcomp gives (unadjusted) components of variation for each species or site separately.

Examples

data(mite)
data(mite.env)
data(mite.pcnm)

## See detailed documentation:
## Not run:
# vegandocs("partition")
## End(Not run)

# Two explanatory matrices -- Hellinger-transform Y
# Formula shortcut "~ ." means: use all variables in 'data'.
mod <- varpart(mite, ~ ., mite.pcnm, data=mite.env, transfo="hel")
mod
showvarparts(2)
plot(mod)

# Alternative way of to conduct this partitioning
# Change the data frame with factors into numeric model matrix
# Test fraction [a] using RDA:
mod <- varpart(decostand(mite, "hel"), mm, mite.pcnm)
rda.result <- rda(decostand(mite, "hell"), mm, mite.pcnm)
anova(rda.result, step=200, perm.max=200)

# Three explanatory matrices
mod <- varpart(mite, ~ SubsDens + WatrCont, ~ Substrate + Shrub + Topo, mite.pcnm, data=mite.env, transfo="hel")
mod
showvarparts(3)
plot(mod)

# An alternative formulation of the previous model using
# matrices mm1 and mm2 and Hellinger transformed species data
# Use RDA to test fraction [a]
# Matrix can be an argument in formula
rda.result <- rda(mite.hel ~ mm1 + Condition(mm2) +
                  Condition(as.matrix(mite.pcnm))
anova(rda.result, step=200, perm.max=200)

# Four explanatory tables
mod <- varpart(mite, ~ SubsDens + WatrCont, ~Substrate + Shrub + Topo,
                mite.pcnm[,1:11], mite.pcnm[,12:22], data=mite.env, transfo="hel")
mod
plot(mod)
# Show values for all partitions by putting 'cutoff' low enough:
plot(mod, cutoff = -Inf, cex = 0.7)

---

vegandocs

Display Package Documentation

Description

Display package documentation using pager or pdfviewer defined in options.

Usage

vegandocs(doc = c("NEWS", "ChangeLog", "FAQ-vegan.pdf",
"partitioning.pdf"))

Arguments

doc                  The name of the document (partial match, case sensitive).

Note

The function is a kluge, since R does not have this facility (I hope it will come there). Function vignette only works with vignettes.

Author(s)

Jari Oksanen

See Also

vignette.

Examples

## Not run:
vegandocs("Change")

## End(Not run)
Description

The function computes dissimilarity indices that are useful for or popular with community ecologists. All indices use quantitative data, although they would be named by the corresponding binary index, but you can calculate the binary index using an appropriate argument. If you do not find your favourite index here, you can see if it can be implemented using `designdist`. Gower, Bray–Curtis, Jaccard and Kulczynski indices are good in detecting underlying ecological gradients (Faith et al. 1987). Morisita, Horn–Morisita, Binomial and Chao indices should be able to handle different sample sizes (Wolda 1981, Krebs 1999, Anderson & Millar 2004), and Mountford (1962) and Raup-Crick indices for presence–absence data should be able to handle unknown (and variable) sample sizes.

Usage

```r
vegdist(x, method="bray", binary=FALSE, diag=FALSE, upper=FALSE, 
na.rm = FALSE, ...)
```

Arguments

- **x** Community data matrix.
- **method** Dissimilarity index, partial match to "manhattan", "euclidean", "canberra", "bray", "kulczynski", "jaccard", "gower", "altGower", "morisita", "horn", "mountford", "raup", "binomial" or "chao".
- **binary** Perform presence/absence standardization before analysis using `decostand`.
- **diag** Compute diagonals.
- **upper** Return only the upper diagonal.
- **na.rm** Pairwise deletion of missing observations when computing dissimilarities.
- **...** Other parameters. These are ignored, except in method = "gower" which accepts `range.global` parameter of `decostand`.

