Package ‘ecodist’

October 30, 2023

Version 2.1.3
Date 2023-10-30
Title Dissimilarity-Based Functions for Ecological Analysis
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Depends R (>= 3.0.0)
LazyData true
Imports stats, graphics, igraph
Suggests knitr, testthat, markdown
VignetteBuilder knitr
License GPL (>= 2)
BugReports https://github.com/phiala/ecodist/issues
NeedsCompilation yes
Repository CRAN
Date/Publication 2023-10-30 15:40:02 UTC

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**Description**

Dissimilarity-based analysis functions including ordination and Mantel test functions, intended for use with spatial and community data.
Details

This package contains well-established dissimilarity-based ecological analyses, such as nmds and mantel, and experimental/research analyses such as xmantel. Helper functions such as crosstab and cor2m facilitate analysis of community data.

Because many of the analyses are time-consuming, this package includes worked examples that can be loaded using data().

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addord

Example for pmgram

Example for pmgram

Author(s)
Sarah Goslee and Dean Urban
Maintainer: Sarah Goslee <Sarah.Goslee@ars.usda.gov>

addord  \textit{Fit new points to an existing NMDS configuration.}

Description
Uses a brute force algorithm to find the location for each new point that minimizes overall stress.

Usage
addord(origconf, fulldat, fulldist, isTrain, bfstep = 10, maxit = 50, epsilon = 1e-12)

Arguments
origconf  The original ordination configuration.
fulldat  The dataset containing original and new points.
fulldist  A dissimilarity matrix calculated on fulldat.
isTrain  A boolean vector of length nrow(fulldat) indicating which rows were training data used in determining origconf (TRUE), or are new points (FALSE).
bfstep  A tuning parameter for the brute force algorithm describing the size of grid to use.
maxit  The maximum number of iterations to use.
epsilon  Tolerance value for convergence.

Details
A region comprising the original ordination configuration plus one standard deviation is divided into a grid of bfstep rows and columns. For a new point, the grid cell with the lowest stress is identified. That cell is divided into a finer grid, and the lowest-stress cell identified. This process is repeated up to maxit times, or until stress changes less than epsilon.

Value
fullfitconf  The new ordination configuration containing training and new points.
stress  The stress value for each point.
isTrain  The boolean vector indicating training set membership, for reference.
Author(s)

Sarah Goslee

Examples

data(iris)
iris.d <- dist(iris[,1:4])

### nmds() is timeconsuming, so this was generated
### in advance and saved.
### set.seed(1234)
### iris.nmds <- nmds(iris.d, nits=20, mindim=1, maxdim=4)
### save(iris.nmds, file="ecodist/data/iris.nmds.rda")
data(iris.nmds)

# examine fit by number of dimensions
plot(iris.nmds)

# choose the best two-dimensional solution to work with
iris.nmin <- min(iris.nmds, dims=2)

# rotate the configuration to maximize variance
iris.rot <- princomp(iris.nmin)$scores

# rotation preserves distance apart in ordination space
cor(dist(iris.nmin), dist(iris.rot))

# fit the data to the ordination as vectors
### vf() is timeconsuming, so this was generated
### in advance and saved.
### set.seed(1234)
### iris.vf <- vf(iris.nmin, iris[,1:4], nperm=1000)
### save(iris.vf, file="ecodist/data/iris.vf.rda")
data(iris.vf)

# repeat for the rotated ordination
### vf() is timeconsuming, so this was generated
### in advance and saved.
### set.seed(1234)
### iris.vfrot <- vf(iris.rot, iris[,1:4], nperm=1000)
### save(iris.vfrot, file="ecodist/data/iris.vfrot.rda")
data(iris.vfrot)

par(mfrow=c(1,2))
plot(iris.nmin, col=as.numeric(iris$Species), pch=as.numeric(iris$Species), main="NMDS")
plot(iris.rot, col=as.numeric(iris$Species), pch=as.numeric(iris$Species), main="Rotated NMDS")
plot(iris.vf)
plot(iris.vfrot)

######## addord example

# generate new data points to add to the ordination
# this might be new samples, or a second dataset

```r
iris.new <- structure(list(Sepal.Length = c(4.6, 4.9, 5.4, 5.2, 6, 6.5, 6, 6.8, 7.3), Sepal.Width = c(3.2, 3.5, 3.6, 2.3, 2.8, 3, 2.7, 3.1, 3.2), Petal.Length = c(1.2, 1.5, 1.5, 3.5, 4.1, 4.2, 4.8, 5, 5.7), Petal.Width = c(0.26, 0.26, 0.26, 1.2, 1.3, 1.4, 1.8, 2, 2), Species = structure(c(1L, 1L, 1L, 2L, 2L, 2L, 3L, 3L, 3L), .Label = c("setosa", "versicolor", "virginica"), class = "factor")), .Names = c("Sepal.Length", "Sepal.Width", "Petal.Length", "Petal.Width", "Species"), class = "data.frame", row.names = c(NA, -9L))
```

# provide a dist object containing original and new data
# provide a logical vector indicating which samples were used to
# construct the original configuration

```r
iris.full <- rbind(iris, iris.new)
all.d <- dist(iris.full[,1:4])
is.orig <- c(rep(TRUE, nrow(iris)), rep(FALSE, nrow(iris.new)))

### addord() is timeconsuming, so this was generated
### in advance and saved.
### set.seed(1234)
### iris.fit <- addord(iris.nmin, iris.full[,1:4], all.d, is.orig, maxit=100)
### save(iris.fit, file="ecodist/data/iris.fit.rda")
data(iris.fit)

plot(iris.fit$conf, col=iris.full$Species, pch=c(18, 4)[is.orig + 1], xlab="NMDS 1", ylab="NMDS 2")
title("Demo: adding points to an ordination")
legend("bottomleft", c("Training set", "Added point"), pch=c(4, 18))
legend("topright", levels(iris$Species), fill=1:3)
```

---

**bcdist**  
*Bray-Curtis distance*

**Description**

Returns the Bray-Curtis (also known as Sorenson, 1 - percent similarity) pairwise distances for the objects in the data. It is duplicated by functionality within `distance` but remains for backward compatibility and because it is substantially faster.

**Usage**

```r
bcdist(x, rmzero = FALSE)
```

**Arguments**

- `x`  
  matrix or data frame with rows as samples and columns as variables (such as species). Distances will be calculated for each pair of rows.
If rmzero=TRUE, empty rows will be removed from the data before distances are calculated. Otherwise, the distance between two empty rows is assumed to be 0 (the default).

Value

This function returns a column-order lower-triangular distance matrix. The returned object has an attribute, Size, giving the number of objects, that is, nrow(x). The length of the vector that is returned is nrow(x)*(nrow(x)-1)/2.

Author(s)

Sarah Goslee

See Also

dist, distance

Examples

data(graze)
system.time(graze.bc <- bcdist(graze[, -c(1:2)]))
# equivalent to but much faster than:
system.time(graze.bc2 <- distance(graze[, -c(1:2)], "bray-curtis"))

all.equal(graze.bc, graze.bc2)

bump

Nine-bump spatial pattern

Description

A two-dimensional artificial "landscape" illustrating the kind of spatial pattern that might be seen across mountain peaks.

Usage

data(bump)

Format

The format is: int [1:25, 1:25] 2 2 2 2 2 2 2 2 2 2 ... - attr(*, "dimnames")=List of 2 ..$ : chr [1:25] "1" "3" "5" "7" ... ..$ : chr [1:25] "V1" "V3" "V5" "V7" ...

Author(s)

Sarah Goslee
See Also

bump.pmgram, pmgram

Examples

data(bump)
image(bump)

bump.pmgram         Nine-bump spatial pattern

Description

An object of class mgram for use in the example for pmgram. Many of the functions in ecodist take a long time to run, so prepared examples have been included.

Usage

data(bump.pmgram)

Format

See pmgram for current format specification.

Author(s)

Sarah Goslee

See Also

bump, pmgram

Examples

data(bump)

par(mfrow=c(1, 2))
image(bump, col=gray(seq(0, 1, length=5)))

z <- as.vector(bump)
x <- rep(1:25, times=25)
y <- rep(1:25, each=25)

X <- col(bump)
Y <- row(bump)
# calculate dissimilarities for data and space
geo.dist <- dist(cbind(as.vector(X), as.vector(Y)))
value.dist <- dist(as.vector(bump))
### pgram() is time-consuming, so this was generated
### in advance and saved.
### set.seed(1234)
### bump.pmgram <- pmgram(value.dist, geo.dist, nperm=10000)
### save(bump.pmgram, file="ecodist/data/bump.pmgram.rda")

data(bump.pmgram)
plot(bump.pmgram)

---

**cor2m**  
*Two-matrix correlation table*

**Description**

Generate a correlation table between the variables of two data sets, originally for comparing species abundances and environmental variables.

**Usage**

```r
cor2m(x, y, trim = TRUE, alpha = 0.05)
```

**Arguments**

- `x`: A matrix or data frame of environmental (or other) variables matching the sites of `x`
- `y`: A matrix or data frame of species (or other) variables
- `trim`: If `trim` is `TRUE`, set ρ<critical value(alpha) to 0
- `alpha`: Alpha p-value to use with `trim`, by default 0.05

**Details**

`cor2m` generates a correlation table between the variables of two matrices. The original use case is to compare species abundances and environmental variables. It results in a data frame with species (or the first matrix) as columns and environmental variables (or the second matrix) as rows, so it’s easy to scan. Correlations less than a user-specified alpha (0.05 by default) can be set to NA. `cor2m(x, y, trim=FALSE)` is equivalent to `cor(x, y)`.

**Value**

Returns a data frame of correlations between the variables of 2 data frames.
Author(s)

Dean Urban

Examples

data(graze)
speciesdata <- graze[, 3:7]
envdata <- graze[, 1:2]
sppenv.cor <- cor2m(envdata, speciesdata)
print(sppenv.cor, na.print="")

corgen

Generate correlated data

Description

Generate correlated data of a given length.

