Package ‘EcoGenetics’

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Type Package

Title Management and Exploratory Analysis of Spatial Data in Landscape Genetics

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Description Management and exploratory analysis of spatial data in landscape genetics. Easy integration of information from multiple sources with ‘`ecogen’’ objects.

License GPL (>= 2)

URL https://github.com/cran/EcoGenetics,
https://leandroroser.github.io/EcoGenetics-Tutorial

LazyLoad yes

Depends R (>= 3.0), methods

Imports d3heatmap, edgebundleR, ggplot2, grid, htmlwidgets, igraph, jsonlite, magrittr, networkD3, party, pheatmap, plotly, raster, reshape2, rgdal, rkt, SoDA, sp, parallel, doParallel, foreach

Suggests adegenet, testthat, covr, vegan, hierfstat

‘ecogen.3OF6.basic.methods.R’ ‘ecogen.4OF6.brackets.R’
‘ecopop.3OF6.basic.methods.R’ ‘ecopop.4OF6.brackets.R’
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Description

Management and exploratory analysis of spatial data in landscape genetics. Easy integration of information from multiple sources with "ecogen" objects.
I. STRUCTURE OF THE PACKAGE

The spatial analysis module of the package computes global (Moran’s I, Mantel test, etc.) and local (Getis-Ord’s G*, local Moran’s I, etc.) spatial tests, variograms and correlograms (see also this link). These analyses use other tool provided by the package: spatial weights matrices or a sequence of spatial weights matrices.

The package have also special plot methods for each of the analyses. Several conversor of data from/to other programs are available, (as to genepop - an importer tool is also defined for genepop-, SPAGeDi, etc.). Basic manipulation of genetic matrices is allowed by eco.convert and eco.format.

Tools for computation of NDVI in Landsat imagery, post-process of rasters and temporal analysis can be found in eco.NDVI, eco.NDVI.post and eco.theilsen. Several other useful functions are defined in the package.

The results obtained with the main functions defined in EcoGenetics are object of class S4. For these objects, the package defines a "show" method for a general overview of the results, and methods to extract the information (generic accessors and double square brackets ("[]") definitions).

For storing and pre-processing the data, the package defines two special classes: ecogen and ecopop

II. ECOCEN OBJECTS

Landscape genetics research requires the integration of data originated in different sources. The class ecogen has been designed for handling multidimensional data of individuals. Its basic structure is the following:

- An **XY** slot, storing a data frame with geographic coordinates.
- A **P** slot, storing a phenotypic data frame.
- A **G** slot, storing a genotypic data frame.
- An **A** slot containing as allelic frequencies the information of G (only available for codominant markers.)
- An **E** slot, storing an environmental data frame.
- A **S** slot, storing a data frame with classes assigned to the individuals.
- A **C** slot, for a custom data frame.
- An **OUT** slot, containing a list for the storage of the results.

For dominant (presence-absence) markers, the slot A is empty.

The construction of a new "ecogen" object from a data frame is made with the homonymous function.

library("EcoGenetics")
data(eco.test)

eco <- ecogen(XY = coordinates, P = phenotype, G = genotype, E = environment, S = structure, order.G = TRUE)

The package defines several methods for manipulation of ecogen objects: eco.cbind, to bind ecogen objects by column, eco.split, to split ecogen objects by group, eco.rbind, to bind ecogen objects by row and to re-bind previously splitted objects, eco.subset, to select a group of individuals, given a population, eco.merge, to merge ecogen objects with different composition of individuals. The functions nrow, ncol, dim, names, as.list, is.ecogen and show are also defined. Single brackets ("[") and double brackets ("[[") are also defined to select a subset of individuals or to get/set the data of a particular slot, respectively. Conversors from and to genind, from and to spagedi, from and to genepop, to hierfstat and to geneland are included. See the ecogen constructor documentation and the publication of the package, cited at the end of this document, for more information.

III. ECOPOP OBJECTS

The class ecopop is the population analogue of ecogen class: while ecogen rows represent individuals, ecopop rows represent populations. An ecopop object contains aggregated data for each population (mean or other statistics of quantitative variables, and counts for each level of qualitative variables [including allele counts]). In normal situations, an ecopop object is obtained from an ecogen object as follows:

ecopop_object <- ecogen2ecopop(eco, hier = "pop") # where pop is the name of the # column of the slot S with # the population of each # individual

ecopop objects can also be obtained from the ecopop constructor, as detailed in the documentation of the function.

The functions nrow, ncol, dim, names, as.list, is.ecogen and show are defined for ecopop objects. Single brackets ("[") and double brackets ("[[") are also defined to select a subset of individuals or to get/set the data of a particular slot, respectively. See ecopop constructor documentation for more information. Conversion from and to genpop is defined.

Author(s)

Leandro Roser, Juan Vilardi, Beatriz Saidman and Laura Ferreyra

Maintainer: Leandro Roser <learoser@gmail.com>

References


Generate aggregated dataframe

Generate aggregated dataframe
aue.check_class

Usage

aue.aggregated_df(X, hier, fun, factor_to_counts = FALSE, ...)

Arguments

x data frame
hier hierarchy
fun function
factor_to_counts split factor into counts for each level?
... additional parameters passed to fun

Author(s)

Leandro Roser <learoser@gmail.com>

aue.check_class Obtain the classes for each column of a data frame

Description

Obtain the classes for each column of a data frame

Usage

aue.check_class(X)

Arguments

x factor

Author(s)

Leandro Roser <learoser@gmail.com>
aue.dataAngle

**Angles for an XY coordinates matrix**

**Description**

Angles for an XY coordinates matrix

**Usage**

```r
aue.dataAngle(XY, maxpi = FALSE, deg = FALSE, latlon = FALSE)
```

**Arguments**

- **XY**: XY matrix with projected coordinates
- **maxpi**: angles bounded between 0 - pi?
- **deg**: angles in decimal degrees?
- **latlon**: Are the coordinates in decimal degrees format? Default FALSE. If TRUE, the coordinates must be in a matrix/data frame with the longitude in the first column and latitude in the second. The position is projected onto a plane in meters with the function `geoXY`.

**Author(s)**

Leandro Roser <learoser@gmail.com>

---

aue.df2image

**Transforming a data frame into a raster**

**Description**

This is the inverse function of aue.image2df

**Usage**

```r
aue.df2image(x, origin = c("upperleft", "lowerleft"))
```

**Arguments**

- **x**: output of aue.image2df
- **origin**: Origin of the reference for the coordinates. Default: upperleft.

**Author(s)**

Leandro Roser <learoser@gmail.com>
**aue.dummy2af**  
*Conversion from dummy allele matrix to frequencies*

**Description**

Conversion from dummy allele matrix to frequencies

**Usage**

```r
aue.dummy2af(df, loc_fac)
```

**Arguments**

- `df`: data frame with alleles coded in dummy format
- `loc_fac`: factor with the locus of each allele

---

**aue.image2df**  
*Transforming a raster into a data frame with cartesian coordinates*

**Description**

This function returns a data frame with the column number (x), row number (y) and cell value (z) of each pixel in a raster.

**Usage**

```r
aue.image2df(mat, origin = c("upperleft", "lowerleft"),
              out = c("data.frame", "matrix"))
```

**Arguments**

- `mat`: Input raster matrix.
- `out`: output format: "data.frame" (default) or "matrix".

**Author(s)**

Leandro Roser <learoser@gmail.com>
Examples

```r
## Not run:
ras <- matrix(eco["P"],[,1],15,15)
image(ras)
ras.row <- aue.image2df(ras)
ras.row
image(matrix(ras.row[,3], 15, 15))

## End(Not run)
```

---

**aue.phenosimil**  
*Phenotypic similarity for vector, matrix or data frame according to Ritland (1996)*

**Description**

Phenotypic similarity for vector, matrix or data frame according to Ritland (1996)

**Usage**

```r
aue.phenosimil(X)
```

**Arguments**

- `X`  
  Data frame, matrix or vector. If `X` is not a vector, the program returns a list of matrices.

**Author(s)**

Leandro Roser <learoser@gmail.com>

---

**aue.rescale**  
*Scaling a data frame or matrix to [0, 1] or [-1, 1] ranges*

**Description**

This program scales each column of a data frame or a matrix to [0,1] range, computing 

\[(X)_{ij} - (X_{\text{min}})_{i}\) / \text{range}((X)_i) \]

for each individual \(j\) of the variable \(i\) or to [-1,1] range computing 

\[2 * ((X)_{ij} - (X_{\text{min}})_{i}) / \text{range}(\text{X}_i) - 1\]

**Usage**

```r
aue.rescale(dfm, method = c("zero.one", "one.one"))
```
### Arguments

- **dfm**: Data frame, matrix or vector to scale.
- **method**: Scaling method: "zero.one" for scaling to [0, 1], "one-one" for scaling to [-1,1].

### Author(s)

Leandro Roser <learoser@gmail.com>

### Examples

```r
# Not run:
data(eco.test)
require(adegenet)
pc <- dudi.pca(eco[,"P"], scannf = FALSE, nf = 3)
pc <- pc$li
pc <- aue.rescale(pc)
plot(eco[,"XY"][-1,], eco[,"XY"][-2,], col = rgb(pc), pch = 16,
     cex = 1.5, xlab = "X", ylab = "Y")
```

### Description

This program takes a matrix and orders the content of each cell. It was specially designed for genetic data, but can be used with any data that can be rearranged by the function `order`. The arguments ploidy and ncode determine the mode of ordering the data. The cells corresponding to each individual `i` and loci `j` are ordered in ascending order in default option (it can be passed decreasing = TRUE as argument, if descending order is desired). For example, a locus with ploidy = 2 and ncod =1, coded as 51, for an individual, will be recoded as 15. A locus with ploidy = 3 and coded as 143645453, will be recoded as 143453645 (alleles 143, 454 and 645).

### Usage

```r
aue.sort(X, ncod = NULL, ploidy = 2, sep.loc = "", chk.plocod = TRUE, ...
```
Arguments

- **X** Any matrix with content to order.
- **ncod** Number of digits coding each allele (e.g., 1: x, 2: xx, 3: xxx, etc.). If NULL, ncode will be obtained from the ploidy and the maximum number of characters in the data cells.
- **ploidy** Ploidy of the data.
- **sep.loc** Character string separating alleles.
- **chk.plocod** Default TRUE. The function checks coherence in ploidy and number of digits coding alleles.
- **...** Additional arguments passed to `order`.

Author(s)

Leandro Roser < learoser@gmail.com >

Examples

```r
## Not run:

# Example 1-----------------------------

geno <- c(12, 52, 62, 45, 54, 21)
geno <- matrix(geno, 3, 2)

# ordering the data
aue.sort(geno, ploidy = 2)

# decreasing sort order
aue.sort(geno, ploidy = 2, decreasing = TRUE)

# Example 2-----------------------------

geno2 <- c(123456, 524556, 629359, 459459, 543950, 219405)
geno2 <- matrix(geno2, 3, 2)

# ordering the data as diploid
aue.sort(geno2, ploidy = 2) # the data is ordered using blocks of 3 characters

# ordering the data as triploid
aue.sort(geno2, ploidy = 3) # the data is ordered using blocks of 2 characters

# error: the ploidy and the number of characters are not congruent
aue.sort(geno2, ploidy = 5)

# error: the ploidy and the number of characters are not congruent
aue.sort(geno2, ploidy = 5)
```
aue.split_categorical

Description

Split categorical variable into levels, using a second factor (hierarchy) to aggregate the data

Usage

aue.split_categorical(x, hier)

Arguments

x
factor
hier
hierarchy

Author(s)

Leandro Roser <learoser@gmail.com>

data(eco.test)

cartesian coordinates

Description

Data frame with cartesian coordinates of 225 simulated individuals.

Usage

data(eco.test)
coordinates

Author(s)

Leandro Roser <learoser@gmail.com>
Description

data frame with simulated environmental data of 173 individuals.

Usage

data(eco4)
E

Author(s)
Leandro Roser<learoser@gmail.com>

description
ecogen object with simulated data of 225 individuals, with codominant markers

Usage

data(eco.test)
eco

Author(s)
Leandro Roser<learoser@gmail.com>

description
Allelic frequency histograms for an ecogen genetic data frame

Usage

eco.alfreq(eco, grp = NULL)
Chi-square and Fisher's exact test for association of loci and alleles with a factor

Description
Chi-square and Fisher's exact test for association of loci and alleles with a factor

Usage
eco.association(eco, assoc = c("within", "between"), x, method = c("fisher.test", "chisq.test"), nrep = 99, adjust = "none", ndig = NA)

Arguments

eco Object of class "ecogen".
assoc "between" if the association test should be performed between a factor and a loci, or "within" if the association test should be performed between a factor and alleles within loci. For haploid data, use option "within".
x The name of the S slot column with the groups for the association test.
eco.bearing

**Description**

Construction of angular spatial weights

**Usage**

```r
eco.bearing(con, theta, XY = NULL, latlon = FALSE)
```
Arguments

- **con**: an eco.weight or eco.lagweight object
- **theta**: reference angle in degrees, between 0 and 180, counterclockwise, with 0 representing the positive x axis, 90 representing the positive y axis, 180 representing the negative x axis. Note that angles 0 and 180 yield identical results.
- **XY**: Matrix/data frame with projected coordinates. Default NULL.
- **latlon**: Are the coordinates in decimal degrees format? Default FALSE. If TRUE, the coordinates must be in the XY matrix/data frame with longitude in the first column and latitude in the second. The position is projected onto a plane in meters with the function `geoXY`.

Details

This program computes an angular weights object (AW) (or a list of AW). If a weights object is passed as argument ("con") the program computes an AW with this element. If XY is passed, the program first computes a matrix of N x N, where N is the number of rows in XY, and then uses the matrix as input to compute the AW. Each element in the weights matrices is then weighted by the squared cosine of the angle formed with the x positive axis by a line connecting the pair of points. Note that this method assumes that the distances in the eco.weight or eco.lagweight object are projected as great-circle distances (for example, using latlon = TRUE during weights construction or UTM coordinates for elements passed with "con", or latlon set TRUE in this function for a coordinates element passed with XY).

Note also that when angular weights are constructed for XY coordinates, the output consists of a weights object with values bounded between 0 and 1, being 1 if the if the direction pointed by the vector V connecting the elements i, j in the matrix points in the same direction of the reference vector R (with and angle theta with the positive x axis), and 0 if V is perpendicular to R.

Value

An object of class eco.weight with a bearing weights matrix

**ACCESS TO THE SLOTS** The content of the slots can be accessed with the corresponding accessors, using the generic notation of EcoGenetics (<ecoslot.> + <name of the slot> + <name of the object>). See help("EcoGenetics accessors") and the Examples for the function eco.weight and eco.lagweight.

Author(s)

Leandro Roser <learoser@gmail.com>

References

eco.cbind

Combining ecogen objects by column

Description
Combining ecogen objects by column

Usage
eco.cbind(eco1, eco2, ..., missing = c("0", "MEAN", "NA"))

Arguments
eco1 Object of class "ecogen".
eco2 Object of class "ecogen".
... Other "ecogen" objects to combine and the specification of the data frames to combine. Can be any of the following(s): "P","G", "E", "S", "C", or "ALL" (default). If a "G" data frame is provided, the program also generates the A slot coding the missing data as "0" in default option (see the argument "missing"). The XY slot is generated automatically if present.
missing Missing data manipulation. It can take three values ("0", "NA" or "MEAN"- i.e, the mean frequency of the corresponding allele). Missing elements are coded as 0 in the default option.

Author(s)
Leandro Roser <learoser@gmail.com>
Examples

## Not run:

data(eco.test)
eco.example <- eco.cbind(eco,eco,"ALL")
eco.example
eco.example2 <- eco.cbind(eco, eco,"P", "G", missing="NA")
eco.example2

## End(Not run)

---

eco.clear

Clearing the working environment, maintaining only the specified objects

Description

This function removes all the elements of the working environment, with the exception of those included in the argument of the function. Hidden elements can also be removed by setting all = TRUE.

Usage

eco.clear(..., all = FALSE)

Arguments

... Objects to retain.
all Remove also hidden elements? Default FALSE.

Author(s)

Leandro Roser <learoser@gmail.com>

Examples

## Not run:

data(eco.test)
ls()
eco.clear(eco)
ls()

## End(Not run)
Description

This function interconverts genetic data among matrix format (one locus or one allele per column) and list format (one locus or one allele per column).

Usage

```r
eco.convert(X, input = c("matrix", "alleles.matrix", "list", "alleles.list"), output = c("matrix", "alleles.matrix", "list", "alleles.list"), ncod = NULL, ploidy = 2, sep.in, sep.out, chk.names = TRUE, chk.plocod = TRUE)
```

Arguments

- `X`: Input data.
- `input`: Input data format.
- `output`: Output data format.
- `ncod`: Number of digits coding each allele.
- `ploidy`: Ploidy of the data.
- `sep.in`: Character separating alleles in the input data if present.
- `sep.out`: Character separating alleles in the output data. Default option do not separate alleles.
- `chk.names`: Default TRUE. The function makes checks of individuals and loci names during conversion.
- `chk.plocod`: Default TRUE. The function checks coherence in/between ploidy and number of digits coding alleles for loci data during conversion.

Author(s)

Leandro Roser <learoser@gmail.com>

Examples

```r
# Not run:
data(eco3)

# One allele per column
loc2al <- eco.convert(eco3["G"], "matrix", "alleles.matrix", ploidy = 2)
loc2al

# Inverse operation (collapse alleles into locus)
```
Description

This program computes a Mantel correlogram for the data M, or a partial Mantel correlogram for the data M conditioned on MC, with P-values or bootstrap confidence intervals.

Usage

eco.cormantel(M, XY, MC = NULL, int = NULL, smin = 0, smax = NULL, nclass = NULL, seqvec = NULL, size = NULL, bin = c("sturges", "FD"), nsim = 99, classM = c("dist", "simil"), method = c("pearson", "spearman", "kendall"), test = c("permutation", "bootstrap"), alternative = c("auto", "two.sided", "greater", "less"), adjust = "holm", sequential = TRUE, latlon = FALSE, angle = NULL, as.deg = TRUE, ...)

Arguments

M Distance or similarity matrix.
XY  Data frame or matrix with individual’s positions (projected coordinates).
MC  Distance or similarity matrix (optional).
int Distance interval in the units of XY.
smin Minimum class distance in the units of XY.
smax Maximum class distance in the units of XY.
class Number of classes.
seqvec Vector with breaks in the units of XY.
size Number of individuals per class.
bin Rule for constructing intervals when a partition parameter (int, nclass or size) is
  not given. Default is Sturges’s rule (Sturges, 1926). Other option is Freedman-
  Diaconis method (Freedman and Diaconis, 1981).
nsim Number of Monte-Carlo simulations.
classM Are M and MC distance or similarity matrices? Default option is classM = "dist"
  (distance). For similarity, classM = "simil". An incorrect option selected will
  generate an inverted plot.
method Correlation method used for the construction of the statistic ("pearson", "spear-
  man" or "kendall"). Kendall’s tau computation is slow.
test If test = "bootstrap", the program generates a bootstrap resampling and the associ-
  ated confidence intervals. If test = "permutation" (default) a permutation test
  is made and the P-values are computed.
alternative The alternative hypothesis. If "auto" is selected (default) the program determines
  the alternative hypothesis. Other options are: "two.sided", "greater" and "less".
adjust Correction method of P-values for multiple tests, passed to p.adjust. Defalut
  is "holm".
sequential Should be performed a Holm-Bonberroni (Legendre and Legendre, 2012) ad-
  justment of P-values? Defalult TRUE.
latlon Are the coordinates in decimal degrees format? Defalut FALSE. If TRUE, the
  coordinates must be in a matrix/data frame with the longitude in the first column
  and latitude in the second. The position is projected onto a plane in meters with
  the function geoXY.
angle for computation of bearing correlogram (angle between 0 and 180). Default
  NULL (omnidirectional).
as.deg in case of bearing correlograms for multiple angles, generate an output for each
  lag in function of the angle? Default TRUE.
... Additional arguments passed to cor.

Value

The program returns an object of class "eco.correlog" with the following slots:

> OUT analysis output
> IN input data of the analysis
> BEAKS breaks
eco.cormantel

> CARDINAL number of elements in each class
> NAMES variables names
> METHOD analysis method
> DISTMETHOD method used in the construction of breaks
> TEST test method used (bootstrap, permutation)
> NSIM number of simulations
> PADJUST P-values adjust method for permutation tests

ACCESS TO THE SLOTS The content of the slots can be accessed with the corresponding accessor, using the generic notation of EcoGenetics (<ecoslot.> + <name of the slot> + <name of the object>). See help("EcoGenetics accessors") and the Examples section below.