Details

Jaccard ("jaccard"), Mountford ("mountford"), Raup–Crick ("raup"), Binomial and Chao indices are discussed later in this section. The function also finds indices for presence/ absence data by setting `binary = TRUE`. The following overview gives first the quantitative version, where \( x_{ij} \) and \( x_{ik} \) refer to the quantity on species (column) \( i \) and sites (rows) \( j \) and \( k \). In binary versions \( A \) and \( B \) are the numbers of species on compared sites, and \( J \) is the number of species that occur on both compared sites similarly as in `designdist` (many indices produce identical binary versions):

- **euclidean**
  \[
  d_{jk} = \sqrt{\sum_i (x_{ij} - x_{ik})^2}
  \]
  binary: \( \sqrt{A + B - 2J} \)
- **manhattan**
  \[
  d_{jk} = \sum_i |x_{ij} - x_{ik}|
  \]
binary: $A + B - 2J$

**gower**
$$d_{jk} = (1/M) \sum_i |x_{ij} - x_{ik}|$$
binary: $(A + B - 2J)/M$, where $M$ is the number of columns (excluding missing values)

**altGower**
$$d_{jk} = (1/NZ) \sum_i |x_{ij} - x_{ik}|$$
where $NZ$ is the number of non-zero columns excluding double-zeros (Anderson et al. 2006).

**canberra**
$$d_{jk} = \frac{1}{NZ} \sum_i \frac{|x_{ij} - x_{ik}|}{x_{ij} + x_{ik}}$$
where $NZ$ is the number of non-zero entries.

**bray**
$$d_{jk} = \frac{\sum_i |x_{ij} - x_{ik}|}{\sum_i (x_{ij} + x_{ik})}$$
binary: $(A + B - 2J)/A + B$

**kulczynski**
$$d_{jk} = 1 - 0.5 \left( \sum_i \min(x_{ij}, x_{ik}) + \sum_i \min(x_{ij}, x_{ik}) \right)$$
binary: $1 - (J/A + J/B)/2$

**morisita**
$$d_{jk} = 1 - \frac{2}{(\lambda_j + \lambda_k)} \sum_i x_{ij} \sum_i x_{ik},$$
where
$$\lambda_j = \sum_i x_{ij} \sum_i (x_{ij}-1)$$
binary: cannot be calculated

**horn**
Like morisita, but $\lambda_j = \sum_i x_{ij}^2 / (\sum_i x_{ij})^2$
binary: $(A + B - 2J)/A + B$

**binomial**
$$d_{jk} = \sum_i |x_{ij} \log(\frac{x_{ij}}{n_i}) + x_{ik} \log(\frac{x_{ik}}{n_i}) - n_i \log(\frac{1}{2})| / n_i,$$
where $n_i = x_{ij} + x_{ik}$
binary: $\log(2) \times (A + B - 2J)$

Jaccard index is computed as $2B/(1 + B)$, where $B$ is Bray–Curtis dissimilarity.

Binomial index is derived from Binomial deviance under null hypothesis that the two compared communities are equal. It should be able to handle variable sample sizes. The index does not have a fixed upper limit, but can vary among sites with no shared species. For further discussion, see Anderson & Millar (2004).

Mountford index is defined as $M = 1/\alpha$ where $\alpha$ is the parameter of Fisher’s logseries assuming that the compared communities are samples from the same community (cf. `fisherfit, fisher.alpha`). The index $M$ is found as the positive root of equation $\exp(aM) + \exp(bM) = 1 + \exp[(a + b - j)M]$, where $j$ is the number of species occurring in both communities, and $a$ and $b$ are the number of species in each separate community (so the index uses presence–absence information). Mountford index is usually misrepresented in the literature: indeed Mountford (1962) suggested an approximation to be used as starting value in iterations, but the proper index is defined as the root of the equation above. The function `vegdist` solves $M$ with the Newton method. Please note that if either $a$ or $b$ are equal to $j$, one of the communities could be a subset of the other, and the dissimilarity is $0$ meaning that non-identical objects may be regarded as similar and the index is non-metric. The Mountford index is in the range $0 \ldots \log(2)$, but the dissimilarities are divided by $\log(2)$ so that the results will be in the conventional range $0 \ldots 1$.