Usage

corgen(len, x, r, population = FALSE, epsilon = 0)

Arguments

len	Length of vectors.

x	Independent data. If x is specified, the population parameter is automatically set to TRUE.

r	Desired correlation between data vectors.

population	TRUE for vectors drawn from two populations with correlation r, otherwise r is the sample correlation.

epsilon	Desired tolerance.

Details

Either x or len must be specified. If epsilon = 0, it has no effect, otherwise the sampling process is repeated until the sample correlation is within epsilon of r. This option allows the production of exactly-correlated data, within the limits of epsilon. Setting epsilon > 0 invalidates the population setting; data will be correlated within that range, rather than sampled from that population. If epsilon = 0, it has no effect, otherwise the sampling process is repeated until the sample correlation is within epsilon of r. This option allows the production of exactly-correlated data, within the limits of epsilon. Setting epsilon > 0 invalidates the population setting; data will be correlated within that range, rather than sampled from that population. If epsilon = 0, it has no effect, otherwise the sampling process is repeated until the sample correlation is within epsilon of r. This option allows the production of exactly-correlated data, within the limits of epsilon. Setting epsilon > 0 invalidates the population setting; data will be correlated within that range, rather than sampled from that population.
**Value**

- **x**: First data vector, either generated by corgen or given by the user.
- **y**: Second data vector.

**Author(s)**

Sarah Goslee

**Examples**

```r
# create two random variables of length 100 with correlation # of 0.10 +/- 0.01
xy <- corgen(len=100, r=.1, epsilon=0.01)
with(xy, cor(x, y))

# create two random variables of length 100 drawn from a population with # a correlation of -0.82
xy <- corgen(len=100, r=-0.82, population=TRUE)
with(xy, cor(x, y))

# create a variable y within 0.01 of the given correlation to x
x <- 1:100
y <- corgen(x=x, r=.5, epsilon=.01)$y
cor(x, y)
```

---

**crosstab**

**Data formatting**

**Description**

Converts field data of the form site, species, observation into a site by species data frame.

**Usage**

```r
crosstab(rowlab, collab, values, type = "sum", data, allrows, allcols,
na.as.0 = TRUE, check.names = TRUE, ...)
```

**Arguments**

- **rowlab**: row labels, e.g. site names.
- **collab**: column labels, e.g. species names.
- **values**: data values.
- **data**: optional data frame from which to take rowlab, collab and/or values.
- **type**: function to use to combine data, one of "sum" (default), "min", "max", "mean", "count".
allrows  optional, list of all desired row names that may not appear in rowlab.
allcols  optional, list of all desired column names that may not appear in collab.
na.as.0  if TRUE, all NA values are replaced with 0.
check.names  if FALSE, data frame names are not checked for syntactic validity, so that they match the input categories. Otherwise make.names() is used to adjust them.
...  optional arguments to the function specified in type, such as na.rm=TRUE

Details

Field data are often recorded as a separate row for each site-species combination. This function reformats such data into a data frame for further analysis based on unique row and column labels. The three vectors should all be the same length (including duplicates). The three vectors may also be provided as names of columns in the data frame specified by the data argument.

If allrows or allcols exists, rows and/or columns of zeros are inserted for any elements of allrows/allcols not present in rowlab/collab.

If values is missing the number of occurrences of combinations of rowlab and collab will be returned. Thus, crosstab(rowlab, collab) is equivalent to table(rowlab, collab).

If type is "count", the unique combinations of rowlab, collab and values will be returned.

Value

data frame with rowlab as row headings, collab as columns, and values as the data.

Author(s)

Sarah Goslee

Examples

# Make a random example
plotnames <- rep(1:5, each = 6)
speciesnames <- rep(c("A", "B", "C"), 10)
freqdata <- runif(30)

# number of samples of each species and plot
crosstab(plotnames, speciesnames)

# can use the data argument
speciesdata <- data.frame(plots = plotnames, species = speciesnames, freq = freqdata, stringsAsFactors=FALSE)

# mean frequency by species and plot
crosstab(plots, species, freq, data=speciesdata, type="mean")

# can specify additional possible row or column levels
crosstab(plots, species, freq, data=speciesdata, type="mean", allcols=LETTERS[1:5])
Description

Returns NULL for the dimensions of a distance object.

Usage

```r
## S3 method for class 'dist'
dim(x)
```

Arguments

- `x` object of class `dist`

Details

The spdep package overwrites the base R behavior of `dim.dist()` to return `c(n, n)` where `n` is the size of the full matrix. The base R behavior returns NULL. This function restores base R behavior within ecodist, because otherwise spdep being loaded breaks ecodist functionality.

Value

NULL

Author(s)

Sarah Goslee

Examples

```r
data(graze)
dim(dist(graze))
```
distance

Calculate dissimilarity/distance metrics

distance

Description

This function calculates a variety of dissimilarity or distance metrics. Although it duplicates the functionality of dist() and bcdist(), it is written in such a way that new metrics can easily be added. distance() was written for extensibility and understandability, and is not necessarily an efficient choice for use with large matrices.

Usage

distance(x, method = "euclidean", sprange=NULL, spweight=NULL, icov, inverted = FALSE)

Arguments

x
  matrix or data frame with rows as samples and columns as variables (such as species). Distances will be calculated for each pair of rows.

method
  Currently 7 dissimilarity metrics can be calculated: "euclidean", "bray-curtis", "manhattan", "mahalanobis" (squared Mahalanobis distance), "jaccard", "difference", "sorensen", "gower", "modgower10" (modified Gower, base 10), "modgower2" (modified Gower, base 2). Partial matching will work for selecting a method.

sprange
  Gower dissimilarities offer the option of dividing by the species range. If sprange=NULL no range is used. If sprange is a vector of length nrow(x) it is used for standardizing the dissimilarities.

spweight
  Euclidean, Manhattan, and Gower dissimilarities allow weighting. If spweight=NULL, no weighting is used. If spweight="absence", then W=0 if both species are absent and 1 otherwise, thus deleting joint absences.

icov
  Optional covariance matrix; only used if method="mahalanobis" since Mahalanobis distance requires calculating the variance-covariance matrix for the entire dataset. Providing icov directly makes it possible to calculate distances for a subset of the full dataset.

inverted
  If TRUE, the optional covariance matrix for method="mahalanobis" is not inverted before solving. Providing an inverted matrix may speed up calculations.

Value

Returns a lower-triangular distance matrix as an object of class "dist".

Author(s)

Sarah Goslee

See Also

dist, bcdist
Examples

```r
data(iris)
iris.bc <- distance(iris[, 1:4], "bray-curtis")

# The effect of specifying icov:

# calculate Mahalanobis distance for the full iris dataset
iris.md <- full(distance(iris[, 1:4], "mahal"))
iris.md[1, 2] # Mahalanobis distance between samples 1 and 2

# calculate Mahalanobis for just one species
setosa.md <- full(distance(iris[iris$Species == "setosa", 1:4], "mahal"))
setosa.md[1, 2] # Mahalanobis distance between samples 1 and 2

# use the covariance matrix for the full dataset to scale for one species
setosa.scaled.md <- full(distance(iris[iris$Species == "setosa", 1:4], "mahal", icov=var(iris[,1:4])))
setosa.scaled.md[1, 2] # Mahalanobis distance between samples 1 and 2
```

---

**fixdmat**

Distance matrix conversion

Description

Convert a row-order lower-triangular distance matrix to a full symmetric matrix.

Usage

```r
fixdmat(v)
```

Arguments

- `v` lower-triangular distance matrix in row order.

Details

R distance functions such as dist and bcdist return a lower triangular distance matrix in column order. Some other programs return the lower triangular matrix in row order. To use this matrix in R functions, it must be converted from row order to column order.

Value

full symmetric distance matrix.

Author(s)

Sarah Goslee
See Also

lower, full

Examples

```r
x.vec <- seq_len(6)
x.vec

# Make an R-style column order symmetric matrix
full(x.vec)

# Extract the lower triangle from a symmetric matrix
# in column order
lower(full(x.vec))

# Convert to or from a row order symmetric matrix
fixdmat(x.vec)
lower(fixdmat(x.vec))

fixdmat(c(1, 2, 4, 3, 5, 6))
```

Description

Convert a column order distance matrix to a full symmetric matrix.

Usage

```r
full(v)
```

Arguments

- `v`: lower-triangular column order distance matrix.

Details

Converts a column order lower-triangular distance matrix as written by R functions into a symmetric matrix. Note that `lower()` used on a 1x1 matrix will return the single element, which may not be the correct behavior in all cases, while `full()` used on a single element will return a 2x2 matrix.

Value

full symmetric matrix.

Author(s)

Sarah Goslee
See Also

lower, fixdmat

Examples

# Given a vector:
x.vec <- seq_len(6)
x.vec

# Make an R-style column order symmetric matrix
full(x.vec)

# Extract the lower triangle from a symmetric matrix
# in column order
lower(full(x.vec))

# Convert to or from a row order symmetric matrix
fixdmat(x.vec)
lower(fixdmat(x.vec))

fixdmat(c(1, 2, 4, 3, 5, 6))

---

graze

*Site information and grazed vegetation data.*

Description

This data frame contains site location, landscape context and dominant plant species abundances for 25 of the plant species found in 50 grazed pastures in the northeastern United States. Elements are the mean values for canopy cover for ten 0.5 x 2 m quadrats.

Usage

data(graze)

Format

A data frame with 50 observations on the following 25 variables.