Author(s)
Leandro Roser <learoser@gmail.com>

References

Examples

```r
## Not run:

data(eco.test)
require(ggplot2)

# Omnidiirectional correlogram

corm <- eco.cormantel(M = dist(eco["P"]), size=1000, smax=7, XY = eco["XY"], nsim = 99)
eco.plotCorrelog(corm)
```
corm <- eco.cormantel(M = dist(eco["P"]), size=1000, smax=7, XY = eco["XY"], nsim = 99, test = "bootstrap")
eco.plotCorrelog(corm)

# A directional approach based in bearing correlograms

corm_b <- eco.cormantel(M = dist(eco["P"]), size=1000, smax=7, XY = eco["XY"],
nsim = 99, angle = seq(0, 170, 10))

# use eco.plotCorrelogB for this object
eco.plotCorrelogB(corm_b)

# plot for the first distance class,
# use a number between 1 and the number of classes to select the corresponding class
eco.plotCorrelogB(corm_b, interactivePlot = FALSE, var = 1)

# partial Mantel correlogram

corm <- eco.cormantel(M = dist(eco["P"]), MC = dist(eco["E"]),
size=1000, smax=7, XY = eco["XY"], nsim = 99)
eco.plotCorrelog(corm)

# standard correlogram plots support the use of ggplot2 syntax
require(ggplot2)
mantelplot <- eco.plotCorrelog(corm, interactivePlot = FALSE)
mantelplot <- mantelplot + theme_bw() + theme(legend.position="none")
mantelplot

# ACCESSORS USE EXAMPLE
#
# the slots are accessed with the generic format
# (ecoslot. + name of the slot + name of the object).
# See help("EcoGenetics accessors")

ecoslot.OUT(corm) # slot OUT
ecoslot.BREAKS(corm) # slot BREAKS

## End(Not run)
Description

This program computes Moran’s, Geary’s and bivariate Moran’s correlograms, for single or multipel variables, with P-values or bootstrap confidence intervals. Correlograms can be omnidirectional or directional, the latter based in the bearing method (Rosenberg, 2000). The program allows high flexibility for the construction of intervals. For detailed information about the range partition methods see eco.lagweight

Usage

eco.correlog(Z, XY, Y = NULL, int = NULL, smin = 0, smax = NULL, nclass = NULL, size = NULL, seqvec = NULL, method = c("I", "C", "CC"), nsim = 99, test = c("permutation", "bootstrap"), alpha = 0.05, alternative = c("auto", "two.sided", "greater", "less"), adjust = "holm", sequential = ifelse((as.deg), FALSE, TRUE), include.zero = TRUE, cumulative = FALSE, bin = c("sturges", "FD"), row.sd = FALSE, latlon = FALSE, angle = NULL, as.deg = TRUE)

Arguments

Z Vector, matrix or data frame with variable/s (in matrix or data frame formats, variables in columns).
XY Data frame or matrix with individual’s positions (projected coordinates).
Y Vector with the second variable for Mantel’s Ixy cross-correlograms. If Z has multiple variables, the program will compute the cross-correlograms for each with Y.
int Distance interval in the units of XY.
smin Minimum class distance in the units of XY.
smax Maximum class distance in the units of XY.
nclass Number of classes.
size Number of individuals per class.
seqvec Vector with breaks in the units of XY.
method Correlogram method. It can be I for Moran’s I, C for Geary’s C and CC for Bivariate Moran’s Ixy. If method = "CC", the program computes for the first interval (d = 0) the corresponding P-value and CI with cor.test.
nsim Number of Monte-Carlo simulations.
test If test = “bootstrap”, the program generates a bootstrap resampling and the associated confidence intervals of the null hypothesis. If test = “permutation” (default) a permutation test is made and the P-values are computed.
alpha Value for alpha (significance level). Default alpha = 0.05.
alternative The alternative hypothesis. If "auto" is selected (default) the program determines the alternative hypothesis. Other options are: "two.sided", "greater" and "less".
adjust P-values correction method for multiple tests. The selected method is passed as argument to p.adjust (default = "holm"). For bearing correlograms, the corrections (and permutation tests) are performed for individual correlograms of fixed variables (i.e., angles fixed [distances variable] or distances fixed [angles variable]).
Should a Holm-Bonferroni correction of P-values (Legendre and Legendre, 2012) be performed? Default TRUE (only available for omnidirectional correlograms or correlograms for fixed angles).

Should be included the distance = 0 in cross correlograms (i.e., the intra-individual correlation)? Default TRUE.

Should be constructed a cumulative correlogram?

Rule for constructing intervals when a partition parameter (int, nclass or size) is not given. Default is Sturges’ rule (Sturges, 1926). Other option is Freedman-Diaconis method (Freedman and Diaconis, 1981).

Logical. Should be row standardized the matrix? Default FALSE (binary weights).

Are the coordinates in decimal degrees format? Default FALSE. If TRUE, the coordinates must be in a matrix/data frame with the longitude in the first column and latitude in the second. The position is projected onto a plane in meters with the function geoxy.

for computation of bearing correlogram (angle between 0 and 180). Default NULL (omnidirectional).

in case of bearing correlograms for multiple angles, generate an output for each lag in function of the angle? Default TRUE.

The program returns an object of class "eco.correlog" with the following slots:

> OUT analysis output
> IN analysis input data
> BEAKS breaks
> CARDINAL number of elements in each class
> NAMES variables names
> METHOD analysis method
> DISTMETHOD method used in the construction of breaks
> TEST test method used (bootstrap, permutation)
> NSIM number of simulations
> PADJUST P-values adjust method for permutation tests

ACCESS TO THE SLOTS The content of the slots can be accessed with the corresponding accessor, using the generic notation of EcoGenetics (<ecoslot.> + <name of the slot> + <name of the object>). See help("EcoGenetics accessors") and the Examples section below.

Author(s)

Leandro Roser <learoser@gmail.com>
References


Geary R. 1954. The contiguity ratio and statistical mapping. The incorporated statistician, 115-146.


Examples

```r
# Not run:
data(eco.test)
require(ggplot2)

#################################################
# Moran's I correlogram
#################################################

## single test with phenotypic traits
moran <- eco.correlog(z=eco[["P"]][,1], X=Y=eco[["XY"]],
method = "1", smax=10, size=1000)

# interactive plot via plotly
eco.plotCorrelog(moran)

# standard plot via ggplot2
eco.plotCorrelog(moran, interactivePlot = FALSE)
```

#---------------------------------------------------------------
## A directional approach based in bearing correlograms

```r
moran_b <- eco.correlog(Z=eco["P"],XY = eco["XY"],
method = "I", smax = 10, size = 1000, angle = seq(0, 175, 5))
```

# use eco.plotCorrelogB for this object

```r
eco.plotCorrelogB(moran_b)
```

# plot for the first distance class,
# use a number between 1 and the number of classes to select the corresponding class

```r
eco.plotCorrelogB(moran_b, var = 1)
```

## Multivariable correlograms

```r
moran2 <- eco.correlog(Z=eco["P"],XY = eco["XY"],
method = "I", smax=10, size=1000)
```

```r
eco.plotCorrelog(moran2, var ="P2")## single plots
eco.plotCorrelog(moran2, var ="P3")## single plots
```

## Multivariable interactive plot with mean correlogram
## and jackknifed confidence intervals.

```r
graf <- eco.plotCorrelog(moran2, meanplot = TRUE)
```

# Only mean

```r
graf$mean.correlog
```

# Mean and variables

```r
graf$multi.correlog
```

# Information
- correlogram data for individual variables
- manhattan distance matrix
- mean correlogram data
- method used for analysis
- names and numbers (column in data frame) of significant variables

```r
graf$data
```

# plot only alleles

```r
graf <- eco.plotCorrelog(moran2, meanplot = FALSE)
graf
```

# Both plots can also be constructed using ggplot2
C single test with genotypic traits

# eco["A"] is a matrix with the genetic data of "eco"
# as frequencies for each allele in each individual. Each allele
# can be analyzed as single traits.
head(eco["A"])  # head of the matrix

# analyzing allele 1
moran <- eco.correlog(Z=eco["A"], X=eco["XY"], method = "I", smax=10, size=1000)
eco.plotCorrelog(moran)

# multiple tests with genotypic traits.
# nsim is set to 10 only for speed in the example
moran2 <- eco.correlog(Z = eco["A"], X = eco["XY"],
method = "I", smax=10, size=1000, nsim=99)

## multiple plot with mean
## correlogram and jackknifed
## confidence intervals.
graf <- eco.plotCorrelog(moran2, meanplot = TRUE)

## the same example, but with nsim = 99.
moran3 <- eco.correlog(Z = eco["A"], X = eco["XY"], method = "I",
smax=10, size=1000, nsim=99)

## plot for alleles with at least one significant value after
```r
## Bonferroni-Holm sequential P correction
## (set adjust "none" for no family-wise
## P correction in "eco.correlog"

eco.plotCorrelog(moran3, meanplot = TRUE, significant.M = TRUE)

# ACCESSORS USE EXAMPLE

# the slots are accessed with the generic format
# (ecoslot. + name of the slot + name of the object).
# See help("EcoGenetics accessors")

ecoslot.OUT(moran)  # slot OUT
ecoslot.BREAKS(moran)  # slot BREAKS

geary <- eco.correlog(Z = eco["P"][,1], XY = eco["XY"], method = "C",
smax=10, size=1000)
# Interactive plot
eco.plotCorrelog(geary)
# ggplot2 plot
eco.plotCorrelog(geary, interactivePlot = FALSE)

cross <- eco.correlog(Z=eco["P"][,1], XY = eco["XY"], Y = eco["P"][, 1],
method = "CC", int= 2, smax=15)
# Interactive plot
eco.plotCorrelog(cross)
# ggplot2 plot
eco.plotCorrelog(cross, interactivePlot = FALSE)

## End(Not run)
```

**eco.detrend**

*Detrending spatial data with polynomial interpolation*
Description

This program performs a Trend Surface Analysis (Borcard et al. 2011, Legendre and Legendre 2012, Lichstein et al 2002) for the data Z and the given coordinates, projected or in decimal degrees format, in which case will be projected with geoXY.

Usage

eco.detrend(Z, XY, degree = TRUE, scale = FALSE, raw = FALSE, latlon = FALSE)

Arguments

Z       Data frame, matrix or vector with dependent variables.
XY      Data frame, matrix or vector with projected coordinates (X, XY or XYZ). For longitude-latitude data in decimal degrees format, use the option latlon = TRUE.
degree  Polynomial degree.
center  Should the data be centered? Default TRUE
scale   Should the data be scaled? Default FALSE
raw     Use raw and not orthogonal polynomials? Default FALSE
latlon  Are the coordinates in decimal degrees format? Default FALSE. If TRUE, the coordinates must be in a data.frame/matrix with the longitude in the first column and latitude in the second. The position is projected onto a plane in meters with the function geoXY.

Value

An object of class ’eco.detrend’ with the following slots:
> POLY.DEG polynomial degree used in the analysis
> RES detrended data
> XY projected coordinates
> MODEL models selected with the Akaike criterion
> ANALYSIS object of class ’eco.mlm’ with the regression results for each variable

ACCESS TO THE SLOTS The content of the slots can be accessed with the corresponding accessors, using the generic notation of EcoGenetics (<ecoslot.> + <name of the slot> + <name of the object>). See help("EcoGenetics accessors") and the Examples section below.

Author(s)

Leandro Roser <learoser@gmail.com>
References


Examples

```r
## Not run:

data(eco2)

# original data
data1 <- matrix(eco2[['P']][,1], 30, 30)
image(data1)

# original data + trend
data2 <- matrix(eco2[['P']][,2], 30, 30)
image(data2)

# data detrending
data2.det <- eco.detrend(Z = eco2[['P']][,2], XY = eco2[['XY']], degree = 1)

# ACCESSORS USE EXAMPLE
# the slots are accessed with the generic format
# (ecoslot. + name of the slot + name of the object).
# See help("EcoGenetics accessors")
data2.det <- ecoslot.RES(data2.det) # detrended data in slot RES
data2.det <- matrix(data2.det[,1], 30, 30)
image(data2.det)

## End(Not run)
```

---

eco.dom.af: Compute allele frequencies for dominant data using different methods
**Description**

Compute allele frequencies for dominant data using different methods.

**Usage**

```r
eco.dom.af(x, method = c("zhivor", "zhivonu", "rawfreq"))
```

**Arguments**

- `x` : ecopop or genpop object, or matrix/data.frame with allele frequencies.
- `method` : Method used to compute the allelic frequencies. Can be one of 'zhivor' (Zhivotovsky 1999, with uniform prior), 'zhivonu' (Zhivotovsky 1999, with non-uniform prior), 'rawfreq' (square root method).

**Author(s)**

Juan Vilardi <vilardi@ege.fcen.uba.ar>

**References**


**Examples**

```r
## Not run:
data(eco.test)
my_ecopop <- ecogen2ecopop(eco, "pop")
eco.dom.af(my_ecopop)

## End(Not run)
```

---

**eco.fill_ecogen_with_df**

*Importation of data frames to ecogen*

**Description**

This function imports into an ecogen object the population data contained in a series of data frames. These data frames can be used to fill the slots XY, P, E and C.

**Usage**

```r
eo.fill_ecogen_with_df(from = pop, pop_levels = NULL, XY = NULL, P = NULL,
E = NULL, C = NULL, bind_columns = FALSE)
```
Arguments

from ecogen object
pop Name of the column of the slot S with populations.
pop_levels Vector with the name of the populations for each row of the input data frames. These populations must correspond to the levels of the column of the slot S used for the argument pop.
XY Population data for slot XY. Default NULL.
P Population data for slot P. Default NULL.
E Population data for slot E. Default NULL.
C Population data for slot C. Default NULL.
bind_columns Bind columns of the generated tables with the preexisting in the ecogen slots? Default FALSE (overwrite the slot).

Author(s)

Leandro Roser <learoser@gmail.com>

See Also

eco.fill_ecogen_with_ecopop

Examples

## Not run:

data(eco.test)

# create some population data
to_ecopop <- ecogen2ecopop(eco, "pop")
XY_pop <- to_ecopop["XY"]
P_pop <- to_ecopop["P"]
E_pop <- to_ecopop["E"]

# Add only XY data to the ecogen object
object_with_pop_data <- eco.fill_ecogen_with_df(eco, "pop", c(1,2,3,4),
       XY = XY_pop)

# Add all the population data to the ecogen object
object_with_pop_data <- eco.fill_ecogen_with_df(eco, "pop", c(1,2,3,4),
       XY = XY_pop, P = P_pop, E = E_pop)

## End(Not run)
**eco.fill_ecogen_with_ecopop**

Importation of ecopop to ecogen

**Description**

This function imports into an ecogen object the population data contained in ecopop object. The function assign the values of the data to each individual, according to the population of the individual.

**Usage**

```r
eco.fill_ecogen_with_ecopop(from, to, pop, what = c("all", "XY", "P", "E", "C"), bind_columns = FALSE)
```

**Arguments**

- `from` ecopop object.
- `to` ecogen object.
- `pop` Column in slot S of ecogen object, with the population of each individual.
- `what` Data frames to add into the the ecogen object. Can be one of c("all", "XY", "P", "E", "C")
- `bind_columns` Bind columns of the generated tables with the preexisting in the ecogen slots?

**Author(s)**

Leandro Roser <learoser@gmail.com>

**See Also**

- `eco.fill_ecogen_with_df`

**Examples**

```r
## Not run:

data(eco.test)

# Example 1: add population data to ecogen object
result <- eco.fill_ecogen_with_ecopop(my_ecopop, eco, "pop")

# Example 2: Create ecogen object only with population data
out <- ecogen(S = eco["S"],)
out <- eco.fill_ecogen_with_ecopop(my_ecopop, out, "pop")

# add the allele frequency data into the slot C with the function eco.add_popdata_into_ecogen
```
Description

This program generates a forest plot for the confidence interval of each individual of the input data (as row number) and the corresponding observed value of the statistic.

Usage

eco.forestplot(input, xlabel = NULL, ylabel = NULL, titlelabel = NULL, legendlabel = NULL, interactivePlot = TRUE, ...)

## S4 method for signature 'eco.lsa'
eco.forestplot(input, rescaled = FALSE, xlabel, ylabel, titlelabel, legendlabel, interactivePlot = TRUE)

## S4 method for signature 'dataframeORMatrix'
eco.forestplot(input, xlabel, ylabel, titlelabel, legendlabel, interactivePlot = TRUE)

Arguments

- **input**: Matrix/data frame, with three columns in the following order: observed value, lower and upper values of the confidence interval.
- **xlabel**: Optional label for x axis.
- **ylabel**: Optional label for y axis.
- **titlelabel**: Optional title label.
- **legendlabel**: Optional legend label.
- **interactivePlot**: Show an interactive plot via plotly? (default: TRUE)
- **rescaled**: rescale values to [-1, 1] range?

Author(s)

Leandro Roser <learoser@gmail.com>
Examples

```r
## Not run:

require(ggplot2)
# simulated confidence intervals for the null hypothesis of a variable "a"
set.seed(8)
a <- runif(10, -2, 2)
infer <- runif(10, -1, 1)
super <- runif(10, -1, 1)
infer2 <- pmin(infer, super)
super2 <- pmax(infer, super)
data <- data.frame(a, infer2, super2)
forest <- eco.forestplot(data)
forest

# the forestplot method support the use of ggplot2 syntax with ggplot2 graphs
forest <- eco.forestplot(data, interactivePlot = FALSE)
forest <- forest + theme_bw() + theme(legend.position="none")
forest

## End(Not run)
```

eco.format

Format tool for genetic data

Description

Format tool for genetic data

Usage

```r
eco.format(data, ncod = NULL, nout = 3, ploidy = 2, sep.in = "",
            sep.out = "", fill.mode = c("last", "first", "none"), recode = c("none",
            "all", "column", "paired"), replace_in = NULL, replace_out = NULL,
            show.codes = FALSE)
```

Arguments

data: Genetic data frame.
ncod: Number of digits coding each allele in the input file.
nout: Number of digits in the output.
ploidy: Ploidy of the data.
sep.in: Character separating alleles in the input data if present.
sep.out: Character separating alleles in the output data. Default
fill.mode Add zeros at the beginning ("first") or the end ("last") of each allele. Default = "last".
recode Recode mode: "none" for no recoding (default), "all" for recoding the data considering all the individuals values at once (e.g., protein data), "column" for recoding the values by column (e.g., microsatellite data), "paired" for passing the values of allelic states and corresponding replacement values, using the replace_in and replace_out arguments (e.g. replace_in = c("A", "T", "C", "G"), replace_out = c(1,2,3,4)).
replace_in vector with states of the data matrix to be replaced, when recode = "paired". This argument must be used in conjunction with the argument "replace_out".
replace_out vector with states of the data matrix used for replacement, when recode = "paired". This argument must be used in conjunction with the argument "replace_in".
show.codes May we returned tables with the equivalence between the old and new codes when recode = "all" or recode = "column"?

Details
The function can format data with different ploidy levels. It allows to: - add/remove zeros at the beginning/end of each allele - separate alleles with a character - divide alleles into columns - bind alleles from separate columns - transform character data into numeric data
"NA" is considered special character (not available data).
Note that the function can also work with other type of data as well, where the "alleles" represent the different states of the variables.