Raup–Crick dissimilarity (method = "raup") is a probabilistic index based on presence/absence data. It is defined as $1 - \text{prob}(j)$, or based on the probability of observing at least $j$ species in shared in compared communities. Legendre & Legendre (1998) suggest using simulations to as-
sess the probability, but the current function uses analytic result from hypergeometric distribution (phyper) instead. This probability (and the index) is dependent on the number of species missing in both sites, and adding all-zero species to the data or removing missing species from the data will influence the index. The probability (and the index) may be almost zero or almost one for a wide range of parameter values. The index is nonmetric: two communities with no shared species may have a dissimilarity slightly below one, and two identical communities may have dissimilarity slightly above zero.

Chao index tries to take into account the number of unseen species pairs, similarly as in method = "chao" in specpool. Function vegdist implements a Jaccard type index defined as 
\[ d_{jk} = 1 - \frac{U_j U_k}{U_j + U_k - U_j U_k}, \]
where 
\[ U_j = C_j/N_j + (N_k - 1)/N_k \times a_1/(2a_2) \times S_j/N_j, \]
and similarly for \( U_k \). Here \( C_j \) is the total number of individuals in the species of site \( j \) that are shared with site \( k \), \( N_j \) is the total number of individuals at site \( j \), \( a_1 \) (and \( a_2 \)) are the number of species occurring in site \( j \) that have only one (or two) individuals in site \( k \), and \( S_j \) is the total number of individuals in the species present at site \( j \) that occur with only one individual in site \( k \) (Chao et al. 2005).

Morisita index can be used with genuine count data (integers) only. Its Horn–Morisita variant is able to handle any abundance data.

Euclidean and Manhattan dissimilarities are not good in gradient separation without proper standardization but are still included for comparison and special needs.

Bray–Curtis and Jaccard indices are rank-order similar, and some other indices become identical or rank-order similar after some standardizations, especially with presence/absence transformation of equalizing site totals with decostand. Jaccard index is metric, and probably should be preferred instead of the default Bray-Curtis which is semimetric.

The naming conventions vary. The one adopted here is traditional rather than truthful to priority. The function finds either quantitative or binary variants of the indices under the same name, which correctly may refer only to one of these alternatives For instance, the Bray index is known also as Steinhaus, Czekanowski and Sørensen index. The quantitative version of Jaccard should probably called Ružička index. The abbreviation "horn" for the Horn–Morisita index is misleading, since there is a separate Horn index. The abbreviation will be changed if that index is implemented in vegan.

Value

Should provide a drop-in replacement for dist and return a distance object of the same type.

Note

The function is an alternative to dist adding some ecologically meaningful indices. Both methods should produce similar types of objects which can be interchanged in any method accepting either. Manhattan and Euclidean dissimilarities should be identical in both methods. Canberra index is divided by the number of variables in vegdist, but not in dist. So these differ by a constant multiplier, and the alternative in vegdist is in range (0,1). Function daisy (package cluster) provides alternative implementation of Gower index that also can handle mixed data of numeric and class variables. There are two versions of Gower distance ("gower", "altGower") which differ in scaling: "gower" divides all distances by the number of observations (rows) and scales each column to unit range, but "altGower" omits double-zeros and divides by the number of pairs with at least one above-zero value, and does not scale columns (Anderson et al. 2006). You can
use \texttt{decostand} to add range standardization to "altGower" (see Examples). Gower (1971) suggested omitting double zeros for presences, but it is often taken as the general feature of the Gower distances. See Examples for implementing the Anderson et al. (2006) variant of the Gower index.