- sitelocation: Site location along a geographic gradient.
- forestpct: Percentage forest cover within a 1-km radius.
- ACM12: Percentage canopy cover.
- ANOD: Percentage canopy cover.
- ASSY: Percentage canopy cover.
- BRIN2: Percentage canopy cover.
- CIAR4: Percentage canopy cover.
CIIN  Percentage canopy cover.
CIVU  Percentage canopy cover.
DAGL  Percentage canopy cover.
ELRE4 Percentage canopy cover.
GAMO  Percentage canopy cover.
LOAR10 Percentage canopy cover.
LOC06  Percentage canopy cover.
LOPE  Percentage canopy cover.
OXST  Percentage canopy cover.
PLMA2  Percentage canopy cover.
POPR  Percentage canopy cover.
PRVU  Percentage canopy cover.
RAAC3  Percentage canopy cover.
RUCR  Percentage canopy cover.
SORU2  Percentage canopy cover.
STGR  Percentage canopy cover.
TAOF  Percentage canopy cover.
TRPR2  Percentage canopy cover.
TRRE3  Percentage canopy cover.
VEOF2  Percentage canopy cover.

Details

Site locations fall along a southwest-northeast transect through the northeastern United States. This is a synthetic gradient calculated from latitude and longitude. Forest landcover is taken from the USGS 1992 National Land Cover Dataset. All forest classes were combined, and the percentage within 1 km of the sample sites was calculated using a GIS.

Author(s)

Sarah Goslee

Source

Details of these data are available in Tracy and Sanderson (2000) and Goslee and Sanderson (2010). The 1992 NLCD data can be obtained from http://www.mrlc.gov/. Species codes are from http://plants.usda.gov (2010).

References


Examples

data(graze)

iris.fit

Example of adding to an ordination

Description

A fitted ordination for use in the example for `addord`. Many of the functions in ecodist take a long time to run, so prepared examples have been included.

Usage

data(iris.fit)

Format

The format of this object is a list with:
- `X1, X2, etc`: ordination configuration: coordinates for each point.
- `stress`: goodness of fit for each point.
- `isTrain`: logical vector indicating whether each point was used in the original ordination.

Author(s)

Sarah Goslee

See Also

`nmds`, `addord`

Examples

```
data(iris)
iris.d <- dist(iris[,1:4])

### nmds() is timeconsuming, so this was generated
### in advance and saved.
### set.seed(1234)
### iris.nmds <- nmds(iris.d, nits=20, mindim=1, maxdim=4)
### save(iris.nmds, file="ecodist/data/iris.nmds.rda")
data(iris.nmds)

# choose the best two-dimensional solution to work with
iris.nmin <- min(iris.nmds, dims=2)

# generate new data points to add to the ordination
# this might be new samples, or a second dataset
iris.new <- structure(list(Sepal.Length = c(4.6, 4.9, 5.4, 5.2, 6, 6.5, 6, 6.8, 7.3),
                           Sepal.Width = c(3.2, 3.5, 3.6, 2.3, 2.8, 3, 2.7, 3.1, 3.4)),
```
# provide a dist object containing original and new data
# provide a logical vector indicating which samples were used to
# construct the original configuration

iris.full <- rbind(iris, iris.new)
all.d <- dist(iris.full[,1:4])
is.orig <- c(rep(TRUE, nrow(iris)), rep(FALSE, nrow(iris.new)))

### addord() is timeconsuming, so this was generated
### in advance and saved.
### set.seed(1234)
### iris.fit <- addord(iris.nmin, iris.full[,1:4], all.d, is.orig, maxit=100)
### save(iris.fit, file="ecodist/data/iris.fit.rda")
data(iris.fit)

plot(iris.fit$conf, col=iris.full$Species, pch=c(18, 4)[is.orig + 1],
     xlab="NMDS 1", ylab="NMDS 2")
title("Demo: adding points to an ordination")
legend("bottomleft", c("Training set", "Added point"), pch=c(4, 18))
legend("topright", levels(iris$Species), fill=1:3)
### iris.vf

#### Example for vector fitting on ordination

**Description**

An object of class vf for use in the examples for nmds and vf. Many of the functions in ecodist take a long time to run, so prepared examples have been included.

**Usage**

```r
data(iris.vf)
```

**Format**

See vf for current format specification.

**Author(s)**

Sarah Goslee

**See Also**

nmds, vf
**Examples**

```r
data(iris)
iris.d <- dist(iris[,1:4])

### nmds() is timeconsuming, so this was generated
### in advance and saved.
### set.seed(1234)
### iris.nmds <- nmds(iris.d, nits=20, mindim=1, maxdim=4)
### save(iris.nmds, file="ecodist/data/iris.nmds.rda")
data(iris.nmds)

# examine fit by number of dimensions
plot(iris.nmds)

# choose the best two-dimensional solution to work with
iris.nmin <- min(iris.nmds, dims=2)

# rotate the configuration to maximize variance
iris.rot <- princomp(iris.nmin)$scores

# rotation preserves distance apart in ordination space
cor(dist(iris.nmin), dist(iris.rot))

# fit the data to the ordination as vectors
### vf() is timeconsuming, so this was generated
### in advance and saved.
### set.seed(1234)
### iris.vf <- vf(iris.nmin, iris[,1:4], nperm=1000)
### save(iris.vf, file="ecodist/data/iris.vf.rda")
data(iris.vf)

plot(iris.nmin, col=as.numeric(iris$Species), pch=as.numeric(iris$Species), main="NMDS")
plot(iris.vf)
```

---

**iris.vfrot**  
*Example for vector fitting on rotated ordination*

**Description**

An object of class `vf` for use in the examples for `nmds` and `vf`. Many of the functions in ecodist take a long time to run, so prepared examples have been included.

**Usage**

```r
data(iris.vfrot)
```

**Format**

See `vf` for current format specification.
Examples

data(iris)
iris.d <- dist(iris[,1:4])

### nmds() is timeconsuming, so this was generated
### in advance and saved.
### set.seed(1234)
### iris.nmds <- nmds(iris.d, nits=20, mindim=1, maxdim=4)
### save(iris.nmds, file="ecodist/data/iris.nmds.rda")
data(iris.nmds)

# examine fit by number of dimensions
plot(iris.nmds)

# choose the best two-dimensional solution to work with
iris.nmin <- min(iris.nmds, dims=2)

# rotate the configuration to maximize variance
iris.rot <- princomp(iris.nmin)$scores

# rotation preserves distance apart in ordination space
cor(dist(iris.nmin), dist(iris.rot))

# fit the data to the ordination as vectors
### vf() is timeconsuming, so this was generated
### in advance and saved.
### set.seed(1234)
### iris.vf <- vf(iris.nmin, iris[,1:4], nperm=1000)
### save(iris.vf, file="ecodist/data/iris.vf.rda")
data(iris.vf)

# repeat for the rotated ordination
### vf() is timeconsuming, so this was generated
### in advance and saved.
### set.seed(1234)
### iris.vfrot <- vf(iris.rot, iris[,1:4], nperm=1000)
### save(iris.vfrot, file="ecodist/data/iris.vfrot.rda")
data(iris.vfrot)

par(mfrow=c(1,2))
plot(iris.nmin, col=as.numeric(iris$Species), pch=as.numeric(iris$Species), main="NMDS")
plot(iris.vf)
plot(iris.rot, col=as.numeric(iris$Species), pch=as.numeric(iris$Species), main="Rotated NMDS")
plot(iris.vfrot)


**lower**

*Lower-triangular matrix*

**Description**

Convert a symmetric distance matrix to a column order lower triangular matrix.

**Usage**

```r
lower(m)
```

**Arguments**

- `m` a symmetric distance matrix.

**Details**

Converts a symmetric matrix, for example a dissimilarity matrix, into a column order lower-triangular matrix. This may be useful to format the input for certain clustering and ordination functions. Note that `lower()` used on a 1x1 matrix will return the single element, which may not be the correct behavior in all cases, while `full()` used on a single element will return a 2x2 matrix.

**Value**

column order lower triangular matrix.

**Author(s)**

Sarah Goslee

**See Also**

`full`, `fixdmat`

**Examples**

```r
x.vec <- seq_len(6)
x.vec

# Make an R-style column order symmetric matrix
full(x.vec)

# Extract the lower triangle from a symmetric matrix
# in column order
lower(full(x.vec))

# Convert to or from a row order symmetric matrix
```
mantel

\begin{verbatim}
fixdmat(x.vec)
lower(fixdmat(x.vec))
fixdmat(c(1, 2, 4, 3, 5, 6))
\end{verbatim}

---

### Mantel test

**Description**

Simple and partial Mantel tests, with options for ranked data, permutation tests, and bootstrapped confidence limits.

**Usage**

\begin{verbatim}
mantel(formula = formula(data), data, nperm = 1000,
       mrank = FALSE, nboot = 500, pboot = 0.9, cboot = 0.95)
\end{verbatim}

**Arguments**

- `formula`: formula describing the test to be conducted. For this test, \( y \sim x \) will perform a simple Mantel test, while \( y \sim x + z_1 + z_2 + z_3 \) will do a partial Mantel test of the relationship between \( x \) and \( y \) given \( z_1, z_2, z_3 \). All variables can be either a distance matrix of class dist or vectors of dissimilarities.
- `data`: an optional dataframe containing the variables in the model as columns of dissimilarities. By default the variables are taken from the current environment.
- `nperm`: number of permutations to use. If set to 0, the permutation test will be omitted.
- `mrank`: if this is set to FALSE (the default option), Pearson correlations will be used. If set to TRUE, the Spearman correlation (correlation ranked distances) will be used.
- `nboot`: number of iterations to use for the bootstrapped confidence limits. If set to 0, the bootstrapping will be omitted.
- `pboot`: the level at which to resample the data for the bootstrapping procedure.
- `cboot`: the level of the confidence limits to estimate.