Author(s)
Leandro Roser <learoser@gmail.com>

Examples
```r
# Not run:
data(eco.test)

# Adding zeros
eample <- as.matrix(genotype[1:10,])
mode(eample) <- "character"
# example data
eample
recoded <- eco.format(eample, ncod = 1, ploidy = 2, nout = 3)
# recoded data
recoded

# Tetraploid data, separating alleles with a "/
tetrap <- as.matrix(eample)
# simulated tetraploid example data
tetrap <- matrix(paste(eample,eample, sep = ""), ncol = ncol(eample))
```

```
recoded <- eco.format(tetrap, ncod = 1, ploidy = 4, sep.out = "/")
# recoded data
recoded

# Example with a single character
ex <- c("A", "T", "G", "C")
ex <- sample(ex, 100, rep=T)
ex <- matrix(ex, 10, 10)
colnames(ex) <- letters[1:10]
rownames(ex) <- LETTERS[1:10]
# example data
ex
recoded <- eco.format(ex, ploidy = 1, nout = 1, recode = "all", show.codes = TRUE)
# recoded data
recoded

# Example using values-replacement pairs
samp1 <- sample(c("A", "T", "G", "C"), 100, replace = TRUE)
samp2 <- sample(c("A", "T", "G", "C"), 100, replace = TRUE)
paired <- matrix(paste0(samp1, samp2), 10, 10)
# Generate some NAs
paired[sample(1:100, 10)] <- NA
out <- eco.format(paired, recode = "paired", replace_in = c("A", "T", "G", "C"),
                  replace_out = c(1, 2, 3, 4))
out

# Example with two strings per cell and missing values:
        "Val", "Trp")
ex1 <- sample(ex, 100, rep=T)
ex2 <- paste(ex1, ex1, sep="")
missing.ex2 <- sample(1:100, 20)
ex2[missing.ex2] <- NA
ex2 <- matrix(ex2, 10, 10)
colnames(ex2) <- letters[1:10]
rownames(ex2) <- LETTERS[1:10]
# example data
ex2
recoded <- eco.format(ex2, ncod = 3, ploidy = 2,
                      nout = 2, recode = "column")
# recoded data
recoded

# Example with a vector, following the latter example:
ex1 <- as.data.frame(ex1)
# example data
ex1
recoded <- eco.format(ex1, ploidy = 1, recode = "all")
# recoded data
recoded
eco.formula

Formula construction for ecogen objects

Description
When a data frame is present in a slot of an object, any individual column can be accessed using the notation: my_object@my_data_frame[, column_to_be_accessed]. The later constitutes an explicit name for the variable present in the object. The present function generalizes this concept, allowing to construct a formula for the variables present in an ecogen object (columns of the data frames in slots). The function creates an explicit formula that can be used to parse ecogen objects into other functions (see examples below). For this purpose, each name in the formula is substituted with explicit names of columns if:
- The name corresponds to the name of an individual column in the data frames of the ecogen object, or
- The name is surrounded by U() and corresponds to the name of a slot. Complete data frames (as P, E, etc.) or subsets (as U[, 1:5]) can be passed with this method and all the columns will be explicitly included in the formula.
In other situations, names are not replaced.

Usage
eco.formula(eco, formula, out.mode = c("formula", "expression"),
expand.tables = "+")

Arguments
eco Object of class "ecogen".
formula Formula with names of columns from the slots XY, P, G, A, E, or C
out.mode Output results explicit formula (default) or expression.
expand.tables method for tables coercion. Default is "+"

Author(s)
Leandro Roser <learoser@gmail.com>

Examples
## Not run:
require(vegan)
data(eco.test)

# Note that in this example "Condition" is not replaced;
# the function Condition has a special meaning in rda, indicating conditioning variables; in eco.formula it is only text.
form <- eco.formula(eco, P1 + P2 + P3 + U(A) ~ E1 + E2 + Condition(X+Y))
rda(form)
form2 <- eco.formula(eco, P1 + P2 + P3 + U(A) ~ E1 + E2 + X + Y)
lm(form2)

# parsing with magrittr
eco.formula(eco, P1 + P2 + P3 + U(A) - U(E) + Condition(X+Y)) %>% rda

## End(Not run)

---

**eco.gsa**

*Global spatial analysis*

---

**Description**

Univariate and multivariable global spatial analysis. This program computes Moran’s I, Geary’s C, Bivariate Moran’s I or Join-count statistics with P-values. The program allows the analysis of a single variable or multiple variables. In this latter last case, the variables must be in columns and the individuals in rows.

In join-count analysis, a ploidy argument must be supplied. The data is then ordered with the function `aue.sort`. This step is required for the analysis of genotypic data. An individual with the alleles A and B, coded as AB, is identical to other coded as BA. The order step ensures the AB and BA will be considered the same genotype. For the analysis of frequencies of single alleles, the input is count data (ploidy times the frequency, as provided by the slot A of an ecogen object; the count data A’ can be obtained as A’ <- ploidy * A), using the function with the arguments ploidy = 1.

**Usage**

```r
eco.gsa(Z, con, Y = NULL, method = c("I", "C", "CC", "JC"), nsim = 99,
alternative = c("auto", "two.sided", "greater", "less"), ploidy = 1,
adjust = "fdr", plotit = TRUE)
```

**Arguments**

- **Z** Vector with a variable, or matrix/data frame with variables in columns.
- **con** An object of class eco.weight obtained with the function `eco.weight`, a listw object or a matrix, giving the spatial weights for the analysis. If "con" is a matrix, an attribute "xy" including the projected coordinates is required.
- **Y** Vector with the second variable for Moran’s Ixy. If Z has multiple variables, the program will compute the coefficient for each with Y.

nsim: Number of Monte-Carlo simulations.

alternative: The alternative hypothesis. If "auto" is selected (default) the program determines the alternative hypothesis. Other options are: "two.sided", "greater" and "less".

ploidy: For join count analysis: number of elements for the values of the vector passed, given value: for example, if ploidy=1, "13" and "31" are considered a same level ("31" is sorted by the program as "13"); if ploidy = 1, "13" and "31" represent two different levels.

adjust: Correction method of P-values for multiple tests, passed to p.adjust. Default is "fdr".

plotit: should be generated a plot for univariate results?

Value

The program returns an object of class "eco.gsa" with the following slots:

> METHOD method used in the analysis
> OBS observed value when a single variable is tested
> EXP expected value when a single variable is tested
> PVAL P-value when a single variable is tested
> ALTER alternative hypothesis when a single variable is tested
> NSIM number of simulations
> MULTI table with observed and expected values, P-values and alternative hypotheses when multiple variables are tested

ACCESS TO THE SLOTS The content of the slots can be accessed with the corresponding accessor, using the generic notation of EcoGenetics (<ecoslot.> + <name of the slot> + <name of the object>). See help("EcoGenetics accessors") and the Examples section below.

Author(s)

Leandro Roser <learoser@gmail.com>

References

Geary R. 1954. The contiguity ratio and statistical mapping. The incorporated statistician, 115-146.
Examples

```r
## Not run:

data(eco.test)

# Moran's I

### one test
con <- eco.weight(eco["XY"], method = "circle", d1 = 0, d2 = 2)
global <- eco.gsa(Z = eco["P"], 1, con = con, method = "I", nsim = 200)
global

require adegenet
con2<-chooseC(eco["XY"], type = 1, result.type = "listw", plot.nb = FALSE)
global <- eco.gsa(Z = eco["P"], 1, con = con2, method = "I", nsim = 200)
global

# ACCESSORS USE EXAMPLE

# the slots are accessed with the generic format
# (ecoslot. + name of the slot + name of the object).
# See help("EcoGenetics accessors")

# observed value
ecoslot.OBS(global)

# p-value
ecoslot.PVAL(global)

# multiple tests

# data(eco3)
con <- eco.weight(eco3["XY"], method = "circle", d1 = 0, d2 = 500)
global <- eco.gsa(Z = eco3["P"], con = con, method = "I", nsim = 200)
global

# Plot method for multivariable eco.gsa objects:
eco.plotGlobal(global)

# accessor use in multiple tests

ecoslot.MULTI(global)

# Geary's s C
```
Kinship and relationship estimation following Hardy (2003).

```r
con <- eco.weight(eco["XY"], method = "circle", d1 = 0, d2 = 2)
global.C <- eco.gsa(Z = eco["P"], 1), con = con, method = "C", nsim = 200)
global.C

#----------------------------------------------

# Bivariate's Moran's Ixy

con <- eco.weight(eco["XY"], method = "circle", d1 = 0, d2 = 2)
global.Ixy <- eco.gsa(Z = eco["P"], 1), Y = eco["E"], 1),
con = con, method = "CC", nsim = 200)
global.Ixy

#----------------------------------------------

# Join-count

## using the allelic frequency matrix of an ecogen object.
## The data is diploid. Frequencies are transformed into counts
## as ploidy * frequency_matrix:

Z = 2* eco["A"]

jc <- eco.gsa(Z, 1), con = con, method = "JC")
eco.plotGlobal(jc)

# multiple tests
# using the first ten alleles of the matrix

global.JC <- eco.gsa(Z, 1:10), con = con, method = "JC", nsim = 99)
global.JC

# plot method for multivariable join-count
eco.plotGlobal(global.JC)

# counting joins between genotypes in the first locus the G matrix:

global.JC <- eco.gsa(Z = eco["G"], 1), ploidy = 2, con = con, method = "JC", nsim = 99)
global.JC
eco.plotGlobal(global.JC)

## End(Not run)
```

Description

Kinship and relationship estimation following Hardy (2003).
eco.kin.loiselle

Usage

eco.kin.hardy(x, fi)

Arguments

x ecogen, genind matrix or data.frame. In case of matrix or data frame, the data consists in a table with individuals in rows and allele counts in columns.

fi Assumed Fi

Value

List with three slots containing, respectively, the heretability of each locus, relationship and kinship values

Author(s)

Juan Vilardi <vilardi@ege.fcen.uba.ar>

References


Examples

```r
# Not run:

data(eco.test)
loiselle <- eco.kin.loiselle(eco)
loiselle[1:5, 1:5]

# End(Not run)
```

eco.lagweight  

Obtention of a list of spatial weights for classes defined by inter-individual distances or nearest-neighbors

Description

This program returns a list of weights matrices (binary or row-standardized), one for each spatial class. For a given maximum and minimum inter-individual distance (IID), the data can be partitioned in different ways. The program set as default the highest IID as the maximum, and the lowest as the minimum. These values can be changed with "smax" and "smin", respectively. Intervals may be generated with the parameters "int" (which divides the range each int distance units), "nclass" (which divides the range in n-classes) and "size" (a fixed size of pairs included in each class). When a partition argument is not given (int, nclass or size) the program determines the number of classes using the Sturge's rule (default) or the Freedman- Diaconis method. Two additional methods can be used: a list with nearest-neighbors matrices, from 1 to k nearest-neighbors, may be generated with the argument "kmax". A custom vector with breaks may be provided by the user with the argument "seqvec". See the examples.

Usage

```r
eco.lagweight(XY, int = NULL, smin = 0, smax = NULL, kmax = NULL, 
nclass = NULL, seqvec = NULL, size = NULL, bin = c("sturges", "FD"), 
cumulative = FALSE, row.sd = FALSE, self = FALSE, latlon = FALSE)
```

Arguments

<table>
<thead>
<tr>
<th>XY</th>
<th>Matrix/data frame with projected coordinates.</th>
</tr>
</thead>
<tbody>
<tr>
<td>int</td>
<td>Distance interval in the units of XY.</td>
</tr>
<tr>
<td>smin</td>
<td>Minimum class distance in the units of XY.</td>
</tr>
<tr>
<td>smax</td>
<td>Maximum class distance in the units of XY.</td>
</tr>
<tr>
<td>kmax</td>
<td>Number of nearest-neighbors.</td>
</tr>
<tr>
<td>nclass</td>
<td>Number of classes.</td>
</tr>
<tr>
<td>seqvec</td>
<td>Vector with breaks in the units of XY.</td>
</tr>
</tbody>
</table>
size
bin
cumulative
row.sd
self
latlon

Value

The program returns an object of class "eco.lagweight" with the following slots:

> PAR parameters used for the construction of breaks
> PAR.VAL values of the parameters used for the construction of breaks
> ROW.SD row standardization (logical)
> SELF data self-included (logical)
> W weights list
> XY coordinates
> MEAN mean class distances
> LOGMEAN mean of the class distances logarithm
> CARDINAL number of elements in each class
> BREAKS breaks
> METHOD breaks construction method

ACCESS TO THE SLOTS The content of the slots can be accessed with the corresponding accessors, using the generic notation of EcoGenetics (<ecoslot.> + <name of the slot> + <name of the object>). See help("EcoGenetics accessors") and the Examples section below.

Author(s)

Leandro Roser <learoser@gmail.com>

References


Examples

```r
## Not run:
data(eco.test)

# method sturges-smax: in this case, the program generates
# classes using the Sturge's rule.
# As smax and smin are undefined, the program uses the default
# options (smin = 0, and smax = maximum inter-individual distance)
classlist <- eco.lagweight(eco["XY"])
classlist

# method sturges-smax: idem, but smax = 16
classlist <- eco.lagweight(eco["XY"], smax=16)
## using smax <16 in this case generates empty classes,
## which is not allowed

classlist <- eco.lagweight(eco["XY"], smin = 3, smax = 15)

# method sturges-smax: complete range,
# and cumulative = TRUE (instead of using
# lower and upper limits for each class, only the upper is used in turn)
classlist <- eco.lagweight(eco["XY"], cumulative = TRUE)

# method n.classes-smax: complete range partitioned in 4 classes
classlist <- eco.lagweight(eco["XY"], nclass = 4)

# method n.classes-smax: idem, but smax = 15
classlist <- eco.lagweight(eco["XY"], nclass = 4, smax = 15)

# method int-smax: the complete range partitioned each <int> units
# of inter-individual distance
classlist <- eco.lagweight(eco["XY"], int = 2)

# method int-smax: idem, but smax = 15 and smin = 3
classlist <- eco.lagweight(eco["XY"], int = 2, smin = 3, smax = 15)

# method equal.size: n individuals in each class,
# partitioning the complete range.
classlist <- eco.lagweight(eco["XY"], size = 1000)

## In the latter example, as an inter-individual distance
## appear more than one time (different individuals pairs, 
## identical distances), with a size <700 the limits
## of some classes cannot be defined, and this is not allowed

# method equal.size: n individuals in each class,
# but smax = 15
classlist <- eco.lagweight(eco["XY"], size = 1000, smax = 15)

# method kmax: sequence from k = 1 to k = n, in this case, n = 3
```

eco.listw2ew <- eco.lagweight(eco["XY"], kmax = 3)

# method kmax: idem, but elements self-included
# (i.e., the pairs i-i, for all individuals i, are included)
classlist <- eco.lagweight(eco["XY"], kmax = 3, self = TRUE)

# method seqvec: a vector with the breaks is used
vec <- seq(0, 10, 2)
classlist <- eco.lagweight(eco["XY"], seqvec = vec)

# ACCESSORS USE EXAMPLE
#----------------------------------
# the slots are accessed with the generic format
# (ecoslot. + name of the slot + name of the object).
# See help("EcoGenetics accessors")
ecoslot.BREAKS(classlist) # information about breaks. It includes the upper and lower limits

## End (Not run)

---

eco.listw2ew Conversion from listw to ecoweight

Description
Conversion from listw to ecoweight

Usage
eco.listw2ew(X)

Arguments

X A listw object

Author(s)
Leandro Roser <learoser@gmail.com>
Description

This program fits for each dependent variable, a Multiple Linear Regression model calling the function \texttt{step} for choosing the best model by AIC criterion, or a Multiple Classification and Regression Trees model, using the package \texttt{party}. The summary of the model returns information about the significance of the models, F-statistics and degrees of freedom, when is fitted a "mlm"; otherwise, when the model fitted is a "mctree", the summary returns the plots of those trees with significant splits.

Usage

\begin{verbatim}
eco.lmtree(df1, df2, analysis = c("mlm", "mctree"), mod.class = "+",
          fact = NULL, ...)
\end{verbatim}

Arguments

\begin{itemize}
\item \texttt{df1} -- Data frame with dependent variables as columns.
\item \texttt{df2} -- Data frame with independent variables as columns.
\item \texttt{analysis} -- Class of analysis to perform. "mlm" for multiple linear regression analysis, or "mctree" for a multiple classification tree analysis.
\item \texttt{mod.class} -- "+" for additive model, "\*" for model with interaction, in both cases, these models will include all terms in the dependent data frame. If other model than these two is desired, it could be specified as a string with the names of those columns of the independent variable that should be used as terms. This string corresponds to the right side "x" of a formula \(y \sim x\) (see examples).
\item \texttt{fact} -- Optional factor for estimating the frequencies of individuals from different levels in each node, when the analysis performed is "mctree".
\item ... -- Further arguments passed to \texttt{lm} or \texttt{ctree}
\end{itemize}

Value

When the analysis selected is "mlm", the output object has three main slots:
\begin{itemize}
\item \texttt{MLM}: the results of the model
\item \texttt{SUMMARY.MLM}: the summary for each variable returned by the \texttt{lm} function
\item \texttt{ANOVA.MLM}: with the ANOVAs results.
\end{itemize}
When the analysis selected is "mctree", the output object has also three main slots:
\begin{itemize}
\item \texttt{TREES}: Trees returned by the multiple \texttt{ctree} analysis.
\item \texttt{PREDICTIONS}: Predictions of the analysis.
\item \texttt{FREQUENCIES}: Number of individuals predicted in each node.
\end{itemize}
ACCESS TO THE SLOTS The content of the slots can be accessed with the corresponding accessors, using the generic notation of EcoGenetics (<ecoslot.> + <name of the slot> + <name of the object>). See help("EcoGenetics accessors") and the Examples section below

Author(s)
Leandro Roser <learoser@gmail.com>

References

Examples

```r
## Not run:

data(eco.test)

# mml additive model
mod <- eco.lmtree(df1 = eco3["P"], df2 = eco3["E"],
                  analysis = "mlm")
mod
summary(mod)

# mctree additive model
mod <- eco.lmtree(df1 = eco3["P"], df2 = eco3["E"],
                  analysis = "mctree", fact = eco3["S"]$pop)
summary(mod)

# ACCESSORS USE EXAMPLE

# the slots are accessed with the generic format
# (ecoslot. + name of the slot + name of the object).
# See help("EcoGenetics accessors")

summary(mod)

ecoslot.FREQUENCIES(mod)  # slot FREQUENCIES

# frequency table with counts of individuals in populations x terminal nodes
tabfreq <- do.call(cbind, ecoslot.FREQUENCIES(mod))
nametab <- lapply(ecoslot.FREQUENCIES(mod), ncol)
nametab <- lapply(nametab, rep)
nametab <- rep(names(nametab), nametab)
colnames(tabfreq) <- nametab
tabfreq

## mlm custom model

```
eco.lock, ecogen-method

Lock rows in an ecogen object

Description

This method locks the rows in an ecogen object. When rows are locked, the object requires rows with identical individuals in the non-empty data frames, and identity in the row names of the data frames.

Usage

## S4 method for signature 'ecogen'
eco.lock(object, set.names = NULL, valid.names = FALSE, order.df = FALSE)

Arguments

<table>
<thead>
<tr>
<th>Argument</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>object</td>
<td>object of class ecogen</td>
</tr>
<tr>
<td>set.names</td>
<td>Character vector with names for the rows of the non-empty data frames. This argument is incompatible with valid.names</td>
</tr>
<tr>
<td>valid.names</td>
<td>Logical. Create valid row names? This argument is incompatible with set.names. The program will name individuals with valid tags I.1, I.2, etc.</td>
</tr>
<tr>
<td>order.df</td>
<td>Order individuals of data frames by row? (all data frames with a same order in row names). This option is only available when the 'lock.rows' parameter is TRUE. If the names of the data frames are not used (i.e., set.names and valid.names are not NULL), setting this parameter to TRUE/FALSE has no effect in the function. Default TRUE. If FALSE, the row names of all the data frames must be ordered. The use of data frames with row names in different order will return an error. In both cases, the program sets an internal names attribute of the object using the row names of the first non-empty data frame found in the following order: XY, P, G, E, S, C. This attribute is used as reference to order rows when order.df = TRUE.</td>
</tr>
</tbody>
</table>

Examples

```r
## Not run:
data(eco.test)
eco2 <- eco.unlock(eco)
is.locked(eco2)
eco3 <- eco.lock(eco2)
is.locked(eco3)

## End(Not run)
```

Description

This method locks the rows in an ecogen object. When rows are locked, the object requires rows with identical individuals in the non-empty data frames, and identity in the row names of the data frames.

Usage

```r
## S4 method for signature 'ecopop'
eco.lock(object, set.names = NULL, valid.names = FALSE, 
          order.df = FALSE)
```

Arguments

- **object**: ecopop object
- **set.names**: Character vector with names for the rows of the non-empty data frames. This argument is incompatible with valid.names
- **valid.names**: Logical. Create valid row names? This argument is incompatible with set.names. The program will name individuals with valid tags I.1, I.2, etc.
- **order.df**: Order individuals of data frames by row? (all data frames with a same order in row names). This option is only available when the 'lock.rows' parameter is TRUE. If the names of the data frames are not used (i.e., set.names and valid.names are not NULL), setting this parameter to TRUE/FALSE has no effect in the function. Default TRUE. If FALSE, the row names of all the data frames must be ordered. The use of data frames with row names in different order will return an error. In both cases, the program sets an internal names attribute of the object using the row names of the first non-empty data frame found in the following order: XY, P, G, E, S, C. This attribute is used as reference to order rows when order.df = TRUE.
Examples

```R
## Not run:
data(eco.test)
my_ecopop2 <- eco.unlock(my_ecopop)
is.locked(my_ecopop2)
my_ecopop3 <- eco.lock(my_ecopop)
is.locked(my_ecopop3)

## End(Not run)
```

### eco.lsa

**Local spatial analysis**

**Description**

Univariate and multivariable local spatial analysis. This program computes Getis-Ord G and G*, and LISA's (local Moran and local Geary) statistics for the data Z, with P-values or bootstrap confidence intervals.

**Usage**

