Package ‘vegclust’

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This package contains functions used to perform fuzzy and hard clustering of vegetation data under different models.

This package is intended to be helpful for generating and maintaining classifications of vegetation. There are two main functions: (1) `vegclust` is used to create a fuzzy classification from an initial data set (a set of relevés of community composition); (2) `vegclass` is used to classify new data (i.e.
new relevés) into previous classification structures.

Author(s)

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References


Examples

```r
## Loads data
data(wetland)

## This equals the chord transformation
## (see also \code{\link{decostand}} in package 'vegan')
wetland.chord = as.data.frame(sweep(as.matrix(wetland), 1, sqrt(rowSums(as.matrix(wetland)^2)), "/"))

## Create noise clustering with 3 clusters. Perform 10 starts from random seeds
## and keep the best solution
wetland.nc = vegclust(wetland.chord, mobileCenters=3, m = 1.2, dnoise=0.75,
                      method="NC", nstart=10)
```

---

### as.memb

**Turns into membership matrix**

**Description**

Attempts to turn its cluster vector argument into a membership matrix
Usage

as.memb(cluster)

Arguments

cluster   A vector indicating the hard membership of each object in x to a set of groups. Can contain NA values.

Value

An matrix with as many rows as the length of cluster and as many columns as different cluster levels. NA values will have zero membership to all clusters.

Author(s)

Miquel De Cáceres, Forest Science Center of Catalonia.

See Also

vegclust, vegclass

Examples

as.memb(factor(c(1,2,NA)))

as.vegclust                  Turns into vegclust objects

Description

Attempts to turn its arguments into a vegclust object

Usage

as.vegclust(x, y, method="KM", m=1.0, dnoise=NULL, eta=NULL)

Arguments

x          A site-by-species data matrix (raw mode), or a site-by-site distance matrix (distance mode).

y          A vector indicating the cluster that each object in x belongs to. Alternatively, a fuzzy/hard site-by-group matrix of membership values.

method     A clustering model from which y was obtained (normally "KM"). Current accepted models are:

  • "KM": K-means or hard c-means (MacQueen 1967)
  • "KMdd": Hard c-medoids (Krishnapuram et al. 1999)
  • "FCM": Fuzzy c-means (Bezdek 1981)
• "FCMdd": Fuzzy c-medoids (Krishnapuram et al. 1999)
• "NC": Noise clustering (Dave and Krishnapuram 1997)
• "NCdd": Noise clustering with medoids
• "HNC": Hard noise clustering
• "HNCdd": Hard noise clustering with medoids
• "PCM": Possibilistic c-means (Krishnapuram and Keller 1993)
• "PCMdd": Possibilistic c-medoids

m
The fuzziness exponent to be used, relevant for all fuzzy models (FCM, FCMdd, NC, NCdd, PCM and PCMdd)
dnoise
The distance to the noise cluster, relevant for noise clustering models (NC, HNC, NCdd and HNCdd).
eta
A vector of reference distances, relevant for possibilistic models (PCM and PCMdd).

Details
This function is used to generate vegclust objects which can then be used in vegclass to classify new data. If the input classification is hard (i.e. yes/no membership), cluster centers are calculated as multivariate means, and the method for assigning new data is assumed to be k-means ("KM"), i.e. plots will be assigned to the nearest cluster center. If community data is given as site-by-species data matrix the cluster centroids are added as mobileCenters in the vegclust object. Centroids will not be computed if community data is given as a site-by-site dissimilarity matrix. Moreover, current implementation does not allow y to be a membership matrix when x is a distance matrix.

Value
An object of class vegclust.

Author(s)
Miquel De Cáceres, Forest Science Center of Catalonia.

See Also
vegclust, vegclass

Examples

```r
## Loads data
data(wetland)

## This equals the chord transformation
## (see also \code{\link{decostand}} in package vegan)
wetland.chord = as.data.frame(sweep(as.matrix(wetland), 1, sqrt(rowSums(as.matrix(wetland)^2)), "/"))

## Splits wetland data into two matrices of 30x27 and 11x22
wetland.30 = wetland.chord[1:30,]
wetland.30 = wetland.30[,colSums(wetland.30)>0]
```
dim(wetland.30)
wetland.11 = wetland.chord[31:41,]
wetland.11 = wetland.11[,colSums(wetland.11)>0]
dim(wetland.11)

## Performs a K-means clustering of the data set with 30 sites
wetland.km = kmeans(wetland.30, centers=3, nstart=10)

## Transforms the 'external' classification of 30 sites into a 'vegclust' object
wetland.30.vc = as.vegclust(wetland.30, wetland.km$cluster)