**Details**

If only one independent variable is given, the simple Mantel r (r12) is calculated. If more than one independent variable is given, the partial Mantel r (rxy|x1 ...) is calculated by permuting one of the original dissimilarity matrices. The bootstrapping is actually resampling without replacement, because duplication of samples is not useful in a dissimilarity context (the dissimilarity of a sample with itself is zero). Resampling within dissimilarity values is inappropriate, just as for permutation.
Value

- `mantelr` Mantel coefficient.
- `pval1` one-tailed p-value (null hypothesis: \( r \leq 0 \)).
- `pval2` one-tailed p-value (null hypothesis: \( r \geq 0 \)).
- `pval3` two-tailed p-value (null hypothesis: \( r = 0 \)).
- `llim` lower confidence limit.
- `ulim` upper confidence limit.

Author(s)

Sarah Goslee

References


See Also

- `mgram`
- `mgroup`

Examples

data(graze)

grasses <- graze[, colnames(graze) %in% c("DAGL", "LOAR10", "LOPE", "POPR")]
legumes <- graze[, colnames(graze) %in% c("LOCO6", "TRPR2", "TRRE3")]

grasses.bc <- bcdist(grasses)
legumes.bc <- bcdist(legumes)

space.d <- dist(graze$sitelocation)
forest.d <- dist(graze$forestpct)

# Mantel test: is the difference in forest cover between sites
# related to the difference in grass composition between sites?
mantel(grasses.bc ~ forest.d)

# Mantel test: is the geographic distance between sites
# related to the difference in grass composition between sites?
mantel(grasses.bc ~ space.d)
# Partial Mantel test: is the difference in forest cover between sites related to the difference in grass composition once the linear effects of geographic distance are removed?
mantel(grasses.bc ~ forest.d + space.d)

# Mantel test: is the difference in forest cover between sites related to the difference in legume composition between sites?
mantel(legumes.bc ~ forest.d)

# Mantel test: is the geographic distance between sites related to the difference in legume composition between sites?
mantel(legumes.bc ~ space.d)

# Partial Mantel test: is the difference in forest cover between sites related to the difference in legume composition once the linear effects of geographic distance are removed?
mantel(legumes.bc ~ forest.d + space.d)

# Is there nonlinear pattern in the relationship with geographic distance?
par(mfrow=c(2, 1))
plot(mgram(grasses.bc, space.d, nclass=8))
plot(mgram(legumes.bc, space.d, nclass=8))

---

**Mantel correlogram**

### Description
Calculates simple Mantel correlograms.

### Usage
```r
mgram(species.d, space.d, breaks, nclass, stepsize, equiprobable = FALSE, nperm = 1000,
      mrank = FALSE, nboot = 500, pboot = 0.9, cboot = 0.95,
      alternative = "two.sided", trace = FALSE)
```

### Arguments
- `species.d`: lower-triangular dissimilarity matrix.
- `space.d`: lower-triangular matrix of geographic distances.
- `breaks`: locations of class breaks. If specified, overrides nclass and stepsize.
- `nclass`: number of distance classes. If not specified, Sturge's rule will be used to determine an appropriate number of classes.
- `stepsize`: width of each distance class. If not specified, nclass and the range of space.d will be used to calculate an appropriate default.
equivarable  if TRUE, create nclass classes of equal number of distances; if FALSE, create nclass classes of equal width

ncolm  number of permutations to use. If set to 0, the permutation test will be omitted.

nrank  if this is set to FALSE (the default option), Pearson correlations will be used. If set to TRUE, the Spearman correlation (correlation ranked distances) will be used.

nboot  number of iterations to use for the bootstrapped confidence limits. If set to 0, the bootstrapping will be omitted.

pboot  the level at which to resample the data for the bootstrapping procedure.

clim  the level of the confidence limits to estimate.

alternative  default is "two.sided", and returns p-values for H0: rM = 0. The alternative is "one.sided", which returns p-values for H0: rM <= 0.

trace  if TRUE, returns progress indicators.

Details

This function calculates Mantel correlograms, and tests the hypothesis that the mean compositional dissimilarity within a distance class differs from the mean of all the other distance classes combined. The Mantel correlogram is essentially a multivariate autocorrelation function. The Mantel r represents the dissimilarity in variable composition (often species composition) at a particular lag distance, and significance is tested in reference to all distance classes.

Value

Returns an object of class mgram, which is a list with two elements. mgram is a matrix with one row for each distance class and 6 columns:

lag  midpoint of the distance class.

ngroup  number of distances in that class.

mantelr  Mantel r value.

pval  p-value for the test chosen.

llim  lower bound of confidence limit for mantelr.

ulim  upper bound of confidence limit for mantelr.

resids is NA for objects calculated by mgram().

Author(s)

Sarah Goslee

References


See Also

mantel, plot.mgram, pmgram
Examples

# generate a simple surface
x <- matrix(1:10, nrow=10, ncol=10, byrow=FALSE)
y <- matrix(1:10, nrow=10, ncol=10, byrow=TRUE)
z <- x + 3*y
image(z)

# analyze the pattern of z across space
space <- cbind(as.vector(x), as.vector(y))
z <- as.vector(z)
space.d <- distance(space, "eucl")
z.d <- distance(z, "eucl")
z.mgram <- mgram(z.d, space.d, nperm=0)
plot(z.mgram)

#
data(graze)
space.d <- dist(graze$sitelocation)
forest.d <- dist(graze$forestpct)
grasses <- graze[, colnames(graze) %in% c("DAGL", "LOAR10", "LOPE", "POPR")]
legumes <- graze[, colnames(graze) %in% c("LOCO6", "TRPR2", "TRRE3")]
grasses.bc <- bcdist(grasses)
legumes.bc <- bcdist(legumes)

# Does the relationship of composition with distance vary for
# grasses and legumes?
par(mfrow=c(2, 1))
plot(mgram(grasses.bc, space.d, nclass=8))
plot(mgram(legumes.bc, space.d, nclass=8))


---

mgroup  

Mantel test for groups

Description

Mantel test across one or more group contrasts.

Usage

mgroup(edist, groups, nperm = 1000, mrank = FALSE)

Arguments

edist  
a dist object or lower triangular distance vector.
groups a vector of group memberships (numeric, character, or factor), or a matrix or
data frame with columns describing multiple sets of groups.
nperm number of permutations to use. If set to 0, the permutation test will be omitted.
mrank if this is set to FALSE (the default option), Pearson correlations will be used. If set to TRUE, the Spearman correlation (correlation ranked distances) will be used.

Details
mgroup returns the Mantel correlations for group contrast matrices computed from cluster groups
across a range of clustering levels.

Value
nclust Number of groups tested.
mantelr Mantel coefficient.
pval1 one-tailed p-value (null hypothesis: r <= 0).

Author(s)
Sarah Goslee

References

See Also
mantel

Examples

# Using a model matrix to test group membership
data(iris)
iris.d <- dist(iris[,1:4])
mgroup(iris.d, iris[,5])

# clustering-based example
data(graze)
graze.d <- dist(graze[, -c(1:2)])
graze.hclust <- hclust(graze.d)

clust.groups <- data.frame(  
k2 = cutree(graze.hclust, k = 2),
  k4 = cutree(graze.hclust, k = 4),
  k6 = cutree(graze.hclust, k = 6),
  k8 = cutree(graze.hclust, k = 8))
min.nmds

mgroup(graze.d, clust.groups, nperm=1000)

---

min.nmds

Find minimum stress configuration

Description

Finds minimum stress configuration from output of nmds()

Usage

## S3 method for class 'nmds'
min(..., na.rm = FALSE, dims = 2)
nmds.min(x, dims = 2)

Arguments

... output from nmds()

x output from nmds()

dims desired dimensionality of result. If dims = 0 then the overall minimum stress configuration is chosen. By default, the best two-dimensional configuration is returned.

na.rm Not used; there should be no NA values in a NMDS configuration.

Details

For back-compatibility, the nmds.min function remains.

Value

Configuration of minimum stress ordination (dataframe of coordinates). The stress and r^2 for the minimum stress configuration are stored as attributes.

Author(s)

Sarah Goslee

See Also

nmds
Examples

data(iris)
iris.d <- dist(iris[,1:4])

### nmds() is timeconsuming, so this was generated
### in advance and saved.
### set.seed(1234)
### iris.nmds <- nmds(iris.d, nits=20, mindim=1, maxdim=4)
### save(iris.nmds, file="ecodist/data/iris.nmds.rda")
data(iris.nmds)

# examine fit by number of dimensions
plot(iris.nmds)

# choose the best two-dimensional solution to work with
iris.nmin <- min(iris.nmds, dims=2)

MRM

Multiple Regression on distance Matrices

Description

Multiple regression on distance matrices (MRM) using permutation tests of significance for regression coefficients and R-squared.

Usage

MRM(formula = formula(data), data, nperm = 1000,
method = "linear", mrank = FALSE)

Arguments

formula formula describing the test to be conducted.
data an optional dataframe containing the variables in the model as columns of dissimilarities. By default the variables are taken from the current environment.
nperm number of permutations to use. If set to 0, the permutation test will be omitted.
mrank if this is set to FALSE (the default option), Pearson correlations will be used. If set to TRUE, the Spearman correlation (correlation ranked distances) will be used.
method if "linear", the default, uses multiple regression analysis. If "logistic", performs logistic regression with appropriate permutation testing. Note that this may be substantially slower.
Details

Performs multiple regression on distance matrices following the methods outlined in Legendre et al. 1994. Specifically, the permutation test uses a pseudo-t test to assess significance, rather than using the regression coefficients directly.

Value

c coef A matrix with regression coefficients and associated p-values from the permutation test (using the pseudo-t of Legendre et al. 1994).

r.squared Regression R-squared and associated p-value from the permutation test (linear only).

F.test F-statistic and p-value for overall F-test for lack of fit (linear only).

dev Residual deviance, degrees of freedom, and associated p-value (logistic only).