```R
eco.lsa(var, con, method = c("Gx", "G", "I", "C"), zerocon = NA,
        nsim = 99, conditional = c("auto", "TRUE", "FALSE"),
        test = c("permutation", "bootstrap"), alternative = c("auto", "two.sided",
                     "greater", "less"), adjust = "none", multi = c("matrix", "list"),
        pop = NULL)
```

**Arguments**

- **var**: Vector, matrix or data frame for the analysis. Multiple variables in columns.
- **con**: An object of class eco.weight obtained with the function `eco.weight`, a "listw" object, or a matrix, containing the weights for the analysis. If a matrix, an attribute "xy" with the projected coordinates is required.
- **method**: Method of analysis: "G" for Getis-Ord G, "G*" for Getis-Ord G*, "I" for local Moran's I or "C" for local Geary's C.
- **zerocon**: If zerocon = 0 the program assigns the value 0 to those individuals with no connections; if zerocon = NA the program assigns NA. Default is NA.
- **nsim**: Number of Monte-Carlo simulations.
- **test**: If test = "bootstrap", for each individual test, the program generates a bootstrap resampling and the associated confidence intervals of the null hypothesis. If test = "permutation" (default) a permutation test is made and the P-value is computed.
alternative
The alternative hypothesis for "permutation" test. If "auto" is selected (default) the program determines the alternative hypothesis in each individual test. Other options are: "two.sided", "greater", and "less".

adjust
Correction method of P-values for multiple tests, passed to `p.adjust`. Default is "none" (no correction).

multi
multiple output format results. "list" for object with a list of individual test for each variable, or "matrix" for results as matrices of multiples variables.

pop
numeric factor with the population of each individual. Optional for multiple tests with multi = "matrix".

Value
For single test, the program returns an object of class "eco.lsa" with the following slots:

> OUT results - table with output results.

-> If test = "permutation": observed value of the statistic, null confidence interval and rescaled observed value to [-1, 1] range, as in Sokal (2006)

-> If test = "bootstrap": observed and expected value of the statistic, alternative hypothesis, null confidence interval and rescaled observed value to [-1, 1] range, as in Sokal (2006)

> METHOD method (coefficient) used in the analysis
> TEST test method used (bootstrap, permutation)
> NSIM number of simulations
> PADJUST P-values adjust method for permutation tests
> COND conditional randomization (logical)
> XY input coordinates

For multiple test, if the parameter multi = "list", the program returns a list of eco.lsa objects (one element for each variable).

For multiple test, if the parameter multi = "matrix", the program returns an object of class "eco.multilsa" with the following slots:

> METHOD method used in the analysis
> TEST test method used (bootstrap, permutation)
> NSIM number of simulations
> PADJUST P-values adjust method for permutation tests
> COND conditional randomization (logical)
> XY input coordinates
> OBS observed value
> EXP expected value
> ALTER test alternative
> PVAL pvalue for permutation test
> LWR lower confidence interval bound of the null hypothesis
> UPPR upper confidence interval bound of the null hypothesis
> OBS.RES rescaled observed value to [-1, 1] range, as in Sokal (2006)

**ACCESS TO THE SLOTS** The content of the slots can be accessed with the corresponding accessors, using the generic notation of EcoGenetics (<ecoslot.> + <name of the slot> + <name of the object>). See help("EcoGenetics accessors") and the Examples section below

**Author(s)**

Leandro Roser <learoser@gmail.com>

**References**


**Examples**

```r
## Not run:
data(eco.test)
#---------------------------------------------------------------#

#-------------------------
# LOCAL MORAN'S I
#-------------------------

DETAILED EXAMPLE

#-----------------------------
# TESTING PHENOTYPIC DATA-
#-----------------------------

set.seed(10)

# test for a single variable-----------------------------
#computing weights

con <- eco.weight(eco["XY"], method = "knearest", k = 4, row.sd = TRUE)
# row standardized weights = TRUE
```
```r
# test for the first trait of the data frame p
localmoran <- eco.lsa(eco[["P"]][, 1], con, method = "I", nsim = 99)

# "rankplot" graph
eco.plotLocal(localmoran)

# test for several variables
eco <- eco[order(eco[["S"]][, 1])]

# computing weights with the ordered object
con <- eco.weight(eco[["XY"]], method = "knearest", k = 4, row.sd = TRUE)
# row standardized weights = TRUE

all.traits <- eco.lsa(eco[["P"]], con, method = "I", nsim = 99)

# Plot of the phenotypic spatial patterns
eco.plotLocal(all.traits)

# in grey: non significant results (P > 0.05)
# set significant = FALSE for showing significant and no significant results
eco.plotLocal(all.traits, significant = FALSE)

# single plots using "rankplot" graphs
all.single.traits <- eco.lsa(eco[["P"]], con, method = "I", nsim = 99, multi="list")
eco.plotLocal(all.single.traits)

# removing legends for a better visualization
eco.plotLocal(all.single.traits, legend = FALSE)
# - individual plots support ggplot2 sintax (plot equivalent to the previous):
eco.plotLocal(all.single.traits) + ggplot2::theme(legend.position="none")

# TESTING GENOTYPIC DATA-

# eco[["A"]]) is a matrix with the genetic data of "eco"
# as frequencies for each allele in each individual.

head(eco[["A"]])  # head of the matrix - 40 alleles

# ordering the factor "pop" in increasing order and the object "eco"
# in relation to this ordered factor prior to the multivariate analysis.
# This step is important for "localplot" graphs
```
data(eco.test) # for security this resets the data (unordered)

eco <- eco[order(eco["S"],[1])] # ordering

# computing weights with the ordered object

con <- eco.weight(eco["XY"], method = "knearest", k = 4, row.sd = TRUE) # row standardized weights = TRUE

# test for a single allele
localmoran.geno <- eco.lsa(eco["A"], con, method = "I", nsim = 99)

# test for several alleles - 40 alleles (it runs in less than 1 min
# for 99 simulations per allele; 999 simulations takes ~ 11 s per allele,
# less than 8 min in total.)
all.alleles <- eco.lsa(eco["A"], con, method = "I", nsim = 99)

# plot all alleles to get an overview of the spatial patterns
eco.plotLocal(all.alleles)

# in grey: non significant results (P > 0.05)
# set significant = FALSE for showing significant and no significant results
eco.plotLocal(all.alleles, significant = FALSE)

# counting individuals with P < 0.05 for each allele (5 * 225 / 100 ~ 12 significant tests
# by random)
signif <- apply(ecoslot.pval.all.alleles, 2, function(x) sum(x < 0.05))

# filtering alleles, loci with > 12 significant individual tests
A.local <- eco["A"], signif > 12] #filtered matrix

all.alleles.f <- eco.lsa(eco["A"], signif > 12], con, method = "I", nsim = 99)

# Plot of the genotypic spatial patterns using "localplot" graphs
eco.plotLocal(all.alleles.f)

## using "rankplot" graphs

all.sf <- eco.lsa(A.local, 2, eco.lsa, con, method = "I", nsim = 99, multi = "list")
eco.plotLocal(all.sf, legend = FALSE)

##########################
# GETIS-ORD'S G* 
##########################

con <- eco.weight(eco["XY"], method = "knearest", k = 4, self = TRUE) # self = TRUE for G*
getis.ak <- eco.lsa(eco["P"], con, method = "G*", nsim = 99, adjust = "none")
getis.ak
### to plot the results, the function "eco.lsa" calls "eco.rankplot"
### (see ?eco.rankplot) when test = "permutation" and "eco.forestplot" (see ?eco.forestplot)
### when test = "bootstrap"

```r
p <- eco.plotLocal(getis.ak)  # rankplot graph
p  # points with colors of the color-scale:
   # points with P < 0.05. Yellow points: points with P > 0.05
p <- eco.plotLocal(getis.ak, significant = FALSE)
p  # all points have a color of the color-scale
```

# ACCESSORS USE EXAMPLE
#---------------------------------------------

# the slots are accessed with the generic format
# (ecoslot + name of the slot + name of the object).
# See help("EcoGenetics accessors")

```r
ecoslot.OUT(getis.ak)
```

## bootstrap example
```r
ggetis.akb <- eco.lsa(eco["P"][, 1], con, method = "Gx", nsim = 99, test = "bootstrap")
p  <- eco.plotLocal(getis.akb)  # forestplot graph
p2 <- eco.plotLocal(getis.akb, interactivePlot = FALSE)
p2 + ggplot2::theme_bw()  # the plot can be modified with ggplot2
   # In this case, the background is modified (white color)
```

#---------------------------------------------

### GETIS-ORD'S G
###---------------------------------------------

```r
con <- eco.weight(eco["XY"], method = "knearest", k = 4)
# self = FALSE for G
ggetis <- eco.lsa(eco["P"][, 1], con, method = "G", nsim = 99)
eco.plotLocal(getis)
```

#---------------------------------------------

### LOCAL GEARY'S C
###---------------------------------------------

```r
con <- eco.weight(eco["XY"], method = "knearest", k = 4, row.sd = TRUE)
# row standardized weights = TRUE
localgeary <- eco.lsa(eco["P"][, 1], con, method = "C", nsim = 99, adjust = "none")
eco.plotLocal(localgeary)
```

## End(Not run)
Description

The program computes, for a kinship matrix, a global multilocus correlogram, or a local analysis. When a kinship matrix is not given as input, the program computes the Loiselle’s Fij (Kalisz et al., 2001; Loiselle et al., 1995). The program can compute a bearing correlogram (Rosenberg 2000, Born et al. 2012) for the obtention of a directional approach in the global test.

Usage

```r
eco.malecot(eco, method = c("global", "local"), kinmatrix = NULL,
          int = NULL, smin = 0, smax = NULL, nclass = NULL, kmax = NULL,
          seqvec = NULL, size = NULL, type = c("knearest", "radialdist"),
          cubic = TRUE, testclass.b = TRUE, testmantel.b = TRUE,
          jackknife = TRUE, cummulative = FALSE, normLocal = TRUE, nsim = 99,
          test = c("permutation", "bootstrap"), alternative = c("auto", "two.sided",
                   "greater", "less"), sequential = TRUE, conditional = c("AUTO", "TRUE",
                   "FALSE"), bin = c("sturges", "FD"), row.sd = FALSE, adjust = "holm",
          latlon = FALSE, angle = NULL)
```

Arguments

- **eco**: Object of class ecogen.
- **method**: Analysis method: "global" or "local".
- **kinmatrix**: Alternative kinship matrix. The program computes the Loiselle’s kinship matrix (codominant data) with the genetic information of the ecogen object if kinmatrix = NULL (Default option).
- **int**: Distance interval in the units of XY.
- **smin**: Minimum class distance in the units of XY.
- **smax**: Maximum class distance in the units of XY.
- **nclass**: Number of classes.
- **kmax**: Number of nearest-neighbors for local analysis.
- **seqvec**: Vector with breaks in the units of XY.
- **size**: Number of individuals per class.
- **type**: Weighting mode for local analysis: "knearest" for nearest neighbors, "radialdist" for radial distances. Default is knearest.
- **cubic**: Should a cubic interpolation (res~ ln(dij)) be performed, for the regression residuals (res) of (kinship)ij ~ ln(dij) ? Default TRUE.
- **testclass.b**: Carry a permutation test within each individual class? Default TRUE.
**testmantel** Should a Mantel test for testing the slope (b) be performed? Default TRUE.

**jackknife** Compute jackknife within each individual class for obtention of the standard deviation (SD) of the coancestry (class) values. Default TRUE.

**cummulative** Should a cummulative correlogram be constructed?.

**normLocal** Normalize the local kinship values ([(local_kinship-mean)/sd]?). Default TRUE

**nsim** Number of Monte-Carlo simulations.

**test** If test = "bootstrap", the program generates a bootstrap resampling and the associated confidence intervals of the null hypothesis. If test = "permutation" (default) a permutation test is made and the P-values are computed.

**alternative** The alternative hypothesis. If "auto" is selected (default) the program determines the alternative hypothesis. Other options are: "two.sided", "greater" and "less".

**sequential** Use the Holm-Bonberroni sequential method for adjustment of P-values (Legendre and Legendre, 2012) in global analysis? Default TRUE.


**bin** Rule for constructing intervals when a partition parameter (int, nclass or size) is not given. Default is Sturge's rule (Sturges, 1926). Other option is Freedman-Diaconis method (Freedman and Diaconis, 1981).

**rowNsd** Logical. Should be row standardized the matrix? Default FALSE (binary weights).

**adjust** P-values correction method for multiple tests passed to `p.adjust`. Default is "holm".

**latlon** Are the coordinates in decimal degrees format? Default FALSE. If TRUE, the coordinates must be in a matrix/data frame with the longitude in the first column and latitude in the second. The position is projected onto a plane in meters with the function `geoxy`.

**angle** direction for computation of a bearing correlogram (angle in degrees between 0 and 180). Default NULL (omnidirectional).

### Details

The GLOBAL ANALYSIS mode, computes a multilocus correlogram, with a detailed summary (see the content of the slot OUT in the "return" section). It also computes (see details about the slot SP in the "return" section): - the slope of the kinship individual values vs the logarithm of the distance, (kinship)ij ~ ln(dij), with a jackknife confidence interval - a Mantel test for testing the association between (kinship)ij and ln(dij) - The Sp statistic (Vekemans and Hardy, 2004) with confidence intervals - A cubic interpolation of (kinship)ij ~ ln(dij) residuals vs ln(dij)

A directional approach is based on the bearing analysis method, and consists in the obtention of a directional correlogram using the method of Rosenberg (2000). A slope is computed for the logarithm of D' (Born et al 2012), where D' is the distance matrix between individuals weighted by cos(alpha - B)^2, being alpha the angle between individuals and B the desired direction angle. With B = 0 the direction analyzed follows the positive x axis, with B = 0 the positive y axis, and with B = 180 the negative x axis, respectively.

The LOCAL ANALYSIS mode, computes a local kinship estimate, based in a weighted mean (for each individual). The significance of each local statistic is computed using a permutation test,
as in eco.lsa (see ?"eco.lsa"). Default option do not adjust the individual P values for multiple comparisons.

**Value**

For the global analysis, the program returns an object of class "eco.IBD" with the following slots:

- **OUT analysis output.**
  - d.mean: mean class distance;
  - d.log: mean logarithm of the class distance;
  - obs, exp, alter: observed, and expected value of the statistic under randomization, alternative, P value;
  - mean.jack, sd.jack, Jack.CI.inf, Jack.CI.sup: jackknifed mean and SD, and confidence intervals for the statistic;
  - null.lwr, nul.uppr: lower and upper bound of the jackknife confidence interval for the statistic;
  - cardinal: number of individuals in each class;

In the permutation test case contains:
- d.mean: mean class distance;
- d.log: mean logarithm of the class distance;
- obs, exp, alter: observed, and expected value of the statistic under randomization, alternative, P value;
- mean.jack, sd.jack, Jack.CI.inf, Jack.CI.sup: jackknifed mean and SD, and confidence intervals for the statistic;
- null.lwr, nul.uppr: lower and upper bound of the jackknife confidence interval for the statistic;
- cardinal: number of individuals in each class;

> **GLOBALTEST** Oden’s (1984) global test of significance for the correlogram. The test consists in checking if the most significant kinship coefficient is significant at a Bonferroni- corrected significance level of alpha’ = alpha/k, where k is the number of distance classes of the correlogram; alpha is set to 0.05. The program return the values: "SIGNIFICANT" or "NOT-SIGNIFICANT"

> **IN analysis input data**

> **SP** Sp statistic results

It contains:
- the regression model;
- information about the distance interval used for the regression (restricted);
- slope (bhat) information (bhat = estimate, SD= bhat jackknife SD, theta = bhat jackknife mean, CI 5% and 95% = 95% confidence interval for bhat);
- X-intercept = dij intercept (in the original units) for the line with slope "bhat", F1 = first class statistic value, and F1 5% and 95% = confidence interval for the first class statistic;
- mantel.obs.b = observed value of the Mantel test between kinship(Fij) and ln(dij); mantel.pval.b = Mantel test P value;
- sp = Sp statistics (sp = Sp observed value, CI 5% and 95% = 95% confidence interval for Sp);
- cubic_model = cubic model for (kinship)ij ~ ln(dij) r esiduals vs ln(dij);

> **BEAKS** breaks

> **CARDINAL** number of elements in each class

> **NAMES** variables names

> **METHOD** analysis method

> **DISTMETHOD** method used in the construction of breaks

> **TEST** test method used (bootstrap, permutation)

> **NSIM** number of simulations

> **PADJUST** P-values adjust method for permutation tests

For the local analysis, the program returns an object of class "eco.lsa" with the following slots:

> **OUT results**
> In the permutation test case it contains:
- d.mean: mean class distance - obs, exp, alter, p.val: observed, and expected value of the statistic under randomization, alternative, P value; - null.lwr, nul.uppr: lower and upper bound of the jackknife confidence interval for the statistic; - cardinal: number of individuals in each class;
> In the bootstrap test case it contains: - d.mean: mean class distance; - obs: observed value of the statistic; - null.lwr, nul.uppr: lower and upper bound of the jackknife confidence interval for the statistic; - cardinal: number of individuals in each class;
> METHOD method (coefficient) used in the analysis
> TEST test method used (bootstrap, permutation)
> NSIM number of simulations
> PADJUST P-values adjust method for permutation tests
> COND conditional randomization (logical)
> XY input coordinates

ACCESS TO THE SLOTS The content of the slots can be accessed with the corresponding accessors, using the generic notation of EcoGenetics (<ecoslot.> + <name of the slot> + <name of the object>). See help("EcoGenetics accessors") and the Examples section below.

Author(s)
Leandro Roser <learoser@gmail.com>

References

Examples

```
## Not run:
data(eco.test)
```
# ---global analysis---

globaltest <- eco.malecot(eco=eco, method = "global", smax=10, size=1000)
eco.plotCorrelog(globaltest)  # Significant mean class coancestry classes at
# individual level (alpha = 0.05, 
# out of the red area),
# and family-wise P corrected values (red-blue
# points, indicated in the legend)

# ecoslot.SP(globaltest) contains:
# - the slope (bhat) and values with confidence intervals
# of the regression reg = kinship - ln(distance_between_individuals)
#- A Mantel test result for assessing the relation between
# between kinship and ln(distance_between_individuals)
#- A cubic interpolation between the residuals of reg and
# ln(distance_between_individuals)
#- the sp statistic and its confidence interval

# ecoslot.OUT(globaltest) contains:
# - In permutation case, the values of mean and log-mean distance
# classes; observed class value; expected + alternative + P value,
# the bootstrap null confidence intervals and
# jackknife statistics (jackknifed mean, jackknifed SD, and
# CI for the class statistic)

# - In bootstrap case, the values of mean and log-mean distance
# classes; the bootstrap null confidence intervals and
# jackknife statistics (jackknifed mean, jackknifed SD, and
# CI for the class statistic)

# A directional approach based in bearing correlograms, 30 degrees
globaltest_30 <- eco.malecot(eco=eco, method = "global", smax=10, size=1000, angle = 30)
eco.plotCorrelog(globaltest)

#--------------------------------------------#
# ---local analysis---

# (using the spatial weights).

# ---local analysis with k nearest neighbors---

localktest <- eco.malecot(eco=eco, method = "local", type = "knearest", kmax = 5, adjust = "none")
eco.plotLocal(localktest)
# ---local analysis with radial distance---

localdtest <- eco.malecot(eco=eco, method = "local",
    type = "radialdist", smax = 3,
    adjust = "none")

eco.plotLocal(localdtest)  # rankplot graphic (see ?"eco.rankplot")
    # Significant values
    # in blue-red scale,
    # non significant
    # values in yellow

eco.plotLocal(localktest, significant = FALSE)  # significant and non
    # significant values
    # in blue-red scale

# The slot OUT of localktest (ecoslot.OUT(localktest)) and localdtest
# (ecoslot.OUT(localdtest)) contains:
# - the mean distance per individual, observed value of the
# statistic, expected + alternative + P value + null hypotesis
# confidence intervals, or bootstrap confidence intervals in
# permutation or bootstrap cases, respectively.

## End(Not run)

---

**eco.mantel**

*Mantel and partial Mantel tests, with truncation option*

---

**Description**

Mantel test or Partial Mantel test for distance matrices d1 and d2, or partial Mantel test for d1 and d2, conditioned on the matrix dc. The test can be performed for truncated distances (Legendre et al. 2015) or for a particular direction (Falsetti and Sokal, 1993) using a weights object generated with *eco.bearing*.

**Usage**