Most dissimilarity indices in \texttt{vegdist} are designed for community data, and they will give misleading values if there are negative data entries. The results may also be misleading or \texttt{NA} or \texttt{NaN} if there are empty sites. In principle, you cannot study species composition without species and you should remove empty sites from community data.

Author(s)

Jari Oksanen, with contributions from Tyler Smith (Gower index) and Michael Bedward (Raup–Crick index).

References


See Also

Function \texttt{designdist} can be used for defining your own dissimilarity index. Alternative dissimilarity functions include \texttt{dist} in base \texttt{R}, \texttt{daisy} (package \texttt{cluster}), and \texttt{dsdist} (package \texttt{labdsv}). Function \texttt{betadiver} provides indices intended for the analysis of beta diversity.

Examples

```r
data(varespec)
vare.dist <- vegdist(varespec)
# Orlóci's Chord distance: range 0 .. sqrt(2)
vare.dist <- vegdist(decostand(varespec, "norm"), "euclidean")
# Anderson et al. (2006) version of Gower
```
vare.dist <- vegdist(decostand(varespec, "log"), "altGower")
# Range standardization with "altGower" (that excludes double-zeros)
vare.dist <- vegdist(decostand(varespec, "range"), "altGower")

---

**vegemite**

*Prints a Compact, Ordered Vegetation Table*

**Description**

The function prints a compact vegetation table, where species are rows, and each site takes only one column without spaces. The vegetation table can be ordered by explicit indexing, by environmental variables or results from an ordination or cluster analysis.

**Usage**

```r
vegemite(x, use, scale, sp.ind, site.ind, zero=".", select ,...)
coverscale(x, scale=c("Braun.Blanquet", "Domin", "Hult", "Hill", "fix","log"), maxabund)
```

**Arguments**

- **x**
  - Vegetation data.

- **use**
  - Either a vector, or an object from `cca`, `decorana` *etc.* or `hclust` or a `dendrogram` for ordering sites and species.

- **sp.ind**
  - Species indices.

- **site.ind**
  - Site indices.

- **zero**
  - Character used for zeros.

- **select**
  - Select a subset of sites. This can be a logical vector (TRUE for selected sites), or a vector of indices of selected sites. The order of indices does not influence results, but you must specify `use` or `site.ind` to reorder sites.

- **scale**
  - Cover scale used (can be abbreviated).

- **maxabund**
  - Maximum abundance used with `scale = "log"`. Data maximum in the selected subset will be used if this is missing.

- **...**
  - Arguments passed to `coverscale` (i.e., `maxabund`).

**Details**

The function prints a traditional vegetation table. Unlike in ordinary data matrices, species are used as rows and sites as columns. The table is printed in compact form: only one character can be used for abundance, and there are no spaces between columns. Species with no occurrences are dropped from the table.

The parameter `use` can be a vector or an object from `hclust`, a `dendrogram` or any ordination result recognized by `scores` (all ordination methods in `vegan` and some of those not in `vegan`). If `use` is a vector, it is used for ordering sites. If `use` is an object from ordination, both sites and species are arranged by the first axis. When `use` is an object from `hclust` or a `dendrogram`,...
the sites are ordered similarly as in the cluster dendrogram. If ordination methods provide species scores, these are used for ordering species. In all cases where species scores are missing, species are ordered by their weighted averages \((\text{wascores})\) on site scores. There is no natural, unique ordering in hierarchic clustering, but in some cases species are still nicely ordered (please note that you can \texttt{reorder.dendrogram} to have such a natural order). Alternatively, species and sites can be ordered explicitly giving their indices or names in parameters \texttt{sp.ind} and \texttt{site.ind}. If these are given, they take precedence over \texttt{use}. A subset of sites can be displayed using argument \texttt{select}, but this cannot be used to order sites, but you still must give \texttt{use} or \texttt{site.ind}.