Author(s)

Sarah Goslee

References


See Also

mantel

Examples

data(graze)

# Abundance of this grass is related to forest cover but not location
MRM(dist(LOAR10) ~ dist(sitelocation) + dist(forestpct), data=graze, nperm=10)

# Abundance of this legume is related to location but not forest cover
MRM(dist(TRRE3) ~ dist(sitelocation) + dist(forestpct), data=graze, nperm=10)

# Compare to presence/absence of grass LOAR10 using logistic regression
LOAR10.presence <- ifelse(graze$LOAR10 > 0, 1, 0)
MRM(dist(LOAR10.presence) ~ dist(sitelocation) + dist(forestpct),
   data=graze, nperm=10, method="logistic")
Non-metric multidimensional scaling

Usage

`nmds(dmat, mindim = 1, maxdim = 2, nits = 10, iconf, epsilon = 1e-12, maxit = 500, trace = FALSE)`

Arguments

dmat: lower-triangular dissimilarity matrix.
mindim: optional, minimum number of dimensions to use.
maxdim: optional, maximum number of dimensions to use.
nits: optional, number of separate ordinations to use.
iconf: optional, initial configuration. If not specified, then a random configuration is used.
epsilon: optional, acceptable difference in stress.
maxit: optional, maximum number of iterations.
trace: if TRUE, will write progress indicator to the screen.

Details

The goal of NMDS is to find a configuration in a given number of dimensions which preserves rank-order dissimilarities as closely as possible. The number of dimensions must be specified in advance. Because NMDS is prone to finding local minima, several random starts must be used. Stress is used as the measure of goodness of fit. A lower stress indicates a better match between dissimilarity and ordination. As of ecodist 1.9, the stress calculation used is the same as in MASS::isoMDS. In previous versions it was monotonically related, so the same configurations were produced, but the absolute value was different.

Value

conf: list of configurations, each in the same units as the original dissimilarities.
stress: list of final stress values.
r2: total variance explained by each configuration.

The first results are for the lowest number of dimensions (total number is (mindim - maxdim + 1) * nits).

Author(s)

Sarah Goslee
nmens

References


See Also

plot.nmds, min.nmds, vf, addord

Examples

data(iris)
iris.d <- dist(iris[,1:4])

### nmds() is timeconsuming, so this was generated
### in advance and saved.
### set.seed(1234)
### iris.nmds <- nmds(iris.d, nits=20, mindim=1, maxdim=4)
### save(iris.nmds, file="ecodist/data/iris.nmds.rda")
data(iris.nmds)

# examine fit by number of dimensions
plot(iris.nmds)

# choose the best two-dimensional solution to work with
iris.nmin <- min(iris.nmds, dims=2)

# rotate the configuration to maximize variance
iris.rot <- princomp(iris.nmin)$scores

# rotation preserves distance apart in ordination space
cor(dist(iris.nmin), dist(iris.rot))

# fit the data to the ordination as vectors
### vf() is timeconsuming, so this was generated
### in advance and saved.
### set.seed(1234)
### iris.vf <- vf(iris.nmin, iris[,1:4], nperm=1000)
### save(iris.vf, file="ecodist/data/iris.vf.rda")
data(iris.vf)

# repeat for the rotated ordination
### vf() is timeconsuming, so this was generated
### in advance and saved.
### set.seed(1234)
### iris.vfrot <- vf(iris.rot, iris[,1:4], nperm=1000)
### save(iris.vfrot, file="ecodist/data/iris.vfrot.rda")
data(iris.vfrot)

par(mfrow=c(1,2))
plot(iris.min, col=as.numeric(iris$Species), pch=as.numeric(iris$Species), main="NMDS")
plot(iris.vf)
plot(iris.rot, col=as.numeric(iris$Species), pch=as.numeric(iris$Species),
      main="Rotated NMDS")
plot(iris.vfrot)

# generate new data points to add to the ordination
# this might be new samples, or a second dataset
iris.new <- structure(list(Sepal.Length = c(4.6, 4.9, 5.4, 5.2, 6, 6.5, 6,
                                  6.8, 7.3), Sepal.Width = c(3.2, 3.5, 3.6, 2.3, 2.8, 3, 2.7, 3.1,
                                  3.2), Petal.Length = c(1.2, 1.5, 1.5, 3.5, 4.1, 4.2, 4.8, 5,
                                  5.7), Petal.Width = c(0.26, 0.26, 0.26, 1.2, 1.3, 1.4, 1.8, 2,
                                  2), Species = structure(c(1L, 1L, 1L, 2L, 2L, 2L, 3L, 3L, 3L), .Label = c("setosa",
                                  "versicolor", "virginica"), class = "factor")), .Names = c("Sepal.Length",
row.names = c(NA, -9L))

# provide a dist object containing original and new data
# provide a logical vector indicating which samples were used to
# construct the original configuration
iris.full <- rbind(iris, iris.new)
all.d <- dist(iris.full[,1:4])
is.orig <- c(rep(TRUE, nrow(iris)), rep(FALSE, nrow(iris.new)))

### addord() is timeconsuming, so this was generated
### in advance and saved.
### set.seed(1234)
### iris.fit <- addord(iris.nmin, iris.full[,1:4], all.d, is.orig, maxit=100)
### save(iris.fit, file="ecodist/data/iris.fit.rda")
data(iris.fit)

plot(iris.fit$conf, col=iris.full$Species, pch=c(18, 4)[is.orig + 1],
     xlab="NMDS 1", ylab="NMDS 2")
title("Demo: adding points to an ordination")
legend("bottomleft", c("Training set", "Added point"), pch=c(4, 18))
legend("topright", levels(iris$Species), fill=1:3)

---

**pathdist**

*Graph extension of dissimilarities*

**Description**

Uses the shortest path connecting sites to estimate the distance between samples with pairwise
distances greater than `maxv`.

**Usage**

`pathdist(v, maxv = 1)`
Arguments

v  lower-triangular distance vector, possibly as produced by dist() or distance().
maxv  cutoff for distances: values greater or equal to this will be estimated from the minimum spanning tree.

Details

Pairwise samples with no species will have distances greater than a cutoff. A distance-weighted graph connecting these samples by way of intermediate samples with some species in common can be used to interpolate distances by adding up the path length connecting those samples. This function will fail if there are completely disconnected subsets.

Value

Returns a lower-triangular distance matrix.

Author(s)

Sarah Goslee

See Also

dist, distance

Examples

# samples 1 and 2, and 3 and 4, have no species in common
x <- matrix(c(1, 0, 1, 0,
              0, 1, 0, 1,
              1, 0, 0, 0,
              0, 1, 1, 1,
              1, 1, 1, 0,
              1, 0, 1, 1,
              0, 0, 1, 1), ncol = 4, byrow = TRUE)

# the maximum Jaccard distance is 1
# regardless of how different the samples are
x.jd <- dist(x, "binary")

# estimate the true distance between those pairs
# by following the shorted path along connected sites
pathdist(x.jd)
Description

Principal coordinates analysis (classical scaling).

Usage

pco(x, negvals = "zero", dround = 0)

Arguments

x

a lower-triangular dissimilarity matrix.

negvals

if = "zero" sets all negative eigenvalues to zero; if = "rm" corrects for negative eigenvalues using method 1 of Legendre and Anderson 1999.

dround

if greater than 0, attempts to correct for round-off error by rounding to that number of places.

Details

PCO (classical scaling, metric multidimensional scaling) is very similar to principal components analysis, but allows the use of any dissimilarity metric.

Value

values

eigenvalue for each component. This is a measure of the variance explained by each dimension.

vectors

eigenvectors. data frame with columns containing the scores for that dimension.

Author(s)

Sarah Goslee

See Also

princomp, nmds

Examples

data(iris)
iris.d <- dist(iris[,1:4])
iris.pco <- pco(iris.d)

# scatterplot of the first two dimensions
plot(iris.pco$vectors[,1:2], col=as.numeric(iris$Species),
      pch=as.numeric(iris$Species), main="PCO", xlab="PCO 1", ylab="PCO 2")
plot.mgram  

Plot a Mantel correlogram

Description

Plot a Mantel correlogram from an object of S3 class mgram, using solid symbols for significant values.

Usage

## S3 method for class 'mgram'
plot(x, pval = 0.05, xlab = "Distance", ylab = NULL, ...)

Arguments

x an object of class mgram
pval cut-off level for statistical significance.
xlab x-axis label.
ylab y-axis label.
... optional, any additional graphics parameters.

Value

draws a plot (graphics device must be active).

Author(s)

Sarah Goslee

See Also

mgram

Examples

# generate a simple surface
x <- matrix(1:10, nrow=10, ncol=10, byrow=FALSE)
y <- matrix(1:10, nrow=10, ncol=10, byrow=TRUE)
z <- x + 3*y
image(z)

# analyze the pattern of z across space
space <- cbind(as.vector(x), as.vector(y))
z <- as.vector(z)
space.d <- distance(space, "eucl")
z.d <- distance(z, "eucl")
z.mgram <- mgram(z.d, space.d, nperm=0)
plot(z.mgram)

#
data(graze)
space.d <- dist(graze$sitelocation)
forest.d <- dist(graze$forestpct)
grasses <- graze[, colnames(graze) %in% c("DAGL", "LOAR10", "LOPE", "POPR")]
legumes <- graze[, colnames(graze) %in% c("LOCO6", "TRPR2", "TRRE3")]
grasses.bc <- bcdist(grasses)
legumes.bc <- bcdist(legumes)

# Does the relationship of composition with distance vary for
# grasses and legumes?
par(mfrow=c(2, 1))
plot(mgram(grasses.bc, space.d, nclass=8))
plot(mgram(legumes.bc, space.d, nclass=8))

---

**plot.nmds**

*Plot information about NMDS ordination*

**Description**

Graphical display of stress and r2 for NMDS ordination along number of dimensions.

**Usage**

```r
## S3 method for class 'nmds'
plot(x, plot = TRUE, xlab = "Dimensions", ...)
```

**Arguments**

- `x` an object of S3 class `nmds`, created by `nmds()`
- `plot` optional, if TRUE a figure is produced
- `xlab` optional, label for x axis of graph
- `...` optional, other graphics parameters

**Value**

Produces a two-panel plot with stress and r2 for ordination by number of dimensions. Points show the mean value; lines indicate minimum and maximum.