```r
eco.mantel(d1, d2, dc = NULL, con = NULL, thres = NULL,
            truncMat = c("d2", "d1", "dc"), method = c("pearson", "spearman",
            "kendall"), nsim = 99, alternative = c("auto", "two.sided", "less",
            "greater"), plotit = TRUE, ...)
```

**Arguments**

- **d1** Distance matrix.
- **d2** Distance matrix.
Distance matrix (optional).

con
Binary eco.weight object used for truncation, or a weights object obtained with eco.bearing.

thres
Threshold distance used for truncation. Distances above the threshold are set as 4 times the threshold. If thres is null, and con is not null, the parameter set to the maximum distance observed in d2.

truncMat
Matrix used for truncation (default = d2)

method
Correlation method used for the construction of the statistic ("pearson", "spearman" or "kendall"). Kendall’s tau computation is slow.

nsim
Number of Monte-Carlo simulations.

alternative
The alternative hypothesis. If "auto" is selected (default) the program determines the alternative hypothesis. Other options are: "two.sided", "greater" and "less".

plotit
Plot a histogram of the simulations?

Value
An object of class "eco.gsa" with the following slots:

> METHOD method used in the analysis
> OBS observed value
> EXP expect value
> PVAL P-value
> ALTER alternative hypotesis
> NSIM number of simulations

ACCESS TO THE SLOTS The content of the slots can be accessed with the corresponding accessors, using the generic notation of EcoGenetics (<ecoslot.> + <name of the slot> + <name of the object>). See help("EcoGenetics accessors") and the Examples section below

Author(s)
Leandro Roser <learoser@gmail.com>

References


Examples

## Not run:

```r
data(eco.test)

### Ordinary Mantel test ###
eco.mantel(d1 = dist(eco["P"])), d2 = dist(eco["E"])), nsim = 99)

### Partial Mantel test ###
pm <- eco.mantel(d1 = dist(eco["P"])), d2 = dist(eco["E"])),
dc = dist(eco["XY"])), nsim = 99)

### Truncated Mantel test ###
# checking threshold in a correlogram:
corm <- eco.cormantel(M = dist(eco["P"])), XY = eco["XY"], nsim = 99)
eco.plotCorrelog(corm)
# Correlation is around 0 when distance between points is > 5

# creating a weights object for truncation
con <- eco.weight(eco@XY, method="circle", d2=5)
# compute a truncated mantel test
eco.mantel(dist(eco["P"])), dist(eco["XY"]), con=con)

### Directional Mantel test ###
# analyzing with a Mantel test, in a direction of 35 degrees counterclockwise from E.
con2 <- eco.bearing(XY = eco["XY"], theta = 37)
eco.mantel(dist(eco["P"])), dist(eco["XY"]), con = con2)

# ACCESSORS USE EXAMPLE
# ACCESSORS USE EXAMPLE

# the slots are accessed with the generic format
# (ecoslot. + name of the slot + name of the object).
# See help("EcoGenetics accessors")
ecoslot.OBS(pm)  # slot OBS (observed value)
ecoslot.PVAL(pm)  # slot PVAL (P-value)

## End(Not run)
**Description**

Merging two ecogen objects. Ordering the rows of an ecogen object according to the rows of another

**Usage**

```r
eco.merge(e1, e2, ...)
```

**Arguments**

- `e1`: Object of class "ecogen".
- `e2`: Object of class "ecogen".
- `...`: Data frames to merge. Could be any combination of the following: "XY", "P", "G", "E" and "C", or "ALL". If a "G" data frame is provided, the program generates also the INT slot by coding the missing data as "0".

**Details**

This program generates an ecogen object by binding the columns of the individuals that have matching row names in the objects `e1` and `e2`. If the objects have different number of rows, the result is a merged data frame with the rows in the order of the first object. If the objects have the same number of rows, but in a different order, the product is an object with the rows ordered as the first object. The algorithm matches sequentially the data frame pairs of each slot that the user wishes to merge.

**Author(s)**

Leandro Roser <learoser@gmail.com>

**Examples**

```r
## Not run:
data(eco.test)
eco
eco1 <- eco[2:28]
merged <- eco.merge(eco, eco1)
merged

## End(Not run)
```

---

**eco.NDVI**

*Generation of atmospherically corrected NDVI and MSAVI2 images for temporal series of Landsat 5 and 7*
Description

This program generates atmospherically corrected images of NDVI and MSAVI2. The images of multiple dates can be processed in a single run. The images for the bands 4 and 3 corresponding to each date (previously subsetted to the region of analysis) must be in the working directory. A table with information for each image as described in the parameter tab (see also the example) is needed for processing the information.

Usage


Arguments

tab data.frame with 7 columns: The date of the images (format: YYYY/MM/DD), the sun elevation (both values could be extracted from Landsat headers), the name of the band 4, the name of the band 3, the starting haze value of the band 4, the starting haze value of the band 3, and the name of the output file. Each row corresponds to an image of different date.

correct Correction method ("COST", "DOS").

method Vegetation index ("NDVI", "MSAVI2").


datatype type of data, see dataType. Default "FLT4S".

Author(s)

Leandro Roser <learoser@gmail.com>

References


Examples

```r
## Not run:
require(raster)

data(tab)

temp <- list()

# we create 4 simulated rasters for the data included in the object tab:

for(i in 1:4) {
  temp[[i]] <- runif(19800, 0, 254)
  temp[[i]] <- matrix(temp[[i]], 180, 110)
  temp[[i]] <- raster(temp[[i]], crs="+proj=utm")
  extent(temp[[i]])<-c(3770000, 3950000, 6810000, 6920000)
}

writeRaster(temp[[1]], "20040719b4.tif", overwrite=T)
writeRaster(temp[[2]], "20040719b3.tif", overwrite=T)
writeRaster(temp[[3]], "20091106b4.tif", overwrite=T)
writeRaster(temp[[4]], "20091106b3.tif", overwrite=T)

# Computing NDVI images:

eco.NDVI(tab, "COST", "NDVI", "LT5")

example <- raster("NDVICOST20040719.tif")
image(example)

file.remove(c("NDVICOST20040719.tif", "NDVICOST20091106.tif",
  "20040719b4.tif", "20040719b3.tif", "20091106b4.tif",
  "20091106b3.tif")

## End(Not run)
```

### Postprocessing for NDVI and MSAVI2 temporal series of Landsat 5 and 7

**Description**

This program must be used sequentially after eco.NDVI. The inputs required (tab, correct, method) are the same described and used in that function. The algorithm stacks the images and save the stack into the working directory with the name "time.tif". If the user wishes, the program can also compute images consisting in the max, min, mean and var for each pixel across the temporal sequence. Default is "mean".
Usage

```r
eco.NDVI.post(tab, correct = c("COST", "DOS"), method = c("NDVI", "MSAVI2"),
               datatype = c("FLT4S", "FLT8S", "INT4U", "INT4S", "INT2U", "INT2S", "INT1U",
                           "INT1S", "LOG1S"), what = c("mean", "max", "min", "var", "none"))
```

Arguments

- `tab`: Table used with `eco.NDVI`.
- `correct`: Correction method used in `eco.NDVI`.
- `method`: The vegetation index used in `eco.NDVI`.
- `datatype`: Type of data, see `dataType`. Default "FLT4S".
- `what`: Functions to apply over the created stack. The allowed values are: "none", "max", "min", "mean" and "var". The functions are implemented with `calc`. For passing more than one function as argument, the following syntax must be used: `c("fun_1", "fun_2", "fun_i")`, where fun_1...fun_n are the functions that you want to compute.

Author(s)

Leandro Roser <learoser@gmail.com>

References


See Also

- `eco.NDVI`
- `extract`
### Examples

```r
## Not run:
require(raster)

data(tab)
data(eco3)
temp <- list()

# we create 4 simulated rasters for the data included in the object tab:

for(i in 1:4) {
  temp[[i]] <- runif(19800, 0, 254)
  temp[[i]] <- matrix(temp[[i]], 180, 110)
  temp[[i]] <- raster(temp[[i]], crs="+proj=utm")
  extent(temp[[i]])<-c(3770000, 3950000, 6810000, 6920000)
}

writeRaster(temp[[1]], "20040719b4.tif", overwrite = T)
writeRaster(temp[[2]], "20040719b3.tif", overwrite = T)
writeRaster(temp[[3]], "20091106b4.tif", overwrite = T)
writeRaster(temp[[4]], "20091106b3.tif", overwrite = T)

# Computing NDVI images:

eco.NDVI(tab, "COST", "NDVI", "LT5")

# Mean NDVI image computed over the NDVI images that we calculated:

eco.NDVI.post(tab, "COST", "NDVI", what = c("mean", "var"))
mean.ndvi <- raster("NDVI.COST.mean.tif")
plot(mean.ndvi)

# Extraction of the mean NDVI for each point in the object eco and plot # of the data:

ndvi <- extract(mean.ndvi, eco3["XY"])
ndvi<- aue.rescale(ndvi)
plot(eco3["XY"], 1, eco3["XY"], 2, col=rgb(ndvi, 0, 0),
pch=15, main = "Mean NDVI", xlab = "X", ylab = "Y")

## End(Not run)

file.remove(c("NDVICOST20040719.tif", "NDVICOST20091106.tif",
"20040719b4.tif", "20040719b3.tif", "20091106b4.tif",
"20091106b3.tif", "NDVI.COST.mean.tif", "NDVI.COST.var.tif",
"NDVICOSTtime.tif"))
```

---

### eco.nei_dist

Estimate Nei distance matrix
 eco.old2new,ecogen-method

Description

Estimate Nei distance matrix. NAs are avoided.

Usage

eco.nei_dist(obj, as_dist = TRUE)

Arguments

obj ecopop or genpop objects, or matrix/data frame with allele frequencies
as_dist Return a dist object or a matrix? default is an object of class "dist"

Author(s)

Juan Vilardi <vilardi@ege.fcen.uba.ar>

Examples

## Not run:
data(eco.test)
eco.nei_dist(my_ecopop)

## End(Not run)

eco.old2new,ecogen-method

Update an old ecogen or ecopop object to version >= 1.5.0-1

Description

Update an old ecogen or ecopop object to version >= 1.5.0-1

Usage

## S4 method for signature 'ecogen'
eco.old2new(object)

Arguments

object ecogen object
Description

Update an old ecogen or ecopop object to a version compatible with EcoGenetics \(>= 1.5.0-1\)

Usage

```
## S4 method for signature 'ecopop'
eco.old2new(object)
```

Arguments

- `object`: ecopop object

Description

Functions deprecated in EcoGenetics version 1.2.0-2

Usage

```
eco.order(...)  # parameters
```

Arguments

- `...`: parameters
eco.pairtest

Kruskall - Wallis + Wilcoxon (Mann-Whitney U) and aov + Tukey-HSD tests for an ecogen object

Description

Kruskall - Wallis + Wilcoxon (Mann-Whitney U) and aov + Tukey-HSD tests for an ecogen object

Usage

eco.pairtest(eco, df = c("P", "E", "A", "C"), x, test = c("wilcoxon", "tukey"), adjust = "fdr", only.p = TRUE, ...)

Arguments

- eco: Object of class "ecogen".
- x: The name of the S slot column with the groups for the analysis.
- test: Test to perform ("wilcoxon", "tukey").
- adjust: P-values correction method for multiple tests passed to p.adjust. Default is "fdr".
- only.p: Should it be just a matrix with P-values returned? Default TRUE.
- ...: Additional arguments passed to wilcox.test or TukeyHSD.

Details

This program returns the Wilcoxon (Mann-Whitney U) or Tukey-HSD statistics and P-values for the multiple comparisons of the variables contained in the selected data frame, among the levels of a factor of the slot "S".

Author(s)

Leandro Roser <learoser@gmail.com>

See Also

wilcox.test TukeyHSD

Examples

```r
# Not run:
data(eco3)
wil <- eco.pairtest(eco = eco3, df = "P", x = "structure")
wil
wil <- eco.pairtest(eco = eco3, df = "E", x = "structure")
```
Wil
wil <- eco.pairtest(eco = eco3, df = "P", x = "structure", only.p = FALSE)
wil
wil <- eco.pairtest(eco = eco3, df = "P", x = "structure", test = "tukey")
wil

## End(Not run)

eco.plotCorrelog

---

### Description

Plot method for correlograms and variograms. For examples, see `eco.correlog` `eco.cormantel` `eco.variogram`

### Usage

```r
eco.plotCorrelog(x, var = NULL, xlabel = NULL, ylabel = NULL, title = NULL, legend = TRUE, background = c("grey", "white"), errorbar = FALSE, intervals = TRUE, significant.S = TRUE, significant.M = FALSE, xlim = NULL, ylim = NULL, nsim = 999, interactivePlot = TRUE, meanplot = TRUE, randtest = c("permutation", "bootstrap", "none"), alpha = 0.05, quiet = FALSE)
```

### Arguments

- **x**: Result of correlogram or variogram analysis
- **var**: Individual variable to plot for multiple analyses with `eco.correlog`. To plot multiple variables in a same plot, use only the argument `x` (see examples)
- **xlabel**: Label for X axis (default: NULL)
- **ylabel**: Label for Y axis (default: NULL)
- **title**: Title of the plot (default: NULL)
- **legend**: Show legends in ggplot graphs? (default: TRUE)
- **background**: Background color ("grey" or "white")
- **errorbar**: Show error-bars? (default: FALSE)
- **intervals**: Show bootstrap CI in kinship analysis? (default: TRUE)
- **significant.S**: With single variables and permutation test: show different colours for significant points? (default: TRUE)
- **significant.M**: With multiple variables: show only significant correlograms? (default: FALSE)
- **xlim**: X axis limits (as vector: c(min, max); default: NULL)
- **ylim**: Y axis limits (as vector: c(min, max); default: NULL)
- **nsim**: Number of simulations for permutation or bootstrap tests.
eco.plotCorrelogB

interactivePlot
Show an interactive plot via plotly? (default: TRUE)

meanplot
Show a line with the mean, when the plot is for multiple variables? (default: TRUE)

randtest
Randomization test (one of: "permutation", "bootstrap", "none")

alpha
significance level for P (or P-adjusted) values (Default alpha = 0.05)

quiet
print quietly? Default FALSE

Author(s)
Leandro Roser <learoser@gmail.com>

See Also
eco.correlog eco.cormantel eco.variogram

description
Plot method for bearing correlograms For examples, see eco.correlog. It constructs an angular correlogram for each distance class taken as fixed.

Usage
eco.plotCorrelogB(x, var = NULL, xlabel = NULL, ylabel = NULL, title = NULL, legend = TRUE, background = c("grey", "white"), significant.N = TRUE, xlim = NULL, ylim = NULL, interactivePlot = TRUE, alpha = 0.05)

Arguments

x
Result of correlogram analysis, with output using angles as independent variables for fixed distances (instead of distances as independent variables)

var
Individual variable to plot; var is a number between 1 and the number of distance classes indicating the corresponding class (for example, with 5 distance classes, the number 3 indicates the third) To plot multiple variables in a same plot, use only the argument x (see examples)

xlabel
Label for X axis (default: NULL)

ylabel
Label for Y axis (default: NULL)

title
Title of the plot (default: NULL)

legend
Show legends in ggplot graphs? (default: TRUE)

background
Background color ("grey" or "white")
significant.S With single variables and permutation test: show different colours for significant points? (default: TRUE)
xlim X axis limits (as vector: c(min, max); default: NULL)
ylim Y axis limits (as vector: c(min, max); default: NULL)
interactivePlot Show an interactive plot via plotly? (default: TRUE)
alpha significance level for P (or P-adjusted) values (Default alpha = 0.05)

Author(s)
Leandro Roser <learoser@gmail.com>

See Also
eco.correlog

description
This function allows to plot results contained in eco.gsa objects. For examples, see eco.gsa

Usage
eco.plotGlobal(input, interactivePlot = TRUE, background = c("grey", "white"), xlabel = NULL, ylabel = NULL, title = NULL, legend = TRUE, rescaled = FALSE, alpha = 0.05)

Arguments
input eco.gsa object
interactivePlot Show an interactive plot via plotly? (default: TRUE)
background background color ("grey" or "white")
xlabel Label for X axis (default: NULL)
ylabel Label for Y axis (default: NULL)
title Title of the plot (default: NULL)
legend Show legends in ggplot graphs? (default: TRUE)
rescaled rescale join-count heatmap?
alpha significance level for the join-count heatmat

Author(s)
Leandro Roser <learoser@gmail.com>
Description

For examples see eco.lsa

SINGLE VARIABLES:
Using permutation test: The function calls eco.rasterplot, who generates a plot for a numeric or factor variable. The X and Y axes in the plot correspond to the rank of the X and Y coordinates, respectively. Additional parameters can be passed to eco.rankplot.

Using bootstrap test: The function calls eco.forestplot, who computes a forest plot for the confidence interval of each individual of the input data (as row number) and the corresponding observed value of the statistic. Additional parameters can be passed to eco.forestplot.

MULTIPLE VARIABLES: multiple output format results. "list" for object with a list of individual test for each variable, or "matrix" for results as matrices of multiples variables.

For results as matrices (option multi = "matrix" in eco.lsa): The function class eco.rasterplot, who generates a multivariate plot for a data matrix (raster). Additional parameters can be passed to eco.rasterplot. The resterplot graph is a flexible tools for multiple data sources (environmental, genetic, phenotypic, etc.).

For results as list (option multi = "list" in eco.lsa): The function generates plots for individual variables calling eco.rankplot. Additional parameters can be passed to eco.rankplot.

Usage

eco.plotLocal(x, interactivePlot = TRUE, multi = c("d3heatmap", "ggplot"),
              significant = TRUE, alpha = 0.05, rescaled = FALSE, limits = NULL,
              title = NULL, z.name = NULL, grp = NULL, vertical = TRUE,
              legend = TRUE, n = 4, nrow = 2, byrow = TRUE, ...)

Arguments

x
  Result of eco.lsa analysis

interactivePlot
  Show an interactive plot via plotly? (default: TRUE)

multi
  for multivariable plot, use d3heatmap or ggplot2? (Default: d3heatmap). In
d3heatmap, NA values are set to 0.

significant
  Show all non significant points with a same colour?

alpha
  significance (alpha) for P (or P-adjusted) values (Default: 0.05)

rescaled
  rescale statistics between -1 and 1? (Default: FALSE)

limits
  When multiple variables are used, values used as limits for computing the gra-
dient for the plot

title
  title of the plot

z.name
  name of the variables axis in multivariable plot (using ggplot2 like plots)
eco.plotWeight

Plot method for an eco.weight object. For examples, see eco.weight

This function can make a static plot with the original coordinates and an additional graph with the coordinates transformed as ranks. It can also construct dynamic plots (force networks and circle networks).

Usage

eco.plotWeight(x, type = c("simple", "igraph", "edgebundle", "network"),
group = NULL, fontSize = 10, ebColor = NULL, vertex.size = 10,
vertex.label = NA, bounded = FALSE, ...)

Arguments

x Connection network
type Plot type: "edgebundle", for a circular network, "network" for a tension network
group Vector with classes assigned to the individuals, in the same original order
fontSize Argument passed to forceNetwork contained in the weight object (which is the order of the table used to construct the weights)
ebColor Vector with edge colors for the groups of the edgebundler plot (Experimental feature)
vertex.size Parameter to plot.igraph
vertex.label Parameter passed to plot.igraph
bounded Logical. Value to enable (TRUE) or disable (FALSE) the bounding box limiting the force network graph extent see forceNetwork.
...

Author(s)

Leandro Roser <learoser@gmail.com>
Author(s)
Leandro Roser <learoser@gmail.com>

Examples

# see the examples in the function eco.weight:
# ?eco.weight

Description
Log posterior probability plot for Geneland repetitions with fixed K

Usage
eco.post.geneland(niter, burnin)

Arguments
niter Number of mcmc iterations per repetition.
burnin Number of mcmc to burn-in.

Details
This program returns, for a series of Geneland repetitions with fixed K, and a specified burn-in value, a plot of the log posterior probability vs the repetition number. This allows to choose the best run. The working directory will be set to the folder containing the results created by Geneland. The program expects each subfolder (run) to have a number as name, that indicates the corresponding number of run. (1, 2, etc., see the example).