If \texttt{scale} is given, \texttt{vegemite} calls \texttt{coverscale} to transform percent cover scale or some other scales into traditional class scales used in vegetation science (\texttt{coverscale} can be called directly, too). Braun-Blanquet and Domin scales are actually not strict cover scales, and the limits used for codes \(r\) and \(+\) are arbitrary. Scale \texttt{Hill} may be inappropriately named, since Mark O. Hill probably never intended this as a cover scale. However, it is used as default 'cut levels' in his \texttt{TWINSPAN}, and surprisingly many users stick to this default, and this is a \textit{de facto} standard in publications. All traditional scales assume that values are cover percentages with maximum 100. However, non-traditional alternative \texttt{log} can be used with any scale range. Its class limits are integer powers of \(1/2\) of the maximum (argument \texttt{maxabund}), with \(+\) used for non-zero entries less than \(1/512\) of the maximum (\texttt{log} stands alternatively for logarithmic or logical). Scale \texttt{fix} is intended for ‘fixing’ 10-point scales: it truncates scale values to integers, and replaces 10 with \(X\) and positive values below 1 with +.

**Value**

The function is used mainly to print a table, but it returns (invisibly) a list with items:

- \texttt{species} Ordered species indices
- \texttt{sites} Ordered site indices

These items can be used as arguments \texttt{sp.ind} and \texttt{site.ind} to reproduce the table. In addition to the proper table, the function prints the numbers of species and sites and the name of the used cover scale at the end.

**Note**

This function was called \texttt{vegetab} in older versions of \texttt{vegan}. The new name was chosen because the output is so compact (and to avoid confusion with the \texttt{vegtab} function in the \texttt{labdsv} package).

**Author(s)**

Jari Oksanen

**References**

The cover scales are presented in many textbooks of vegetation science; I used:


**See Also**

\texttt{cut} and \texttt{approx} for making your own ‘cover scales’, \texttt{wascores} for weighted averages.
Examples

data(varespec)
## Print only more common species
freq <- apply(varespec > 0, 2, sum)
vegemite(varespec, scale="Hult", sp.ind = freq > 10)
## Order by correspondence analysis, use Hill scaling and layout:
dca <- decorana(varespec)
vegemite(varespec, dca, "Hill", zero="-"
## Show one class from cluster analysis, but retain the ordering above
clus <- hclust(vegdist(varespec))
c1 <- cutree(clus, 3)
cl <- vegemite(varespec, use=dca, select = cl == 3, scale="Br")
# Re-create previous
vegemite(varespec, sp=sel$sp, site=sel$site, scale="Hult")

wascres Weighted Averages Scores for Species

Description

Computes Weighted Averages scores of species for ordination configuration or for environmental variables.

Usage

wascres(x, w, expand=FALSE)
eigengrad(x, w)

Arguments

x Environmental variables or ordination scores.
w Weights: species abundances.
expand Expand weighted averages so that they have the same weighted variance as the corresponding environmental variables.

Details

Function `wascres` computes weighted averages. Weighted averages ‘shrink’: they cannot be more extreme than values used for calculating the averages. With `expand = TRUE`, the function ‘dehrsinks’ the weighted averages by making their biased weighted variance equal to the biased weighted variance of the corresponding environmental variable. Function `eigengrad` returns the inverses of squared expansion factors or the attribute `shrinkage` of the `wascres` result for each environmental gradient. This is equal to the constrained eigenvalue of `cca` when only this one gradient was used as a constraint, and describes the strength of the gradient.
Value

Function `wascores` returns a matrix where species define rows and ordination axes or environmental variables define columns. If `expand = TRUE`, attribute `shrinkage` has the inverses of squared expansion factors or `cca` eigenvalues for the variable. Function `eigengrad` returns only the `shrinkage` attribute.

Author(s)

Jari Oksanen

See Also

`isoMDS`, `cca`.