**Author(s)**

Dean Urban
See Also

nmds

Examples

```r
data(iris)
iris.d <- dist(iris[,1:4])

### nmds() is timeconsuming, so this was generated
### in advance and saved.
### set.seed(1234)
### iris.nmds <- nmds(iris.d, nits=20, mindim=1, maxdim=4)
### save(iris.nmds, file="ecodist/data/iris.nmds.rda")
data(iris.nmds)

# examine fit by number of dimensions
plot(iris.nmds)

# choose the best two-dimensional solution to work with
iris.nmin <- min(iris.nmds, dims=2)
```

---

**plot.vf**

*Plots fitted vectors onto an ordination diagram*

**Description**

Add vector fitting arrows to an existing ordination plot.

**Usage**

```r
## S3 method for class 'vf'
plot(x, pval = NULL, r = NULL, cex = 0.8, ascale = 0.9, ...)
```

**Arguments**

- `x` an object of S3 class `vf`, created by `vf()`
- `pval` optional, critical p-value for choosing variables to plot
- `r` optional, minimum Mantel r for choosing variables to plot
- `cex` text size
- `ascale` optional, proportion of plot area to use when calculating arrow length
- `...` optional, other graphics parameters

**Value**

Adds arrows to an existing ordination plot. Only arrows with a p-value less than `pval` are added. By default, all variables are shown.
pmgram

Piecewise multivariate correlogram

Description

This function calculates simple and partial piecewise multivariate correlograms.

Usage

pmgram(data, space, partial, breaks, nclass, stepsize, equiprobable = FALSE, resids = FALSE, nperm = 1000)
Arguments

data  lower-triangular dissimilarity matrix. This can be either an object of class dist (treated as one column) or a matrix or data frame with one or two columns, each of which is an independent lower-triangular dissimilarity in vector form.

space  lower-triangular matrix of geographic distances.

partial  optional, lower-triangular dissimilarity matrix of ancillary data.

breaks  locations of class breaks. If specified, overrides nclass and stepsize.

nclass  number of distance classes. If not specified, Sturge’s rule will be used to determine an appropriate number of classes.

stepsize  width of each distance class. If not specified, nclass and the range of space.d will be used to calculate an appropriate default.

equiprobable  if TRUE, create nclass classes of equal number of distances; if FALSE, create nclass classes of equal width

resids  if resids=TRUE, will return the residuals for each distance class. Otherwise returns 0.

nperm  number of permutations to use. If set to 0, the permutation test will be omitted.

Details

The standard Mantel correlogram calculated by mgram tests the hypothesis that the mean compositional dissimilarity within a distance class differs from the mean of all the other distance classes combined. This function instead produces a piecewise correlogram by testing the relationship between dissimilarities within each distance class on its own, without reference to relationships across other distance classes.

This function does four different analyses: If data has 1 column and partial is missing, calculates a multivariate correlogram for data.

If data has 2 columns and partial is missing, calculates a piecewise Mantel cross-correlogram, calculating the Mantel r between the two columns for each distance class separately.

If data has 1 column and partial exists, calculates a partial multivariate correlogram based on residuals of data ~ partial.

If data has 2 columns and partial exists, does a partial Mantel cross-correlogram, calculating partial Mantel r for each distance class separately.

The Iwt statistic used for the multivariate correlograms is not the standard Mantel r. For one variable, using Euclidean distance, this metric converges on the familiar Moran autocorrelation. Like the Moran autocorrelation function, this statistic usually falls between -1 and 1, but is not bounded by those limits. Unlike the Moran function, this correlogram can be used for multivariate data, and can be extended to partial tests.

The Mantel r is used for piecewise cross-correlograms.

The comparisons in vignette("dissimilarity", package="ecodist") may help.

Value

Returns a object of class mgram, which is a list containing two objects: mgram is a matrix with one row for each distance class and 4 columns:
lag     midpoint of the distance class.
ngroup  number of distances in that class.
piece or Iwt Mantel r value or appropriate statistic (see Details).
pval    two-sided p-value.

resids is a vector of the residuals (if calculated) and can be accessed with the residuals() method.

Author(s)
Sarah Goslee

See Also
mgram, mantel, residuals.mgram, plot.mgram

Examples

data(bump)
par(mfrow=c(1, 2))
image(bump, col=gray(seq(0, 1, length=5)))

z <- as.vector(bump)
x <- rep(1:25, times=25)
y <- rep(1:25, each=25)

X <- col(bump)
Y <- row(bump)
# calculate dissimilarities for data and space
geo.dist <- dist(cbind(as.vector(X), as.vector(Y)))
value.dist <- dist(as.vector(bump))

### pmgram() is time-consuming, so this was generated
### in advance and saved.
### set.seed(1234)
### bump.pmgram <- pmgram(value.dist, geo.dist, nperm=10000)
data(bump.pmgram)
plot(bump.pmgram)

### Partial pmgram example

# generate a simple surface
# with complex nonlinear spatial pattern
x <- matrix(1:25, nrow=25, ncol=25, byrow=FALSE)
y <- matrix(1:25, nrow=25, ncol=25, byrow=TRUE)

# create z1 and z2 as functions of x, y
# and scale them to [0, 1]
z1 <- x + 3*y  
z2 <- y - cos(x)

z1 <- (z1 - min(z1)) / (max(z1) - min(z1))  
z2 <- (z2 - min(z2)) / (max(z2) - min(z2))

z12 <- (z1 + z2*2)/3

# look at patterns
layout(matrix(c(  
1, 1, 2, 2,  
1, 1, 2, 2,  
3, 3, 4, 4,  
3, 3, 5, 5), nrow=4, byrow=TRUE))

image(z1, col=gray(seq(0, 1, length=20)), zlim=c(0,1))  
image(z2, col=gray(seq(0, 1, length=20)), zlim=c(0,1))  
image(z12, col=gray(seq(0, 1, length=20)), zlim=c(0,1))

# analyze the pattern of z across space
z1 <- as.vector(z1)  
z2 <- as.vector(z2)  
z12 <- as.vector(z12)  
z1.d <- dist(z1)  
z2.d <- dist(z2)  
z12.d <- dist(z12)

space <- cbind(as.vector(x), as.vector(y))  
space.d <- dist(space)

# take partial correlogram without effects of z1
### pmgram() is time-consuming, so this was generated  
### in advance and saved.  
### set.seed(1234)  
### z.no <- pmgram(z12.d, space.d, nperm=1000, resids=FALSE)  
### save(z.no, file="ecodist/data/z.no.rda")  
data(z.no)  
plot(z.no)

# take partial correlogram of z12 given z1
### pmgram() is time-consuming, so this was generated  
### in advance and saved.  
### set.seed(1234)  
### z.z1 <- pmgram(z12.d, space.d, z2.d, nperm=1000, resids=FALSE)  
### save(z.z1, file="ecodist/data/z.z1.rda")  
data(z.z1)  
plot(z.z1)
Relativize a compositional data matrix.

Description

Relativizes the range of each column of a data frame or matrix x to 0-1. If globalmin and/or globalmax are provided, those are used to scale the columns, for instance to scale a subset to match a larger sample. If they are NA, the minimum and maximum values for each column are used.

Usage

relrange(x, globalmin = NA, globalmax = NA)

Arguments

x The data frame or matrix to be relativized.

globalmin A value other than the population minimum to be used. Should be the same length as the number of columns of x.

globalmax A value other than the population maximum to be used. Should be the same length as the number of columns of x.

Details

Relativizes the data using the minimum and maximum values. If globalmin and global max are not used, the range will be 0-1 for each variable. This can be useful for putting disparate variables to the same magnitude while keeping all non-negative values.

Value

Returns an object of the same class as x (matrix or data frame) with the columns rescaled.

Author(s)

Sarah Goslee

See Also

scale

Examples

x <- matrix(1:15, ncol = 3)

# uses min and max of the data
relrange(x)

# uses min and max determined by other knowledge of the variables
relrange(x, globalmin = c(0, 0, 0), globalmax = c(6, 10, 20))
Description

Extracts residuals from an S3 object of class `mgram` (only relevant for objects created by `pmgram()`).

Usage

```r
## S3 method for class 'mgram'
residuals(object, ...)
```

Arguments

- `object` an object of class `mgram`
- `...` additional arguments

Value

vector of residuals.

Author(s)

Sarah Goslee

See Also

`pmgram`, `mgram`

Examples

```r
#### Partial pmgram example

# generate a simple surface
# with complex nonlinear spatial pattern
x <- matrix(1:10, nrow=10, ncol=10, byrow=FALSE)
y <- matrix(1:10, nrow=10, ncol=10, byrow=TRUE)

# create z1 and z2 as functions of x, y
# and scale them to [0, 1]
z1 <- x + 3*y
z2 <- y - cos(x)

z1 <- (z1 - min(z1)) / (max(z1) - min(z1))
z2 <- (z2 - min(z2)) / (max(z2) - min(z2))
z12 <- (z1 + z2*2)/3
```
# analyze the pattern of z across space
z1 <- as.vector(z1)
z2 <- as.vector(z2)
z12 <- as.vector(z12)
z1.d <- dist(z1)
z2.d <- dist(z2)
z12.d <- dist(z12)

space <- cbind(as.vector(x), as.vector(y))
space.d <- dist(space)

# take partial correlogram of z12 given z1
z.z1 <- pmgram(z12.d, space.d, z2.d, nperm=0, resids=TRUE)
summary(residuals(z.z1))

---

### rotate2d

#### Rotate a 2D ordination.

**Description**

Rotates a two-dimensional ordination configuration to place the direction indicated along the horizontal axis.

**Usage**

`rotate2d(ord, x)`

**Arguments**

- `ord` A matrix or data frame with two columns, or a vf object, containing the points of an ordination configuration.
- `x` The coordinates of a point in the ordination space.