Author(s)
Leandro Roser <learoser@gmail.com>

Examples

## Not run:
require("Geneland")
data(eco.test)

# We create a folder in the working directory for the results and
# save the data frames of the object "eco" in the format required
# by Geneland:

path.1 <- getwd()
path <- paste(path.1,"/test/", sep="")
```r
dir.create(path)
setwd(path)
ecogen2geneland(eco, ploidy = 2)

# Auxiliar function for running some repetitions with fixed K = 4.
# Each repetition is saved in the folder "test":
simul <- function(i) {
  path <- getwd()
  path <- paste(path, "/", i, sep = "")
  dir.create(path)
  MCMC(coordinates = read.table("XY.txt"),
       geno.dip.codom = read.table("G.txt"),
       vrnnpop = TRUE, npopmin = 4, npopmax = 4, spatial = TRUE,
       freq.model = "Correlated", nit = 500, thinning = 10,
       path.mcmc = path)
}

# 5 repetitions with K = 4
lapply(1:5, simul)

# Check that in the folder "test" are the simulated results.
# Your results must have that appearance.

# Plot of the repetition order number vs the corresponding
# posterior probability, with a burn-in of 10 mcmc:
eco.post.geneland(5, 10)

## End(Not run)
```

### eco.rankplot

**Rankplot graphs**

**Description**

This function generates a plot for a numeric or factor variable. A data frame/matrix with XY coordinates is required. The X and Y axes in the plot correspond to the rank of the X and Y coordinates, respectively.

**Usage**

```r
eco.rankplot(input, XY, xlabel = NULL, ylabel = NULL, title = NULL,
             legendlabel = NULL, background = c("grey", "white"), ...)
```

## S4 method for signature 'eco.lsa,missing,missing'
```
eco.rankplot(input, XY, xlabel, ylabel,
             title, legendlabel, background = c("grey", "white"), significant = TRUE,
             rescaled = FALSE, ns = NULL, interactivePlot = TRUE)
```
eco.rankplot

## S4 method for signature 'numeric, dataframe or matrix, missing'
eco.rankplot(input, XY, xlabel, ylabel, title, legendlabel, background = c("grey", "white"), interactivePlot = TRUE)

## S4 method for signature 'factor, dataframe or matrix, missing'
eco.rankplot(input, XY, xlabel, ylabel, title, legendlabel, background = c("grey", "white"), interactivePlot = TRUE)

**Arguments**

- `input` Numeric/factor variable.
- `XY` Data frame or matrix with X-Y coordinates.
- `xlabel` Optional label for x axis.
- `ylabel` Optional label for y axis.
- `title` Optional title label.
- `legendlabel` Optional legend label.
- `background` Background color ("grey" or "white").
- `...` Additional elements to the generic.
- `significant` Should only the individuals with significant result be colored? This argument can be used with `eco.lsa` results. Default TRUE
- `rescaled` rescale values to [-1, 1] range?
- `ns` Color for non significant individuals, when significant = TRUE. This argument can be used with `eco.lsa` results.
- `interactivePlot` Show an interactive plot via plotly? (default: TRUE)

**Author(s)**

Leandro Roser <learoser@gmail.com>

**Examples**

```r
## Not run:
data(eco3)

# The data set eco3 has 50 points in two sites,
# but points are not visible in a usual X-Y plot,
# due to the small distance among them in relation to the large
# distance between sites

var <- eco3["P",1]
plot(eco3["XY"], col = var)
x <- sample(1:100, 30)
y <- sample(1:100, 30)
```
# in a rankplot graph, the inter-individual distances are
# reduced to a single scale
rankeco3 <- eco.rankplot(var, eco3[["XY"]], interactivePlot = FALSE)
rankeco3 <- rankeco3 + theme_bw() + theme(legend.position="none")
## End(Not run)

## Description

This function generates a multivariate plot for a data matrix (raster), with an option for filtering the data and to plot using groups. The rasterplot graph is a flexible tool useful for different data sources.

## Usage

eco.rasterplot(x, filter = NULL, condition = NULL, grp = NULL,
               limits = NULL, title = NULL, z.name = NULL, vertical = TRUE,
               interactivePlot = TRUE, ...)

## Arguments

x                  Data matrix (raster)
filter             Optional data matrix used as filter
condition          Condition used to filter data
grp                Factor with groups to use in the plot. Default NULL
limits             Values limits used for computing the data gradient for the plot
title              Plot title
z.name             Name for the legend
vertical           Should the populations on the x axis be partitioned? Default TRUE.
interactivePlot    Show an interactive plot via plotly? (default: TRUE)
...                 additional arguments
Examples

```r
## Not run:
data(eco.test)
require(ggplot2)

# using the ecogen object "eco" to perform a multiple-lsa
con <- eco.weight(eco["XY"], method = "knearest", k = 4, row.sd = TRUE)
test.lsa <- eco.lsa(eco["P"], con = con, method = "I", nsim = 99, multi = "matrix")

# the plot method for this object based in ggplot2, is a rasterplot
eco.plotLocal(test.lsa, multi = "ggplot2")

# adding a factor
test.lsa <- eco.lsa(eco["P"], con = con, method = "I",
nsim = 99, multi = "matrix", pop = eco["S"][,1])
eco.plotLocal(test.lsa, multi = "ggplot2")

# The generic rasterplot method requires a data matrix, and, as option, a condition
# and a filter matrix. The condition is an expression, containing the word "filter" and
# logical elements, e.g., "filter < 50", "filter <50 || filter > 2", etc.).
# Filter is used as a logical matrix (TRUE-FALSE, in relation to the passed condition),
# for filtering the data. If a condition is passed but not a filter matrix, the condition
# is applied over the data matrix, also using the word "filter".
# Internally, the multi.lsa plot uses three fundamental elements.
# - a data matrix: in the example, ecoslot.OBS(test.lsa)
# - a filter matrix: in the example, ecoslot.PVAL(test.lsa): i.e.,
# the data matrix will be filtered by P-value using the third element, an expression.
# - an expression: in the example: "filter < 0.05"

# by combining the three elements, the multivariate plot can be manually constructed:
my.plot <- eco.rasterplot(x = ecoslot.OBS(test.lsa),
filter = ecoslot.PVAL(test.lsa), condition = "filter < 0.05")
my.plot

# add population
my.plot <- eco.rasterplot(x = ecoslot.OBS(test.lsa),
filter = ecoslot.PVAL(test.lsa),
condition = "filter < 0.05", grp = ecoslot.POP(test.lsa))
my.plot

# extra manipulation with ggplot2 graphs (ggplot2 commands allowed by rasterplot)
my.plot <- eco.rasterplot(x = ecoslot.OBS(test.lsa),
filter = ecoslot.PVAL(test.lsa), condition = "filter < 0.05",
interactivePlot = FALSE)
my.plot

## rotate plot
```
my.plot + coord_flip()

## change design
my.plot + theme_grey()

# using the data as filter
eco.rasterplot(x= ecoslot.OBS(test.lsa), filter = ecoslot.OBS(test.lsa),
condition = "filter > 0 & filter < 3")

# example of bad syntax (incorrect use of && over matrices)
eco.rasterplot(x= ecoslot.OBS(test.lsa), filter = ecoslot.OBS(test.lsa),
condition = "filter > 0 && filter < 3")

## End(Not run)

eco.rasterplot, eco.multilsa-method

\textit{rasterplot graph for eco.lsa results}

\section*{Description}

Plot method for local spatial analysis

\section*{Usage}

### S4 method for signature 'eco.multilsa'
eco.rasterplot(x, grp = NULL, limits = NULL,
title = NULL, z.name = NULL, vertical = TRUE, significant = TRUE,
rescaled = FALSE, alpha = 0.05, interactivePlot = TRUE, ...)

\section*{Arguments}

\begin{itemize}
  \item \textbf{x} eco.multilsa object returned by \texttt{eco.lsa} or
  \item \textbf{grp} factor with groups to use in the plot. Default NULL
  \item \textbf{limits} values limits used for computing the data gradient for the plot
  \item \textbf{title} plot title
  \item \textbf{z.name} name for the legend
  \item \textbf{vertical} should be partitioned the populations on the x axis? Default TRUE.
  \item \textbf{significant} plot only significant results? Default TRUE
  \item \textbf{rescaled} plot the rescaled observed values ([-1,1] range)?
  \item \textbf{alpha} threshold P value for results with permutation tests. default = 0.05.
  \item \textbf{interactivePlot} Show an interactive plot via plotly? (default: TRUE)
  \item \ldots additional arguments
\end{itemize}
eco.rbind

Author(s)
Leandro Roser <learoser@gmail.com>

See Also
eco.lsa

Description
Combining ecogen objects by row

Usage
eco.rbind(..., check_colnames = TRUE, check_rownames = TRUE)

Arguments
... "ecogen" objects to combine.
check_colnames Check for duplicated column names? Default TRUE.
check_rownames Check for duplicated row names? Default TRUE.

Author(s)
Leandro Roser <learoser@gmail.com>

Examples

## Not run:

data(eco.test)

# split the object "eco" into a list of ecogen objects by population
x <- eco.split(eco, "pop", asList = TRUE)

# re-bind the objects
eco.r <- eco.rbind(eco)

# create a new objects with the first and second population
eco.r <- eco.rbind(x[[1]], x[[3]])

# duplicated row names are not allowed by eco.rbind with default options
eco2 <- eco
eco.rbind(eco, eco2)
eco.remove

eco.remove(eco, ...)  

Arguments

eco  
Object of class "ecogen".

...  
Objects to remove from eco, typed without quotations.

Author(s)

Leandro Roser <learoser@gmail.com>

Examples

## Not run:

data(eco.test)
variog <- eco.variogram(eco["P"][, 1], eco["XY"])

# the assignment of values can be made with the corresponding accessors,
# using the generic notation of EcoGenetics
# (<ecoslot.> + <name of the slot> + <name of the object>).
# See help("EcoGenetics accessors")

ecoslot.OUT(eco) <- variog
we.are.numbers <- c(1:10)
we.are.characters <- c("John Coltrane", "Charlie Parker")
ecoslot.OUT(eco) <- list(w.
we.are.numbers, w.
we.are.characters)
ecoslot.OUT(eco)
eco <- eco.remove(eco, we.
we.are.numbers)
ecoslot.OUT(eco)

## End(Not run)
**Sliding a window along a connection network**

**Description**

This program applies a function defined by the user, over the individuals included in a connection network. For a given variable, the program computes recursively a function for the individuals of the network, using all the individuals connected to each. The function uses a connection network generated with the function `eco.weight`.

**Usage**

```
eco.slide.con(x, con, fun)
```

**Arguments**

- **x**: Input variable or matrix.
- **con**: Connection network.
- **fun**: Function to apply in each focal point.

**Author(s)**

Leandro Roser <learoser@gmail.com>

**Examples**

```r
## Not run:

data(eco2)
myMatrix <- eco2[["P"]]
con <- eco.weight(XY = eco2[["XY"]], method = "knearest", k = 5)
result <- eco.slide.con(myMatrix, con, function(x) mean(x, na.rm = TRUE))

image(matrix(myMatrix[, 1], 30, 30)) # original image
image(matrix(result[, 1], 30, 30)) # smoothed image

data(eco3)
myMatrix2 <- eco3[["P"]]
con <- eco.weight(XY = eco3[["XY"]], method="knearest", k = 5)
eco.plotWeight(con)
# smoothing values in myMatrix2 using the connection network:
result <- eco.slide.con(myMatrix2, con, function(x) mean(x, na.rm = TRUE))

## End(Not run)
```
eco.slide.matrix  

Sliding window for matrix data

Description

This program applies a function defined by the user, using a moving window (circle area or square) and assigning the value to the focal pixel.

Usage

eco.slide.matrix(mat, r, slide, fun, window = c("square", "circle"), within = TRUE)

Arguments

mat  
Input raster matrix.

r  
Half a side for square, radius for circle, diagonal length for rhombus.

slide  
Number of elements between two focal pixels, for column and row dimension

fun  
Function to apply in each focal pixel.

window  
Window type. Default "square".

within  
Should the function be computed in focal pixels of the borders, only if the area is within the matrix? Default TRUE.

Author(s)

Leandro Roser <learoser@gmail.com>

Examples

```r
# Not run:

data(eco.test)
ras <- matrix(eco["P"][, 1], 15, 15)
image(ras)
ras.square <- eco.slide.matrix(ras, 1, 1, mean, "square")
image(ras.square)

# or allowing more control over the function:
ras.square <- eco.slide.matrix(ras, r = 3, slide = 1, function(x) mean(x, na.rm = TRUE), "square")
image(ras.square)

# sliding a circle:
ras.circle <- eco.slide.matrix(ras, r = 3, slide = 1, mean, "circle", within = FALSE)
image(ras.circle)

# End(Not run)
```
ecosplit

eco.split

Splitting an ecogen object by group

Description

The function splits an ecogen object into the groups defined in the slot S. If asList is TRUE, a list
with the objects is created, that can be assigned to a name with regular rules, using the operator
"<-". Otherwise, the function creates in the workspace an ecogen object for each group with the
following nomenclature: <name of ecogen object>.<name of the group>.

Usage

eco.split(eco, hier, name = NULL, overwrite = FALSE, missing = c("0",
"NA", "MEAN"), asList = TRUE)

Arguments

eco Object of class "ecogen".
hier The name of the S slot column with labels assigning individuals to groups.
name Name used for the output objects. Default is the name of the input, followed by
a suffix (see Description).
overwrite Overwrite files with the same name of the output if already present in workspace
when asList = FALSE? Default FALSE.
missing Missing data argument This can take three values ("0", "NA" or "MEAN"), as
described in ecogen. # Missing elements are treated as zeros in the default
option.
asList Return a list with the objects instead of creating objects in workspace? Default
= TRUE

Author(s)

Leandro Roser <learoser@gmail.com>

Examples

## Not run:
data(eco3)
eco3

# list of objects
x <- eco.split(eco3, "structure", asList = TRUE)

# rebinding
eco.bind <- eco.rbind(x)
# Note that different subsets can also be created
S1.3 <- eco.rbind(x[[1]], x[[3]])

# Split and create objects with prefix "eco3"
eco.split(eco3,"structure", asList = FALSE)

# Split and create objects with prefix "newObjects"
eco.split(eco3,"structure", "newObjects", aslist = FALSE)

## End(Not run)

---

**eco.subset**

*Subsetting an ecogen object by group*

### Description

Subsetting an ecogen object by group

### Usage

```r
eco.subset(eco, hier, grp, missing = c("0", "NA", "MEAN"))
```

### Arguments

- `eco` Object of class "ecogen".
- `hier` The name of the S slot column containing labels assigning individuals to groups.
- `grp` Label shared by the subset of individuals, contained in hier.
- `missing` Missing data argument. It can take three values ("0", "NA" or "MEAN"), as described in `ecogen`. Missing elements are treated as zeros in the default option.

### Author(s)

Leandro Roser <learoser@gmail.com>

### Examples

```r
## Not run:
data(eco3)
eco3
eco.sub <- eco.subset(eco3,"structure", 1)
eco.sub
```
eco.theilsen

Theil-sen regression for a raster time series, with parallelization available

Description
This function computes the theil-sen estimator and the associated P-value, for each pixel over time in a stack of images. The output consists of two rasters (one for the estimators and one for the P-values). It is recommended to use a "RasterBrick", which is more efficient in memory management. The program can compute the result using serial (default) or parallel evaluation. For parallel evaluation, the program uses PSOCK cluster for windows, and FORK cluster for other operative systems.

Usage
eco.theilsen(stack, dates, adjust = "none", run_parallel = FALSE, workers = NULL, physical = FALSE, cl_type = NULL)

Arguments
stacked Stacked images ("RasterLayer" or "RasterBrick").
dates Data vector with decimal dates for each image.
adjust P-values correction method for multiple tests. Passed to p.adjust. Default is "none".
run_parallel Run code in parallel? Default FALSE
workers Number of workers used for parallel evaluation. If NULL, the program uses N - 1, where N is the total number of available logical cores.
physical Use only physical cores for parallel evaluation? Default FALSE.
cl_type Cluster type. If not specified, "PSOCK" will be used for windows and "FORK" otherwise. The value is passed as the parameter "type" to the function makeCluster.

Author(s)
Leandro Roser <learoser@gmail.com>

References
See Also

rkt.

Examples

```r
## Not run:
require("raster")
set.seed(6)

temp <- list()
for(i in 1:100) {
  temp[[i]] <- runif(36, -1, 1)
  temp[[i]] <- matrix(temp[[i]], 6, 6)
  temp[[i]] <- raster(temp[[i]])
}
temp <- brick(temp)

writeRaster(temp, "temporal.tif", overwrite=T)
rm(temp)
ndvisim <- brick("temporal.tif")

date <- seq(from = 1990.1, length.out = 100, by = 0.2)

# Parallel evaluation ----
eco.theilsen(ndvisim, date)

slope <- raster("slope.tif")
pvalue <- raster("pvalue.tif")

par(mfrow = c(1, 2))
plot(slope, main = "slope")
plot(pvalue, main = "p-value")

file.remove(c("slope.tif", "pvalue.tif"))

# Serial evaluation ----
eco.theilsen(ndvisim, date)

slope <- raster("slope.tif")
pvalue <- raster("pvalue.tif")

par(mfrow = c(1, 2))
plot(slope, main = "slope")
plot(pvalue, main = "p-value")
file.remove(c("temporal.tif", "slope.tif", "pvalue.tif"))
```
Description

This method unlocks the rows in an ecogen object. This means that different data frames in the object can have different rows, with different row names.

Usage

```r
## S4 method for signature 'ecogen'
eco.unlock(object)
```

Arguments

- `object` object of class ecogen

Examples

```r
## Not run:
data(eco.test)
eco2 <- eco.unlock(eco)
is.locked(eco2)
eco3 <- eco.lock(eco2)
is.locked(eco3)
## End(Not run)
```

Description

This method unlocks the rows in an ecogen object. This means that different data frames in the object can have different rows, with different row names.

Usage

```r
## S4 method for signature 'ecopop'
eco.unlock(object)
```
Arguments

object ecopop object

Examples

```r
## Not run:
data(eco.test)
my_ecopop2 <- eco.unlock(my_ecopop)
is.locked(my_ecopop2)
my_ecopop3 <- eco.lock(my_ecopop)
is.locked(my_ecopop3)
## End(Not run)
```

eco.v variogram

**Empirical variogram**

Description

This program computes the empirical variogram of a selected variable. If the coordinates are in decimal degrees, set latlon = TRUE. The program return a table with the mean class distances (d.mean) and the semivariances (obs) for each class.

Usage

```r
eco.v variogram(Z, XY, int = NULL, smin = 0, smax = NULL, nclass = NULL,
seqvec = NULL, size = NULL, bin = c("sturges", "FD"), row.sd = FALSE,
latlon = FALSE, angle = NULL)
```

Arguments

- **Z**: Vector for the analysis.
- **XY**: Data frame or matrix with the position of individuals (projected coordinates).
- **int**: Distance interval in the units of XY.
- **smin**: Minimum class distance in the units of XY.
- **smax**: Maximum class distance in the units of XY.
- **nclass**: Number of classes.
- **seqvec**: Vector with breaks in the units of XY.
- **size**: Number of individuals per class.
- **bin**: Rule for constructing intervals when a partition parameter (int, nclass or size) is not given. Default is Sturge’s rule (Sturges, 1926). Other option is Freedman-Diaconis method (Freedman and Diaconis, 1981).
- **row.sd**: Logical. Should be row standardized the matrix? Default FALSE (binary weights).
latlon Are the coordinates in decimal degrees format? Default FALSE. If TRUE, the
coordinates must be in a matrix/data frame with the longitude in the first column
and latitude in the second. The position is projected onto a plane in meters with
the function geoXY.

angle Direction for computation of a bearing variogram (angle between 0 and 180).
Default NULL (omnidirectional).

Value

The program returns an object of class "eco.correlog" with the following slots:
> OUT analysis output
> IN analysis input data
> BEAKS breaks
> CARDINAL number of elements in each class
> DISTMETHOD method used in the construction of breaks

ACCESS TO THE SLOTS The content of the slots can be accessed with the corresponding acces-
sors, using the generic notation of EcoGenetics (<ecoslot.> + <name of the slot> + <name of the
object>). See help("EcoGenetics accessories") and the Examples section below

Author(s)

Leandro Roser <learoser@gmail.com>

References

Business Media.
Amsterdam, Netherlands.
autocorrelation. Geographical Analysis, 32: 267-278.

Examples

# Not run:

data(eco.test)
variog <- eco.variogram(Z = eco[["P"]][, 2], XY = eco[["XY"]])
eco.plotCorrelog(variog)

# variogram plots support the use of ggplot2 syntax
require(ggplot2)
variogplot <- eco.plotCorrelog(variog) + theme_bw() + theme(legend.position="none")
variogplot

#------------------------
# ACCESSORS USE EXAMPLE
#------------------------
# the slots are accessed with the generic format
# (ecoslot. + name of the slot + name of the object).
# See help("EcoGenetics accessors")

ecoslot.OUT(variog)       # slot OUT
ecoslot.BREAKS(variog)    # slot BREAKS

## End(Not run)

---

**eco.weight**  

**Spatial weights**

---

### Description

Spatial weights for individuals (nodes) with coordinates XY

### Usage

```r
eco.weight(XY, method = c("circle", "knearest", "inverse", "circle.inverse", "exponential", "circle.exponential"), W = NULL, d1 = 0, d2 = NULL, k = NULL, p = 1, alpha = 1, dist.method = "euclidean", row.sd = FALSE, max.sd = FALSE, self = FALSE, latlon = FALSE, ties = c("unique", "min", "random", "ring", "first"))
```

### Arguments

- **XY**  
  Matrix/data frame with projected coordinates.