Examples

```r
data(varespec)
data(varechem)
library(MASS)  # isoMDS
vare.dist <- vegdist(wisconsin(varespec))
vare.mds <- isoMDS(vare.dist)
vare.points <- postMDS(vare.mds$points, vare.dist)
vare.wa <- wascores(vare.points, varespec)
plot(scores(vare.points), pch="+", asp=1)
text(vare.wa, rownames(vare.wa), cex=0.8, col="blue")
# Omit rare species (frequency <= 4)
freq <- apply(varespec>0, 2, sum)
plot(scores(vare.points), pch="+", asp=1)
text(vare.wa[freq > 4,], rownames(vare.wa)[freq > 4], cex=0.8, col="blue")
# Works for environmental variables, too.
wascores(varechem, varespec)
# And the strengths of these variables are:
eigengrad(varechem, varespec)
```

---

**wcmdscale**

*Weighted Classical (Metric) Multidimensional Scaling*

Description

Weighted classical multidimensional scaling, also known as weighted *principal coordinates analysis*.

Usage

```r
wcmdscale(d, k, eig = FALSE, add = FALSE, x.ret = FALSE, w)
```
Arguments

- **d**: a distance structure such as that returned by `dist` or a full symmetric matrix containing the dissimilarities.
- **k**: the dimension of the space which the data are to be represented in; must be in \{1, 2, ..., \(n - 1\)\}. If missing, all dimensions with above zero eigenvalue.
- **eig**: indicates whether eigenvalues should be returned.
- **add**: logical indicating if an additive constant \(c^*\) should be computed, and added to the non-diagonal dissimilarities such that all \(n - 1\) eigenvalues are non-negative. **Not implemented.**
- **x.ret**: indicates whether the doubly centred symmetric distance matrix should be returned.
- **w**: Weights of points.

Details

Function `wcmdscale` is based on function `cmdscale` (package `stats` of base R), but it uses point weights. Points with high weights will have a stronger influence on the result than those with low weights. Setting equal weights \(w = 1\) will give ordinary multidimensional scaling.

Value

If `eig = FALSE` and `x.ret = FALSE` (default), a matrix with \(k\) columns whose rows give the coordinates of the points chosen to represent the dissimilarities. Otherwise, an object of class `wcmdscale` containing the components that are mostly similar as in `cmdscale`:

- **points**: a matrix with \(k\) columns whose rows give the coordinates of the points chosen to represent the dissimilarities.
- **eig**: the \(n - 1\) eigenvalues computed during the scaling process if `eig` is true.
- **x**: the doubly centred and weighted distance matrix if `x.ret` is true.
- **GOF**: Goodness of fit statistics for \(k\) axes. The first value is based on the sum of absolute values of all eigenvalues, and the second value is based on the sum of positive eigenvalues
- **weights**: Weights.
- **negaxes**: A matrix of scores for axes with negative eigenvalues scaled by the absolute eigenvalues similarly as `points`. This is **NULL** if there are no negative eigenvalues or \(k\) was specified, and would not include negative eigenvalues.

References


See Also

`cmdscale`. Also `isoMDS` and `sammon` in package `MASS`. 
Examples

## Correspondence analysis as a weighted principal coordinates analysis of Euclidean distances of Chi-square transformed data
data(dune)
rs <- rowSums(dune)/sum(dune)
d <- dist(decostand(dune, "chi"))
ord <- wcmdscale(d, w = rs, eig = TRUE)
## Ordinary CA
ca <- cca(dune)
## Eigenvalues are numerically similar
cia$CA$eig - ord$eig
## Configurations are similar when site scores are scaled by eigenvalues in CA
procrustes(ord, ca, choices=1:19, scaling = 1)
plot(procrustes(ord, ca, choices=1:2, scaling=1))
## Reconstruction of non-Euclidean distances with negative eigenvalues
d <- vegdist(dune)
ord <- wcmdscale(d, eig = TRUE)
## Only positive eigenvalues:
cor(d, dist(ord$points))
## Correction with negative eigenvalues:
cor(d, sqrt(dist(ord$points)^2 - dist(ord$negaxes)^2))
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