**Details**

The configuration ord is rotated so that the vector defined by c(0, 0), and x is along the horizontal axis. This can be useful for placing a specific variable, for instance from vf(), in a consistent direction across multiple ordinations. Doing so can facilitate interpretation.

**Value**

A rotated data frame of coordinates of the same size as ord and in the same order. If ord was produced by vf(), the complete vf object is returned.

**Author(s)**

Sarah Goslee
See Also
  vf, nmds

Examples

```r
# Example of multivariate analysis using built-in iris dataset
data(iris)
iris.d <- dist(iris[,1:4])

### nmds() is timeconsuming, so this was generated
### in advance and saved.
### set.seed(1234)
### iris.nmds <- nmds(iris.d, nits=20, mindim=1, maxdim=4)
### save(iris.nmds, file="ecodist/data/iris.nmds.rda")
data(iris.nmds)

# examine fit by number of dimensions
plot(iris.nmds)

# choose the best two-dimensional solution to work with
iris.nmin <- min(iris.nmds, dims=2)

# fit the data to the ordination as vectors
### vf() is timeconsuming, so this was generated
### in advance and saved.
### set.seed(1234)
### iris.vf <- vf(iris.nmin, iris[,1:4], nperm=1000)
### save(iris.vf, file="ecodist/data/iris.vf.rda")
data(iris.vf)

plot(iris.nmin, col=as.numeric(iris$Species), pch=as.numeric(iris$Species), main="NMDS")
plot(iris.vf)

# rotate configuration so Sepal Width is along the horizontal axis
iris.nmin.rot <- rotate2d(iris.nmin, iris.vf[2, 1:2])
iris.vf.rot <- rotate2d(iris.vf, iris.vf[2, 1:2])

plot(iris.nmin.rot, col=as.numeric(iris$Species), pch=as.numeric(iris$Species), main="NMDS")
plot(iris.vf.rot)
```

---

**vf**  
Vector fitting

**Description**

Fits ancillary variables to an ordination configuration.
Usage

vf(ord, vars, nperm = 100)

Arguments

ord          matrix containing a 2-dimensional ordination result with axes as columns.
vars         matrix with ancillary variables as columns.
nperm        number of permutation for the significance test. If nperm = 0, the test will be omitted.

Details

Vector fitting finds the maximum correlation of the individual variables with a configuration of samples in ordination space.

Value

an object of class vf, which is a data frame with the first 2 columns containing the scores for every variable in each of the 2 dimensions of the ordination space. r is the maximum correlation of the variable with the ordination space, and pval is the result of the permutation test.

Author(s)

Sarah Goslee

References


See Also

plot.vf

Examples

# Example of multivariate analysis using built-in iris dataset
data(iris)
iris.d <- dist(iris[,1:4])

### nmds() is timeconsuming, so this was generated
### in advance and saved.
### set.seed(1234)
### iris.nmds <- nmds(iris.d, nits=20, mindim=1, maxdim=4)
### save(iris.nmds, file="ecodist/data/iris.nmds.rda")
data(iris.nmds)

# examine fit by number of dimensions
plot(iris.nmds)
# choose the best two-dimensional solution to work with
iris.nmin <- min(iris.nmds, dims=2)

# fit the data to the ordination as vectors
### vf() is timeconsuming, so this was generated
### in advance and saved.
### set.seed(1234)
### iris.vf <- vf(iris.nmin, iris[,1:4], nperm=1000)
### save(iris.vf, file="ecodist/data/iris.vf.rda")
data(iris.vf)

plot(iris.nmin, col=as.numeric(iris$Species), pch=as.numeric(iris$Species), main="NMDS")
plot(iris.vf)

# rotate configuration so Sepal Width is along the horizontal axis
iris.nmin.rot <- rotate2d(iris.nmin, iris.vf[2, 1:2])
iris.vf.rot <- rotate2d(iris.vf, iris.vf[2, 1:2])

plot(iris.nmin.rot, col=as.numeric(iris$Species), pch=as.numeric(iris$Species), main="NMDS")
plot(iris.vf.rot)

---

**xdistance**

Cross-distance between two datasets.

**Description**

Pairwise dissimilarity calculation between rows of one dataset and rows of another, for instance across different sampling periods for the same set of sites.

**Usage**

`xdistance(x, y, method = "euclidean")`

**Arguments**

- **x**: A site by species or other matrix or data frame.
- **y**: A a second site by species dataset, which must have at least the same columns.
- **method**: This function calls `distance` to do the calculations, and will accept any symmetric method used there, currently: "euclidean", "bray-curtis", "manhattan", "mahalanobis" (squared Mahalanobis distance), "jaccard", "sorensen", "gower", "modgower10" (modified Gower, base 10), "modgower2" (modified Gower, base 2). Partial matching will work for selecting a method. The asymmetric "difference" method will not work for calculating cross-distances.
Details

This function will calculate rowwise dissimilarities between any pair of matrices or data frames with the same number of columns. Note that the cross-dissimilarity functions are for research purposes, and are not well-tested.

Value

A non-symmetric and possibly not square matrix of dissimilarities of class xdist, where \( \text{result} <- \text{xdistance}(x, y) \) produces a matrix with \( \text{result}[a, b] \) containing the dissimilarity between \( x[a, \] and \( y[b, \).

Author(s)

Sarah Goslee

See Also

distance, xmantel, xmgram

Examples

data(graze)

```r
### EXAMPLE 1: Square matrices

# take two subsets of sites with different dominant grass abundances
# use cut-offs that produce equal numbers of sites
dom1 <- subset(graze, POPR > 50 & DAGL < 20) # 8 sites
dom2 <- subset(graze, POPR < 50 & DAGL > 20) # 8 sites

# first two columns are site info
dom.xd <- xdistance(dom1[, -c(1,2)], dom2[, -c(1,2)], "bray")

# environmental and spatial distances; preserve rownames
forest.xd <- xdistance(dom1[, "forestpct", drop=FALSE],
                      dom2[, "forestpct", drop=FALSE])
sitelocation.xd <- xdistance(dom1[, "sitelocation", drop=FALSE],
                             dom2[, "sitelocation", drop=FALSE])

# permutes rows and columns of full nonsymmetric matrix
xmantel(dom.xd ~ forest.xd)
xmantel(dom.xd ~ forest.xd + sitelocation.xd)

plot(xmgram(dom.xd, sitelocation.xd))
```

```r
### EXAMPLE 2: Non-square matrices

# take two subsets of sites with different dominant grass abundances
# this produces a non-square matrix
```
xmantel <- subset(graze, POPR > 45 & DAGL < 20) # 13 sites
dom2 <- subset(graze, POPR < 45 & DAGL > 20) # 8 sites

# first two columns are site info
dom.xd <- xdistance(dom1[, -c(1,2)], dom2[, -c(1,2)], "bray")

# environmental and spatial distances; preserve rownames
forest.xd <- xdistance(dom1[, "forestpct", drop=FALSE],
                      dom2[, "forestpct", drop=FALSE])
sitelocation.xd <- xdistance(dom1[, "sitelocation", drop=FALSE],
                          dom2[, "sitelocation", drop=FALSE])

# permutes rows and columns of full nonsymmetric matrix
xmantel(dom.xd ~ forest.xd, dims=c(13, 8))
xmantel(dom.xd ~ forest.xd + sitelocation.xd, dims=c(13, 8))
plot(xmgram(dom.xd, sitelocation.xd))

---

**xmantel**

*Cross-Mantel test*

**Description**

Simple and partial cross-Mantel tests, with options for ranked data and permutation tests.

**Usage**

```
xmantel(formula = formula(data), data, dims = NA,
        nperm = 1000, mrank = FALSE)
```

**Arguments**

- **formula** formula describing the test to be conducted. For this test, `y ~ x` will perform a simple Mantel test, while `y ~ x + z1 + z2 + z3` will do a partial Mantel test of the relationship between `x` and `y` given `z1, z2, z3`. All variables should be either non-symmetric square cross-dissimilarity matrices of class `xdist`, or vector forms thereof.

- **data** an optional dataframe containing the variables in the model as columns of dissimilarities. By default the variables are taken from the current environment.

- **dims** if the dissimilarity matrices are not square, the dimensions must be provided as `c(nrow, ncol)`

- **nperm** number of permutations to use. If set to 0, the permutation test will be omitted.

- **mrank** if this is set to FALSE (the default option), Pearson correlations will be used. If set to TRUE, the Spearman correlation (correlation ranked distances) will be used.
Details

If only one independent variable is given, the simple Mantel r (r_{12}) is calculated. If more than one independent variable is given, the partial Mantel r (r_{y|x1 ...}) is calculated by permuting one of the original dissimilarity matrices. Note that the cross-dissimilarity functions are for research purposes, and are not well-tested.

Value

- **mantelr**: Mantel coefficient.
- **pval1**: one-tailed p-value (null hypothesis: \( r \leq 0 \)).
- **pval2**: one-tailed p-value (null hypothesis: \( r \geq 0 \)).
- **pval3**: two-tailed p-value (null hypothesis: \( r = 0 \)).