- **method**  
  Method of spatial weight matrix: "circle", "knearest", "inverse", "circle.inverse", "exponential", "circle.exponential".

- **W**  
  Custom weight matrix, with rownames and colnames identical to the XY data frame with coordinates

- **d1**  
  Minimum distance for circle matrices.

- **d2**  
  Maximum distance for circle matrices.

- **k**  
  Number of neighbors for nearest neighbor distance. When equidistant neighbors are present, the program select them randomly.

- **p**  

- **alpha**  
  Alpha value for exponential distance. Default = 1.

- **dist.method**  
  Method used for computing distances passed to `dist`. Default = euclidean.

- **row.sd**  
  Logical. Should be row standardized the matrix? Default FALSE (binary weights).

- **max.sd**  
  Logical. Should be divided each weight by the maximum of the matrix? Default FALSE (binary weights).
self

Should be the individuals self-included in circle or knearest weights? Default FALSE.

latlon

Are the coordinates in decimal degrees format? Default FALSE. If TRUE, the coordinates must be in a matrix/data frame with the longitude in the first column and latitude in the second. The position is projected onto a plane in meters with the function geoXY.

ties

ties handling method for "knearest" method: "unique" (default) for counting the ties as an unique neighbor (i.e), "min" for counting all the ties in a given category but each is counted as a neighbor, "random" for choosing at random a neighbor, "ring" for ring of neighbors, "first" for sequential k values for each neighbor.

Details

This program computes a weights matrix (square matrix with individuals in rows and columns, and weights wij in cells (i and j, individuals)) under the following available methodologies:

- circle: all the connection between individuals i and j, included in a distance radius, higher than d1 and lower than d2, with center in the individual i, have a value of 1 for binary weights. This distance requires the parameters d1 and d2 (default d1 = 0).

- knearest: the connections between an individual and its nearest neighbors of each individual i have a value of 1 for binary weights. This distance requires the parameter k.

- inverse: inverse distance with exponent p (distance = 1/dij^p, with dij the distance between individuals i and j). This distance requires the parameter p (default p = 1).

- circle inverse: combination of "circle" and "inverse". It is the matrix obtained by multiplying each element in a "circle" binary matrix, and an "inverse" matrix. This distance requires the parameters p, d1 and d2 (default p = 1, d1 = 0).

- exponential: inverse exponential distance with parameter alpha (distance = 1/e^(alpha *dij), with dij the distance between individuals i and j). This distance requires the parameter alpha (default alpha = 1).

- circle exponential: combination of "circle" and "exponential". It is the matrix obtained by multiplying each element in a "circle" binary matrix, and an "exponential" matrix. This distance requires the parameters alpha, d1 and d2 (default alpha = 1, d1 = 0).

In addition to these methods, a spatial weight object can be created assigning a custom W matrix ("W" argument). In this case, the "method" is argument automatically set by the program to "custom" (see te example).

In row standardization, each weight wij for the individual i, is divided by the sum of the row weights (i.e., wij / sum(wij), where sum(wij) is computed over an individual i and all individuals j).

When self is TRUE, the connection j = i is also included.

PLOTS FOR ECO.WEIGHT OBJECTS:

A plot method is available (function "eco.plotWeight") showing static or interactive plots, In the case of using the function eco.plotWeight for the argument type="simple", the connections are shown in two plots: an X-Y graph, with the individuals as points, representing the original coordinates, and in a plot with coordinates transformed as ranks (i.e., each coordinate takes an ordered value from
1 to the number of individuals). The other static method (type="igraph") uses the igraph package to generate a visual attractive graph (force network). Two interactive methods are available: type = "network", to plot an interactive force network, and type = "edgebundle" to plot a circular network. For the cases type = "inverse" or type = "exponential", the program generates a plot of weights values vs distance See the examples below.

Value

An object of class eco.weight with the following slots:

> W weights matrix
> XY input coordinates
> METHOD weights construction method
> PAR parameters used for the construction of weights
> PAR.VAL values of the parameters used for the construction of weights
> ROW.SD row standardization (logical)
> SELF data self-included (logical)
> NONZERO percentage of non-zero connections
> NONZEROIND percentage of individuals with non-zero connections
> AVERAGE average number of connection per individual

ACCESS TO THE SLOTS The content of the slots can be accessed with the corresponding accessor, using the generic notation of EcoGenetics (<ecoslot.> + <name of the slot> + <name of the object>). See help("EcoGenetics accessors") and the Examples section below

Author(s)

Leandro Roser <learoser@gmail.com>

Examples

```r
## Not run:

data(eco3)

# 1) "circle" method
con <- eco.weight(eco3[['XY']], method = "circle", d1 = 0, d2 = 500)

#---- Different plot styles for the graph ----#

# simple
eco.plotWeight(con, type = "simple")

# igraph
eco.plotWeight(con, type = "igraph", group = eco3[['S']])$structure)
```
# network (interactive)
## click in a node to see the label
eco.plotWeight(con, type = "network", bounded = TRUE, group = eco3[["S"]]

# edgebundle (interactive)
## in the following plot, the assignment a group factor,
## generates clustered nodes.
## hover over the nodes to see the individual connections
eco.plotWeight(con, type = "edgebundle", fontSize=8, group = eco3[["S"]]

# 2) "knearest" method
con <- eco.weight(eco3[["XY"]], method = "knearest", k = 10)
eco.plotWeight(con)
eco.plotWeight(con, type = "network", bounded = TRUE, group = eco3[["S"]]

# 3) "inverse" method
## scale dependent. In the example, the original coordinates (in km) are converted into m
con <- eco.weight(eco3[["XY"]]/1000, method = "inverse", max.sd = TRUE, p = 0.1)
con
eco.plotWeight(con)

# 4) "circle.inverse" method
con <- eco.weight(eco3[["XY"]], method = "circle.inverse", d2 = 1000)
con
eco.plotWeight(con)

# 5) "exponential" method
## scale dependent. In the example, the original coordinates (in km) are converted into m
con <- eco.weight(eco3[["XY"]]/1000, method = "exponential", max.sd = TRUE, alpha = 0.1)
eco.plotWeight(con)

# 6) "circle.exponential" method
con <- eco.weight(eco3[["XY"]], method = "circle.exponential", d2 = 2000)
con
eco.plotWeight(con)

# 7) CUSTOM WEIGHT MATRIX
## An eco.weight object can be created with a custom W matrix. In this case,
## the rows and the columns of W (weight matrix) must have names,
## that must coincide (also in order) with the name of the XY (position) matrix.

require(igraph)
## this example generates a network with the package igraph
tr <- make_tree(40, children = 3, mode = "undirected")
plot(tr, vertex.size = 10, vertex.label = NA)

## conversion from igraph to weight matrix
weights <- as.matrix(as_adj(tr))
## weight matrix requires named rows and columns
myNames <- 1:nrow(weights)
rownames(weights) <- colnames(weights) <- myNames

## extract coordinates from the igraph object
coord <- layout.auto(tr)
rownames(coord) <- myNames
plot(layout.auto(tr))

## custom weight object
customw <- eco.weight(XY = coord, W = weights)

## simple plot of the object
eco.plotWeight(customw, type = "simple")

## create a vector with groups to have coloured plots
myColors <- c(rep(1, 13), rep(2, 9), rep(3, 9), rep(4, 9))
eco.plotWeight(customw, type = "igraph", group = myColors)

## in the following plot, the argument bounded is set to FALSE,
## but if you have many groups, it probably should be set to TRUE.
# click in a node to see the label
eco.plotWeight(customw, type = "network", bounded = FALSE, group = myColors)

## in the following plot, the assignment a group factor,
# generates clustered nodes.
# hover over the name of the nodes to see the individual connections
eco.plotWeight(customw, type = "edgebundle", group = myColors)

#### CONVERSION FROM LISTW OBJECTS ####
require(adeegenet)
# Delaunay triangulation
temp <- chooseCN(eco3["XY"], type = 1, result.type = "listw", plot.nb = FALSE)
con <- eco.listw2ew(temp)
eco.plotWeight(con, "network", bounded = TRUE, group = eco3["S"]$structure)

# ACCESSORS USE EXAMPLE
# the slots are accessed with the generic format
# (ecoslot. + name of the slot + name of the object).
# See help("EcoGenetics accessors")
ecoslot.METHOD(con)  # slot METHOD
ecoslot.PAR(con)     # slot PAR
ecoslot.PAR.VAL(con) # slot PAR.VAL

## End(Not run)
Description

ekogen object with simulated data of 900 individuals.

Usage

data(eco2)
eco2

Author(s)

Leandro Roser <learoser@gmail.com>

Description

ekogen object with simulated data of 173 individuals.

Usage

data(eco3)
eco3

Author(s)

Leandro Roser <learoser@gmail.com>
Description

data frames with simulated data of 173 individuals. The data frames can be used to construct an ecogen object that includes genetic data separated by a character ad with non uniform of number of characters

Usage

data(eco3)
eco3

Author(s)

Leandro Roser <learoser@gmail.com>

ecogen

Creating a new ecogen object

Description

Creating a new ecogen object

Usage

ecogen(XY = data.frame(), P = data.frame(), G = data.frame(),
E = data.frame(), S = data.frame(), C = data.frame(),
G.processed = TRUE, order.G = FALSE, ploidy = 2,
type = c("codominant", "dominant"), sep = "", ncod = NULL,
missing = c("NA", "0", "MEAN"), NA.char = "NA", poly.level = 5,
rm.empty.ind = FALSE, order.df = TRUE, set.names = NULL,
valid.names = FALSE, lock.rows = TRUE)

Arguments

XY Data frame with m columns (coordinates) and n rows (individuals).
P Data frame with n rows (individuals), and phenotypic data in columns.
G Data of class: "data.frame", with individuals in rows and genotypic data in columns (loci). The ploidy and the type (codominant, dominant) of the data, must be passed with the arguments "ploidy" and "type". Missing data is coded as NA. Dominant data must be coded with binary values (0 for absence - 1 for presence).
E Data frame with n rows (individuals), and environmental data in columns.
S Data frame with n rows (individuals), and groups (factors) in columns. The program converts non-factor data into factor.

C Data frame with n rows (individuals), and custom variables in columns.

G.processed If TRUE, the slot G will include a processed data frame: removed non informative loci (the data non available for all the individuals), or non polymorphic loci (for dominant data).

order.G Genotypes must be ordered in G slot? (codominant data) Default FALSE. If true alleles are ordered in ascending order.


type Marker type: "codominant" or "dominant".

sep Character separating alleles (codominant data). Default option is no character separating alleles.

ncod Number of characters coding each allele (codominant data).

missing Missing data treatment ("NA", "0", or "MEAN") for the A slot. Missing elements are set to NA in the default option. missing elements are recoded as 0 or the mean allelic frequency across individuals in "0" and "MEAN" options, respectively.

NA.char Character symbolizing missing data in the input. Default is "NA".

poly.level Polymorphism threshold in percentage (0 - 100), for remotion of non polymorphic loci (for dominant data). Default is 5 (5%).

rm.empty.ind Remotion of noninformative individuals (row of "NAs"). Default if FALSE. This option is only available when the 'lock.rows' parameter is FALSE.

order.df Order individuals of data frames by row? (all data frames with a same order in row names). This option is only available when the 'lock.rows' parameter is TRUE. If the names of the data frames are not used (i.e., set.names and valid.names are not NULL), setting this parameter to TRUE/FALSE has no effect in the function. Default TRUE. If FALSE, the row names of all the data frames must be ordered. The use of data frames with row names in different order will return an error. In both cases, the program sets an internal names attribute of the object using the row names of the first non-empty data frame found in the following order: XY, P, G, E, S, C. This attribute is used as reference to order rows when order.df = TRUE.

set.names Character vector with names for the rows of the non-empty data frames. This argument is incompatible with valid.names

valid.names Logical. Create valid row names? This argument is incompatible with set.names. The program will name individuals with valid tags I.1, I.2, etc.

lock.rows Turn on row names check. Data frames require identical individuals in rows. Default TRUE.

Details This is a generic function for creation of ecogen objects. In the default option, missing data should be coded as "NA", but any missing data character can be passed with the option NA.char. In all the
cases, the new object will have a slot G coding the missing data as NA. For dominant markers (0/1 coding), the slot A is unnecessary as it is treated by ecogen methods as a symbolic link to G.

ACCESS TO THE SLOTS. MODIFICATION OF ECOKEN OBJECTS

The content of the slots can be extracted with the corresponding accessors ecoslot.XY, ecoslot.P, ecoslot.G, ecoslot.A, ecoslot.E, ecoslot.C and ecoslot.OUT. Accessors can be also used to assign data to the slots. The correct use of ecogen objects requires the implementation of accessors, as they ensure the checking and pre-processing of the data. The use of accessors allows to modify or fill the slots of ecogen objects, without the need of creating a new object each time. See help("EcoGenetics accessors") for a detailed description and examples about ecogen accessors.

OTHER SLOT ACCESS METHODS FOR ECOKEN OBJECTS

The use of brackets is defined for ecogen objects:

- Single bracket: the single bracket ("[") is used to subset all the ecogen data frames (P, G, E, S, A and C) by row, at once. The notation for an object is eco[from:to], where eco is any ecogen object, and from: to is the row range. For example: eco[1:10], subsets the object eco from row 1 to row 10, for all the data frames at once.

- Double square brackets: the double square brackets are symbolic abbreviations of the accessors (i.e., it is a call to the corresponding accessor). The usage is: eco["X"], where X is a slot: eco["P"], eco["G"], eco["A"], eco["E"], eco["S"], eco["C"] and eco["OUT"]. Double square brackets can be used in get/set mode. See Examples below and in help("EcoGenetics accessors").

ABOUT THE CONSTRUCTION OF NEW ECOKEN OBJECTS

A new ecogen object can be constructed in two different ways. First, a new object can be created, incorporating all the information at once. Second, the data can be added in each slot, using the corresponding accessor / ". Accessor/double square brackets methods allow temporal modification of any ecogen object and ensure the modularity of this kind of object. These methods are not only functions used to get/assign values to the slots, they provide a basic pre-processing of the data during assignment, generating a coherent and valid set of information.

LOCKED AND UNLOCKED OBJECTS

Starting from version 1.2.1.5, ecogen and ecopop objects can be "locked" (default) or "unlocked". A "locked" object must have identical number of rows and row names in all the input data frames (or a rule must be provided to construct the row names in case of ecogen objects, with valid.names or set.names arguments). An unlocked objects allows to have a free number of rows in each table, and row names do not need to coincide among tables. See examples below.

Author(s)

Leandro Roser <learoser@gmail.com>
Leandro Roser <learoser@gmail.com>

Examples

## Not run:

# Example with G data of class "data.frame", corresponding to # microsatellites of a diploid organism:
data(eco.test)
eco <- ecogen(XY = coordinates, P = phenotype, G = genotype, E = environment, S = structure)

# Example with G data of class "data.frame", corresponding to a presence - absence molecular marker:
dat <- sample(c(0,1),100,rep = TRUE)
dat <- data.frame(matrix(dat,10,10))
eco <- ecogen(G = dat, type = "dominant")

# DYNAMIC ASSIGNMENT WITH ACCESSORS AND "[["
eco <- ecogen(XY = coordinates, P = phenotype)
eco
ecoslot.G(eco, order.G = TRUE) <- genotype

# this is identical to
eco["G", order.G=TRUE] <- genotype
ecoslot.E(eco) <- environment

# this is identical to
eco["E"] <- environment

# See additional examples in help("EcoGenetics accessors")

# Storing data in the slot OUT
singers <- c("carlos_gardel", "billie_holiday")
ecoslot.OUT(eco) <- singers

# Storing several datasets

golden.number <- (sqrt(5) + 1) / 2
ecoslot.OUT(eco) <- list(singers, golden.number)  # several objects must be passed as a list

# this is identical to:
eco[["OUT"]] <- list(singers, golden.number)

# Locked and unlocked objects
is.locked(eco)  # check if object is locked
eco[["P"]]<- rbind(eco[["P"]], eco[["P"]])  # invalid

# in locked object
ecogen2ecopop

Conversion form ecogen to ecopop

Description
This function creates an ecopop object from an ecogen object.

Usage
ecogen2ecopop(from, hier, factor_to_counts = TRUE, aggregator = function(x) mean(x, na.rm = TRUE), allele_data = c("counts", "frequencies"))

Arguments
from Object of class "ecogen"
hier Name of the level of the slot S with hierarchies
factor_to_counts Convert factors into counts for each level?
aggregator Function used to aggregate data
allele_data Genetic data should be created as counts ("counts") or allele frequencies ("frequencies")?

Default is "counts".

Author(s)
Leandro Roser <learoser@gmail.com>

Examples

## Not run:

data(eco.test)
ecogen2ecopop(eco, hier = "pop")

## End(Not run)
Description

This function creates four data frames (XY.txt, NAMES.txt, P.txt, G.txt) in the indicated directory (default: working directory), which can be loaded in Geneland.

Usage

```r
ecogen2geneland(eco, dir = "", ncod = NULL, ploidy = 2,
    to_numeric = FALSE, nout = 3, recode = c("all", "column", "paired"),
    replace_in = NULL, replace_out = NULL, ...)
```

Arguments

- `eco`: Object of class "ecogen"
- `dir`: output path. Default = "" (current directory).
- `ncod`: Number of digits coding each allele (e.g., 1: x, 2: xx, 3: xxx, etc.).
- `ploidy`: Ploidy of the data.
- `to_numeric`: Recode the genetic data into numeric format? If TRUE, the function performs the correction via `eco.format`. Additional formatting parameters can be passed to this function.
- `nout`: Number of digits in the output when to_numeric = TRUE.
- `recode`: Recode mode when to_numeric = TRUE: "all" for recoding the data considering all the individuals values at once (e.g., protein data), "column" for recoding the values by column (e.g., microsatellite data), "paired" for passing the values of allelic states and corresponding replacement values, using the replace_in and replace_out arguments (e.g. replace_in = c("A", "T", "C", "G"), replace_out = c(1,2,3,4)).
- `replace_in`: vector with states of the data matrix to be replaced, when recode = "paired". This argument must be used in conjunction with the argument "replace_out".
- `replace_out`: vector with states of the data matrix used for replacement, when recode = "paired". This argument must be used in conjunction with the argument "replace_in".
- `...`: Additional parameters passed to `eco.format` when to_numeric = TRUE

Value

- `XY.txt`: Matrix with coordinates.
- `NAMES.txt`: Matrix with row names.
- `P.txt`: Matrix with phenotypic data.
- `G.txt`: Matrix with genotypic data.
Author(s)
Leandro Roser <learoser@gmail.com>

Examples

```r
## Not run:

data(eco.test)
ecogen2geneland(eco, dir = "", ncod=1)

## End(Not run)
```

---

ecogen2geneland  Exporting an ecogen genetic data frame into Genepop format

Description
This function converts the genetic data of an ecogen object into a Genepop input file.

Usage

```r
ecogen2geneland(eco, dir = "", outName = "infile.genepop.txt", grp = NULL, nout = 3, sep = "", recode = c("none", "all", "column", "paired"), replace_in = NULL, replace_out = NULL, ...)
```

Arguments

- `eco` Object of class "ecogen".
- `dir` output path. Default = "" (current directory).
- `outName` The name of the output file.
- `grp` The name of the S slot column with groups in which the sample must be divided (e.g., populations). If groups are not given (grp = NULL), all individuals will be assigned to a single one.
- `nout` Number of digits in the output file.
- `sep` Character separating alleles.
- `recode` Recode mode: "none" for no recoding (default), "all" for recoding the data considering all the individuals values at once (e.g., protein data), "column" for recoding the values by column (e.g., microsatellite data), "paired" for passing the values of allelic states and corresponding replacement values, using the replace_in and replace_out arguments (e.g. replace_in = c("A", "T", "C", "G"), replace_out = c(1,2,3,4)).
- `replace_in` vector with states of the data matrix to be replaced, when recode = "paired". This argument must be used in conjunction with the argument "replace_out".
Conversion from ecogen to genind and genind to ecogen

These functions export from ecogen to genind and viceversa

Usage

```
ecogen2genind(from)
genind2ecogen(from)
```

Arguments

- `from` Object of class "ecogen" / "genind"

Author(s)

Leandro Roser <learoser@gmail.com>
Examples

```r
## Not run:

data(eco.test)

# ecogen to genind
outGenind <- ecogen2genind(eco)
outGenind

# genind to ecogen
outEco <- genind2ecogen(outGenind)

## End(Not run)
```

---

**ecogen2gstudio**  
*Conversion from ecogen to gstudio and gstudio to ecogen*

**Description**

These functions convert the genetic data of an ecogen object in a gstudio data frame or vice versa.