Author(s)

Sarah Goslee

See Also

- xdistance, xmgram

Examples

data(graze)

```r
### EXAMPLE 1: Square matrices

# take two subsets of sites with different dominant grass abundances
# use cut-offs that produce equal numbers of sites
dom1 <- subset(graze, POPR > 50 & DAGL < 20) # 8 sites
dom2 <- subset(graze, POPR < 50 & DAGL > 20) # 8 sites

# first two columns are site info
dom.xd <- xdistance(dom1[, -c(1,2)], dom2[, -c(1,2)], "bray")

# environmental and spatial distances; preserve rownames
forest.xd <- xdistance(dom1[, "forestpct", drop=FALSE],
dom2[, "forestpct", drop=FALSE])
sitelocation.xd <- xdistance(dom1[, "sitelocation", drop=FALSE],
dom2[, "sitelocation", drop=FALSE])

# permutes rows and columns of full nonsymmetric matrix
xmantel(dom.xd ~ forest.xd)
xmantel(dom.xd ~ forest.xd + sitelocation.xd)
plot(xmgram(dom.xd, sitelocation.xd))
```

### EXAMPLE 2: Non-square matrices
# take two subsets of sites with different dominant grass abundances
# this produces a non-square matrix

```r
dom1 <- subset(graze, POPR > 45 & DAGL < 20) # 13 sites
dom2 <- subset(graze, POPR < 45 & DAGL > 20) # 8 sites
```

# first two columns are site info
```
dom.xd <- xdistance(dom1[, -c(1,2)], dom2[, -c(1,2)], "bray")
```

# environmental and spatial distances; preserve rownames
```
forest.xd <- xdistance(dom1[, "forestpct", drop=FALSE],
                      dom2[, "forestpct", drop=FALSE])
sitelocation.xd <- xdistance(dom1[, "sitelocation", drop=FALSE],
                            dom2[, "sitelocation", drop=FALSE])
```

# permutes rows and columns of full nonsymmetric matrix
```
xmantel(dom.xd ~ forest.xd, dims=c(13, 8))
xmantel(dom.xd ~ forest.xd + sitelocation.xd, dims=c(13, 8))
```

plot(xmgram(dom.xd, sitelocation.xd))

---

**xmgram**

*Cross-Mantel correlogram*

**Description**

Calculates simple Mantel correlograms from cross-distance matrices.

**Usage**

```
xmgram(species.xd, space.xd, breaks, nclass, stepsize, equiprobable = FALSE, nperm = 1000,
        mrank = FALSE, alternative = "two.sided", trace = FALSE)
```

**Arguments**

- `species.xd`: non-symmetric square cross-distance matrix.
- `space.xd`: non-symmetric square matrix of geographic distances.
- `breaks`: locations of class breaks. If specified, overrides nclass and stepsize.
- `nclass`: number of distance classes. If not specified, Sturges's rule will be used to determine an appropriate number of classes.
- `stepsize`: width of each distance class. If not specified, nclass and the range of space.d will be used to calculate an appropriate default.
- `equiprobable`: if TRUE, create nclass classes of equal number of distances; if FALSE, create nclass classes of equal width
- `nperm`: number of permutations to use. If set to 0, the permutation test will be omitted.
mranks if this is set to FALSE (the default option), Pearson correlations will be used. If set to TRUE, the Spearman correlation (correlation ranked distances) will be used.

alternative default is "two.sided", and returns p-values for H0: rM = 0. The alternative is "one.sided", which returns p-values for H0: rM <= 0.

trace if TRUE, returns progress indicators.

Details

This function calculates cross-Mantel correlograms. The Mantel correlogram is essentially a multivariate autocorrelation function. The Mantel r represents the dissimilarity in variable composition (often species composition) at a particular lag distance. Note that the cross-dissimilarity functions are for research purposes, and are not well-tested.

Value

Returns an object of class mgram, which is a list with two elements. mgram is a matrix with one row for each distance class and 6 columns:

- lag midpoint of the distance class.
- ngroup number of distances in that class.
- mantelr Mantel r value.
- pval p-value for the test chosen.

resids is NA for objects calculated by mgram().

Author(s)

Sarah Goslee

References


See Also

xdistance xmantel, plot.mgram

Examples

# Need to develop a cross-dissimilarity example
data(graze)

### EXAMPLE 1: Square matrices

# take two subsets of sites with different dominant grass abundances
# use cut-offs that produce equal numbers of sites
dom1 <- subset(graze, POPR > 50 & DAGL < 20) # 8 sites
dom2 <- subset(graze, POPR < 50 & DAGL > 20) # 8 sites
# first two columns are site info
dom.xd <- xdistance(dom1[, -c(1,2)], dom2[, -c(1,2)], "bray")

# environmental and spatial distances; preserve rownames
forest.xd <- xdistance(dom1[, "forestpct", drop=FALSE],
                      dom2[, "forestpct", drop=FALSE])
sitelocation.xd <- xdistance(dom1[, "sitelocation", drop=FALSE],
                           dom2[, "sitelocation", drop=FALSE])

# permutes rows and columns of full nonsymmetric matrix
xmantel(dom.xd ~ forest.xd)

### EXAMPLE 2: Non-square matrices

# take two subsets of sites with different dominant grass abundances
# this produces a non-square matrix

dom1 <- subset(graze, POPR > 45 & DAGL < 20) # 13 sites
dom2 <- subset(graze, POPR < 45 & DAGL > 20) # 8 sites

# first two columns are site info
dom.xd <- xdistance(dom1[, -c(1,2)], dom2[, -c(1,2)], "bray")

# environmental and spatial distances; preserve rownames
forest.xd <- xdistance(dom1[, "forestpct", drop=FALSE],
                      dom2[, "forestpct", drop=FALSE])
sitelocation.xd <- xdistance(dom1[, "sitelocation", drop=FALSE],
                           dom2[, "sitelocation", drop=FALSE])

# permutes rows and columns of full nonsymmetric matrix
xmantel(dom.xd ~ forest.xd, dims=c(13, 8))

plot(xmgram(dom.xd, sitelocation.xd))

---

**z.no**

**Example for pmgram**

**Description**

An object of class mgram for use in the example for `pmgram`. Many of the functions in ecodist take a long time to run, so prepared examples have been included.

**Usage**

```r
data(z.no)
```
Format

See `pmgram` for current format specification.

Author(s)

Sarah Goslee

See Also

`pmgram`, `z.z1`

Examples

```r
#### Partial pmgram example

# generate a simple surface
# with complex nonlinear spatial pattern

x <- matrix(1:25, nrow=25, ncol=25, byrow=FALSE)
y <- matrix(1:25, nrow=25, ncol=25, byrow=TRUE)

# create z1 and z2 as functions of x, y
# and scale them to [0, 1]

z1 <- x + 3*y
z2 <- y - cos(x)

z1 <- (z1 - min(z1)) / (max(z1) - min(z1))
z2 <- (z2 - min(z2)) / (max(z2) - min(z2))

z12 <- (z1 + z2*2)/3

# look at patterns

layout(matrix(c(1, 1, 2, 2,
                1, 1, 2, 2,
                3, 3, 4, 4,
                3, 3, 5, 5), nrow=4, byrow=TRUE))

image(z1, col=gray(seq(0, 1, length=20)), zlim=c(0,1))
image(z2, col=gray(seq(0, 1, length=20)), zlim=c(0,1))
image(z12, col=gray(seq(0, 1, length=20)), zlim=c(0,1))

# analyze the pattern of z across space

z1 <- as.vector(z1)
z2 <- as.vector(z2)
z12 <- as.vector(z12)
z1.d <- dist(z1)
z2.d <- dist(z2)
z12.d <- dist(z12)

space <- cbind(as.vector(x), as.vector(y))
```
space.d <- dist(space)

# take partial correlogram without effects of z1
### pgram() is time-consuming, so this was generated
### in advance and saved.
### set.seed(1234)
### z.no <- pmgram(z12.d, space.d, nperm=1000, resids=FALSE)
### save(z.no, file="ecodist/data/z.no.rda")
plot(z.no)

# take partial correlogram of z12 given z1
### pgram() is time-consuming, so this was generated
### in advance and saved.
### set.seed(1234)
### z.z1 <- pmgram(z12.d, space.d, z2.d, nperm=1000, resids=FALSE)
### save(z.z1, file="ecodist/data/z.z1.rda")
plot(z.z1)

---

z.z1

Example for pmgram

Description

An object of class mgram for use in the example for pmgram. Many of the functions in ecodist take a long time to run, so prepared examples have been included.

Usage

data(z.z1)

Format

See pmgram for current format specification.

Author(s)

Sarah Goslee

See Also

pmgram, z.no

Examples

### Partial pmgram example

# generate a simple surface
# with complex nonlinear spatial pattern
x <- matrix(1:25, nrow=25, ncol=25, byrow=FALSE)
y <- matrix(1:25, nrow=25, ncol=25, byrow=TRUE)

# create z1 and z2 as functions of x, y
# and scale them to [0, 1]
z1 <- x + 3*y
z2 <- y - cos(x)

z1 <- (z1 - min(z1)) / (max(z1) - min(z1))
z2 <- (z2 - min(z2)) / (max(z2) - min(z2))
z12 <- (z1 + z2*2)/3

# look at patterns

layout(matrix(c(1, 1, 1, 1, 2, 2, 2, 2, 3, 3, 3, 3, 4, 4, 5, 5), nrow=4, byrow=TRUE))

image(z1, col=gray(seq(0, 1, length=20)), zlim=c(0,1))
image(z2, col=gray(seq(0, 1, length=20)), zlim=c(0,1))
image(z12, col=gray(seq(0, 1, length=20)), zlim=c(0,1))

# analyze the pattern of z across space
z1 <- as.vector(z1)
z2 <- as.vector(z2)
z12 <- as.vector(z12)
z1.d <- dist(z1)
z2.d <- dist(z2)
z12.d <- dist(z12)

space <- cbind(as.vector(x), as.vector(y))
space.d <- dist(space)

# take partial correlogram without effects of z1
### pgram() is time-consuming, so this was generated
### in advance and saved.
### set.seed(1234)
### z.no <- pmgram(z12.d, space.d, nperm=1000, resids=FALSE)
### save(z.no, file="ecodist/data/z.no.rda")
plot(z.no)

# take partial correlogram of z12 given z1
### pgram() is time-consuming, so this was generated
### in advance and saved.
### set.seed(1234)
### z.z1 <- pmgram(z12.d, space.d, z2.d, nperm=1000, resids=FALSE)
### save(z.z1, file="ecodist/data/z.z1.rda")
plot(z.z1)
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