**Conversion from gstudio to ecogen**

**Usage**

```r
ecogen2gstudio(from, type = c("codominant", "dominant"))
gstudio2ecogen(from, ID = "ID", lat = "Latitude", lon = "Longitude",
               struct = NULL)
```

**Arguments**

- **from**: Input object of class "ecogen" or "gstudio" (depending on the direction of conversion)
- **type**: The type of data: "codominant" (for codominant data); "dominant" for presence-absence data.
- **ID**: name of the column with ID (default "ID")
- **lat**: name of the column with latitude (default "Latitude")
- **lon**: name of the column with longitude (default "Longitude")
- **struct**: vector with name of the columns with structures (default NULL)

**Author(s)**

Leandro Roser <learoser@gmail.com>
### Examples

```r
## Not run:
data(eco.test)
togstudio <- ecogen2gstudio(eco, type = "codominant")
togstudio
toeco <- gstudio2ecogen(togstudio, ID = "ID", lat = "Latitude",
lon = "Longitude", struct = "pop")
toeco
# as ID, Latitude and Longitude are column names in the <togstudio> data frame
# (that match default parameter values for gstudio2ecogen),
# the latter is identical to this:
toeco <- gstudio2ecogen(togstudio, struct = "pop")
toeco
## End(Not run)
```

### Description

This function converts the genetic data of an ecogen object into a hierfstat data frame.

### Usage

```r
ecogen2hierfstat(eco, pop = NULL, to_numeric = FALSE, nout = 3,
recode = c("all", "column", "paired"), replace_in = NULL,
replace_out = NULL, ...)
```

### Arguments

- **eco**: Object of class "ecogen".
- **pop**: The name of the S slot column with the groups for the hierfstat data frame.
- **to_numeric**: Recode the genetic data into numeric format? If TRUE, the functions performs the correction via `eco.format`. Additional formatting parameters can be passed to this function.
- **nout**: Number of digits in the output when to_numeric = TRUE.
- **recode**: Recode mode when to_numeric = TRUE: "all" for recoding the data considering all the individuals values at once (e.g., protein data), "column" for recoding the values by column (e.g., microsatellite data), "paired" for passing the values of allelic states and corresponding replacement values, using the replace_in and replace_out arguments (e.g. replace_in = c("A", "T", "C", "G"), replace_out = c(1,2,3,4)).
ecogen2spagedi

Exporting an ecogen genetic data frame into SPAGeDi format

Description

This function converts the genetic data of an ecogen object in a SPAGeDi input file. When distance classes are required, they can be constructed by combining the parameters "int", "smin", "smax", "nclass", "seqvec" and "size", as described in the function eco.lagweight. A distance matrix can also be included using the "distmat" parameter. Missing data must be coded as a single "NA" in the G data frame.

Usage

ecogen2spagedi(eco, pop = NULL, ndig = NULL, dir = "", outName = "infile.spagedi.txt", smin = 0, smax = NULL, int = NULL, nclass = NULL, seqvec = NULL, size = NULL, bin = c("sturges", "FD"), distmat = NULL, latlon = FALSE, to_numeric = FALSE, nout = 3, recode = c("all", "column", "paired"), replace_in = NULL, replace_out = NULL, ...)

Author(s)

Leandro Roser <learoser@gmail.com>

Examples

## Not run:

data(eco.test)
hiereco <- ecogen2hierfstat(eco, "pop", to_numeric = TRUE)
require("hierfstat")
basic.stats(hiereco)

## End(Not run)
Arguments

- **eco**: Object of class "ecogen".
- **pop**: The name of the S slot column with the groups for the output data. The default option includes all the individuals into a single group.
- **ndig**: Number of digits coding each allele in the output file (e.g., 1: x, 2: xx, or 3: xxx). If NULL, the value will be deduced from the number of digits used for coding alleles in the ecogen object.
- **dir**: output path. Default = "" (current directory).
- **outName**: The name of the output file.
- **smin**: Minimum class distance in the units of the XY slot data.
- **smax**: Maximum class distance in the units of the XY slot data.
- **int**: Distance interval in the units of the XY slot data.
- **nclass**: Number of classes.
- **seqvec**: Vector with breaks in the units of the XY slot data.
- **size**: Number of individuals per class.
- **bin**: Rule for constructing intervals when a partition parameter (int, nclass or size) is not given. Default is Sturges’s rule (Sturges, 1926). Other option is Freedman-Diaconis method (Freedman and Diaconis, 1981).
- **distmat**: Distance matrix to include (optional).
- **latlon**: Are the coordinates in decimal degrees format? Default FALSE. If TRUE, the coordinates must be in a matrix/data frame with the longitude in the first column and latitude in the second. The position is projected onto a plane in meters with the function `geoxy`.
- **to_numeric**: Recode the genetic data into numeric format? If TRUE, the function performs the correction via `eco.format`. Additional formatting parameters can be passed to this function.
- **nout**: Number of digits in the output when to_numeric = TRUE.
- **recode**: Recode mode when to_numeric = TRUE: "all" for recoding the data considering all the individuals values at once (e.g., protein data), "column" for recoding the values by column (e.g., microsatellite data), "paired" for passing the values of allelic states and corresponding replacement values, using the `replace_in` and `replace_out` arguments (e.g. replace_in = c("A", "T", "C", "G"), replace_out = c(1,2,3,4)).
- **replace_in**: vector with states of the data matrix to be replaced, when recode = "paired". This argument must be used in conjunction with the argument "replace_out".
- **replace_out**: vector with states of the data matrix used for replacement, when recode = "paired". This argument must be used in conjunction with the argument "replace_in".
- **...**: Additional parameters passed to `eco.format` when to_numeric = TRUE

Author(s)

Leandro Roser <learoser@gmail.com>
References


Hardy O. and X Vekemans. 2002. SPAGeDi: a versatile computer program to analyse spatial genetic structure at the individual or population levels. Molecular ecology notes, 2: 18-620.


Examples

```r
## Not run:

data(eco.test)
ecogen2spagedi(eco, dir = "", pop = "pop", ndig = 1, int=2, smax=6, outName="infile.spagedi.txt")

## End(Not run)
```

---

**ecogenetics_devel**  
EcoGenetic devel site

**Description**

The function opens the EcoGenetics-devel web site: https://github.com/leandroroser/EcoGenetics-devel

**Usage**

```r
ecogenetics_devel()
```

---

**ecogenetics_tutorial**  
EcoGenetic tutorial site

**Description**

The function opens the EcoGenetics tutorial web site: https://leandroroser.github.io/EcoGenetics-Tutorial

**Usage**

```r
ecogenetics_tutorial()
```
ecopop

Creating a new ecopop object

Description

Creating a new ecopop object

Usage

ecopop(XY = data.frame(), P = data.frame(), AF = data.frame(),
E = data.frame(), S = data.frame(), C = data.frame(),
pop_names_column = 1L, ploidy, type = c("codominant", "dominant"),
order.df = FALSE, allele_data = c("counts", "frequencies"),
lock.rows = TRUE)

Arguments

XY       Data frame with n rows (populations) and m columns (coordinates).
P       Data frame with n rows (populations), and m columns (phenotypic variables).
AF       Data of class: "matrix", with n rows (populations) and m columns (allele counts).
The ploidy and the type (codominant, dominant) of the data, must be passed with the arguments "ploidy" and "type" for consistency with other methods of the package.
E       Data frame with n rows (populations), and n columns (environmental variables).
S       Vector (factor) with n items (population hierarchical levels).
C       Data frame with n rows (populations), and m columns (custom variables).
pop_names_column
Column with the population in the slot S that represents used to create the name of the object. Dafault is the first column.
ploidy    Ploidy of the AF data frame.
type     Marker type: "codominant" or "dominant".
order.df Order populations of data frames by row? (all data frames with a same row order). Default FALSE. The row names of all the data frames must be ordered. In this case, the use of data frames with row names in different order will return an error. In both cases, the program set the content of the S slots as the reference names of the object using the row names of the first non-empty data frame found in the following order: XY, P, AF, E, C. This attribute is used as reference to order rows when order.df = TRUE.
allele_data format for allele data output (slot AF). Can be "counts" or "frequencies".
lock.rows Turn on row names check. Data frames require indentical individuals in rows. Default TRUE.
Details

This is a generic function for creation of ecopop objects. Missing data should be coded as "NA".

ACCESS TO THE SLOTS. MODIFICATION OF ecopop OBJECTS

The content of the slots can be extracted with the corresponding accessors ecoslot.XY, ecoslot.P, ecoslot.AF, ecoslot.E and ecoslot.C. Accessors can be also used to assign data to the slots. The correct use of ecopop objects requires the implementation of accessors, as they ensure the checking and pre-processing of the data. The use of accessors allows to modify or fill the slots of ecopop objects, without the need of creating a new object each time. See help("EcoGenetics accessors") for a detailed description and examples about ecopop accessors.

OTHER SLOT ACCESS METHODS FOR ECOPOP OBJECTS

The use of brackets is defined for ecopop objects:

- Single bracket: the single bracket ("[") is used to subset all the ecopop data frames (P, G, E, S, AF and C) by row, at once. The notation for an object is eco[from:to], where eco is any ecopop object, and from: to is the row range. For example: my_ecopop[1:10], subsets the object my_ecopop from row 1 to row 10, for all the data frames at once.

- Double square brackets: the double square brackets are symbolic abbreviations of the accessors (i.e., it is a call to the corresponding accessor). The usage is: my_ecopop["X"], where X is a slot: my_ecopop["P"], my_ecopop["AF"], my_ecopop["E"], my_ecopop["S"] and my_ecopop["C"]). Double square brackets can be used in get/set mode. See Examples below and in help("EcoGenetics accessors").

ABOUT THE CONSTRUCTION OF NEW ECOPOP OBJECTS

In most cases, a new ecopop object is created from an ecogen object, using the function eco-gen2ecopop. A new ecopop object can also be directly constructed in two different ways. First, a new object can be created, incorporating all the information in one step with the constructor. Second, the data can be added to each slot, using the corresponding accessor or, in an equivalent way, with double brackets notation ("[").

LOCKED AND UNLOCKED OBJECTS # Starting from version 1.2.1.5, ecogen and ecopop objects can be "locked" (default) or "unlocked". A "locked" object must have identical number of rows and row names in all the input data frames (or a rule must be provided to construct the row names in case of ecogen objects, with valid.names or set.names arguments). An unlocked objects allows to have a free number of rows in each table, and row names do not need to coincide among tables. See examples below.

Author(s)

Leandro Roser <learoser@gmail.com>

Examples

```r
## Not run:
data(eco.test)

## Three ways to construct an ecopop object

## 1) ecogen to ecopop
```
my_ecopop <- ecogen2ecopop(eco, hier = "pop")

# Extracting tables with accessors (double brackets notation)
XY_pop <- my_ecopop["XY"]
P_pop <- my_ecopop["P"]
AF_pop <- my_ecopop["AF"]
E_pop <- my_ecopop["E"]
S_pop <- my_ecopop["S"]

## 2) Creating a new ecopop object
my_ecopop2 <- ecopop(XY = XY_pop, P = P_pop, AF = AF_pop, E = E_pop,
                      S = S_pop,
                      ploidy = 2, type = "codominant")

## 3) From an empty object
# new empty object
my_ecopop3 <- ecopop(ploidy = 2, type = "codominant")

set slots, using as example the data generated above
my_ecopop3["XY"] <- XY_pop # The first assignments initializes the S slot
my_ecopop3["P"] <- P_pop
my_ecopop3["AF", ploidy = 2] <- AF_pop
my_ecopop3["E"] <- E_pop
my_ecopop3["S"] <- S_pop

## Subsetting by rows
my_ecopop3[1:10]

# Locked and unlocked objects

is.locked(my_ecopop) # check if object is locked
my_ecopop["P"] <- rbind(my_ecopop["P"], my_ecopop["P"])) # invalid in locked object

my_ecopop_unlocked <- eco.unlock(my_ecopop) # unlocked object
my_ecopop_unlocked["P"] <- rbind(my_ecopop["P"], my_ecopop["P"])) # valid now

new_locked <- eco.lock(my_ecopop_unlocked) # invalid
my_ecopop_unlocked["P"] <- my_ecopop["P"]
new_locked <- eco.lock(my_ecopop_unlocked) # valid now

## End(Not run)
ecopop2genpop

**Conversion form ecopop to genpop and genpop to ecopop**

**Description**

These functions export from ecopop to genpop and vice versa.

```r
genpop2ecpop
```

**Usage**

```r
ecopop2genpop(from)
genpop2ecpop(from)
```

**Arguments**

- `from` Object of class "ecopop" / "genpop"

**Author(s)**

Leandro Roser <learoser@gmail.com>

**Examples**

```r
## Not run:
data(eco.test)
my_ecopop <- ecogen2ecopop(eco, hier = "pop")
my_genpop <- ecopop2genpop(my_ecopop)
my_ecopop2 <- genpop2ecopop(my_genpop)
## End(Not run)
```

desc_countsRaf

**Description**

Conversion from ecopop with genetic data as count, into ecopop with genetic data as allele frequencies.

**Usage**

```r
ecopop_counts2af(from)
```
eco_dom

Arguments
from ecopop object

Examples
## Not run:
data(eco.test)
ecopop_counts2af(my_ecopop)

## End(Not run)

data(eco.test)
eco

Description
ecogen object with simulated data of 225 individuals, with dominant markers

Usage
data(eco.test)
eco

Author(s)
Leandro Roser <learoser@gmail.com>

data(eco.test)
environment

Description
Data frame with simulated environmental variables of 225 individuals.

Usage
data(eco.test)
environment

Author(s)
Leandro Roser <learoser@gmail.com>
Description

data frame with simulated genetic data of 173 individuals.

Usage

data(eco4)
G

Author(s)

Leandro Roser <learoser@gmail.com>

Description

This function converts a Genepop file into an object with a genetic matrix (G) and a structures matrix (S).

Usage

genepop2ecogen(genefile = NULL)

Arguments

genefile Genepop file.

Value

A list with the objects G (genetic matrix) and S (structures matrix).

Author(s)

Leandro Roser <learoser@gmail.com>
**Examples**

```r
## Not run:
# ingpop, file with Genepop format in the folder "/extdata" of the package
ecopepath <- paste(path.package("EcoGenetics"), "/extdata/ingpop", sep = "")
ingpop <- genepop2ecogen(ecopath)
ingpop

## End(Not run)
```

---

**Description**

Data frame with simulated microsatellite data of 225 individuals.

**Usage**

```r
data(eco.test)
genotype
```

**Author(s)**

Leandro Roser <learoser@gmail.com>

---

**Description**

Data frame with simulated dominant data of 225 individuals.

**Usage**

```r
data(eco.test)
genotype_dom
```

**Author(s)**

Leandro Roser <learoser@gmail.com>
grf.multiplot 

Multiple plot function for ggplot

Description

Multiple plot function for ggplot

Usage

grf.multiplot(..., plotlist = NULL, cols = 1, layout = NULL)

Arguments

... ggplot objects
plotlist List of ggplot object
cols Number of columns in layout
layout A matrix specifying the layout. If present, 'cols' is ignored.

grf.seqmultiplot 

Plot a ggplot sequence in layers of n plots arranged in k rows

Description

Plot a ggplot sequence in layers of n plots arranged in k rows

Usage

grf.seqmultiplot(x, n, nrow, byrow = TRUE)

Arguments

x list of ggplot objects
n number of plot in layout
nrow Number of rows in layout
byrow plot by row?

Author(s)

Leandro Roser <learoser@gmail.com>
is.locked, ecogen-method

Test if rows of an ecogen object are locked

Description
Test if rows of an ecogen object are locked

Usage
```r
## S4 method for signature 'ecogen'
is.locked(object)
```

Arguments
- `object` ecogen object

Examples
```r
## Not run:
data(eco.test)
is.locked(eco)
eco2 <- eco.unlock(eco)
is.locked(eco2)
## End(Not run)
```

is.locked, ecopop-method

Test if rows of an ecopop object are locked

Description
Test if rows of an ecopop object are locked

Usage
```r
## S4 method for signature 'ecopop'
is.locked(object)
```

Arguments
- `object` ecopop object
Examples

```r
## Not run:
data(eco.test)
is.locked(my_ecopop)
eco2 <- eco.unlock(eco)
is.locked(eco2)

## End(Not run)
```

Description

ecopop object generated with the object eco

Usage

```r
data(eco.test)
my_ecopop
```

Author(s)

Leandro Roser <learoser@gmail.com>

---

Description

Factor with simulated phenotypic data of 173 individuals.

Usage

```r
data(eco4)
P
```

Author(s)

Leandro Roser <learoser@gmail.com>
phenotype

Description
Data frame with simulated morphometric data of 225 individuals.

Usage
```
data(eco.test)
phenotype
```

Author(s)
Leandro Roser <learoser@gmail.com>

plot.eco.multilsa,ANY-method

Description
Plot method for local spatial analysis

Usage
```
## S4 method for signature 'eco.multilsa,ANY'
plot(x)
```

Arguments
```
x    eco.multilsa object
```

Author(s)
Leandro Roser <learoser@gmail.com>

See Also
```
eco.lsa
```
Description

Factor with simulated groups of 173 individuals.

Usage

data(eco4)
S

Author(s)

Leandro Roser <learoser@gmail.com>

---

spagedi2ecogen

Importing a SPAGeDi file, via conversion to ecogen

Description

This function converts a SPAGeDi file into a ecogen object

Usage

spagedi2ecogen(infile, sep = "", missCode = NULL, type = c("codominant", "dominant"), ...)

Arguments

infile Path to the SPAGeDi file.
sep Character separating alleles (codominant data). Default option is no character separating alleles.
missCode characters to represent missing genotypes in codominant markers. If NULL, is computed as "0" times the number of characters coding alleles.
type Marker type: "codominant" or "dominant".
... additional arguments passed to ecogen

Author(s)

Leandro Roser <learoser@gmail.com>
# Examples

```r
## Not run:

data(eco.test)
ecogen2spagedi(eco, dir = "", pop = "pop", ndig = 1, int=2, smax=6, outName="infile.spagedi.txt")
spagedi2ecogen("infile.spagedi.txt", sep = "")

## End(Not run)
```

## Description

Factor with simulated groups of 225 individuals.

## Usage

```r
data(eco.test)
structure
```

## Author(s)

Leandro Roser <learoser@gmail.com>

## summary,eco.mlm-method

**Summary for eco.lmtree output**

## Description

Summary for eco.lmtree output

## Usage

```r
## S4 method for signature 'eco.mlm'
summary(object)

## S4 method for signature 'eco.mctree'
summary(object)
```

## Arguments

- **object**: Output object of `eco.lmtree`.  

---

*Note: The content is a natural representation of the English text from the document.*
Value

A table with a summary of the analysis for "mlm" analysis, the plot of the trees with significant splits for "mctree" analysis.

Author(s)

Leandro Roser <learoser@gmail.com>

See Also

eco.lmtree

eco.lmtree

Examples

## Not run:

data(eco.test)
# ' mod <- eco.lmtree(DF1 = eco$P, DF2 = eco$E, 
analysis = "mlm")
summary(mod)               #summary for "mlm" analysis

mod <- eco.lmtree(DF1 = eco$P, DF2 = eco$E, 
analysis = "mctree", fact = eco$S$structure)
summary(mod)               #summary for "mctree" analysis

## End(Not run)

Description

Data frame with information of bands 3 and 4 for two dates, corresponding to real Landsat 5 images and used in this package as pedagogic material complementing simulated data. Date and sun elevation data were extracted from the header provided with the image in http://glovis.usgs.gov/. The starting haze values (SHV) were estimated checking the profiles of the bands.

Usage

data(tab)
tab

Author(s)

Leandro Roser <learoser@gmail.com>
**Description**

Allelic frequency table from 50 villages, analyzed in Sokal et al. (1986).

**Usage**

data(sokal1986)
table.sokal

**Author(s)**

Leandro Roser<learoser@gmail.com>

---

<table>
<thead>
<tr>
<th>$XY$</th>
<th>$XY$</th>
</tr>
</thead>
</table>

**Description**

Factor with simulated coordinates of 173 individuals.

**Usage**

data(eco4)

XY

**Author(s)**

Leandro Roser<learoser@gmail.com>
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