## Assigns the second set of sites according to the (k-means) membership rule
## That is, sites are assigned to the cluster whose cluster centroids is nearest.
wetland.11.km = vegclass(wetland.30.vc, wetland.11)

## A similar 'vegclust' object is obtained when using the distance mode...
wetland.d.vc = as.vegclust(dist(wetland.30), wetland.km$cluster)

## which can be also used to produce the assignment of the second set of objects
wetland.d.11 = as.data.frame(as.matrix(dist(wetland.chord))[31:41,1:30]
wetland.d.11.km = vegclass(wetland.d.vc,wetland.d.11)

CAP  Cumulative abundance profile (CAP)

Description

Functions to calculate cumulative abundance profiles (CAPs), to build matrices from them, and to summarize several profiles.

Usage

CAP(x, transform=NULL, verbose=FALSE)
CAP2matrix(CAP, type="cumulative",classWeights=NULL)
CAPcenters(CAP, y=NULL)
CAPquantile(CAP, q = 0.5, y = NULL)

Arguments

x  A stratified vegetation data set (see function stratifyvegdata).
transform  A function or the name of a function to be applied to each cumulative abundance value.
verbose  A logical flag to indicate extra output.
CAP  An object of class 'CAP'.
type  The type of information that the resulting matrix should contain. Either "profile", "abundance" or "volume".
classWeights  A numerical vector containing the weight for size class. If NULL, then all classes are assumed to have the same weight.
y A vector used as a factor to calculate average or quantile profiles per each level. Alternatively, an object of class `vegclust` for which CAP centroids or medoids are desired.

q Probability value for which the quantile is desired. By default the median is given.

Details

Function CAP replaces the abundance value of a size class by the sum of abundances in this and larger size classes (strata). Thus, upper classes contain smaller abundance values than lower classes, creating a cumulative abundance profile. Function CAP2matrix takes an object of class 'CAP' and returns a data matrix, where values differ depending on parameter type: (1) type="cumulative" simply reshapes the 'CAP' object (a list) into a matrix with as many rows as plot records and where columns are organized in blocks (there are as many blocks as species and each block has as many columns as size classes); (2) type="total" returns a plot-by-species matrix where each value is the total abundance of the species in the plot (i.e. the CAP value at the ground level); (3) type="volume" returns a plot-by-species matrix where each value is the sum of CAP values across size classes (a measure of the "volume" occupied by the species in the plot). When provided, `classWeights` are used to weight size classes of the cumulative abundance profiles (for (1) and (3) only). Function CAPcenters calculates the average abundance profile for a set of plot records. If `y` is a factor, it is used to specify groups of samples for which average profiles are to be calculated. If `y` is an object of class 'vegclust' then the function returns the CAP centroids or medoids corresponding to the clustering result. Function CAPquantile calculates a quantile profile for a set of CAPs. The usage of `y` is the same as for CAPcenters.

Value

Function CAP returns an object of class 'CAP', similar to objects of class 'stratifiedvegdata' but where abundance values of upper size classes have been added to those of lower size classes. Function CAP2matrix returns a matrix with species as rows (columns depend on the value of type). Functions CAPcenters and CAPquantile return an object of class 'CAP'.

Author(s)

Miquel De Cáceres, Forest Science Center of Catalonia.

References


De Cáceres, M., Coll, L., Martín-Alcón, S., González-Olabarria, J.R. (submitted) A general method for the classification of forest stands using structure and composition.

See Also

`stratifyvegdata, plot.CAP, vegdiststruct`
Examples

```r
## Load stratified data
data(medreg)

## Check that 'medreg' has correct class
class(medreg)

## Look at the data for the third plot
medreg[[3]]

## Create cumulative abundance profile (CAP) for each plot
medreg.CAP = CAP(medreg)

## Look at the profile of the third plot
medreg.CAP[[3]]

## Create matrix with species abundances
medreg.X = CAP2matrix(medreg.CAP, type="total")
head(medreg.X)

## Generate and plot average profile
average.CAP = CAPcenters(medreg.CAP)
plot(average.CAP)

## Generate and plot median profile
median.CAP = CAPquantile(medreg.CAP, q = 0.5)
plot(median.CAP)
```

---

**Cumulative abundance surface (CAS)**

**Description**

Functions to calculate cumulative abundance surfaces (CASs), to build matrices from them, and to summarize several CASs.

**Usage**

```r
CAS(x, transform=NULL, verbose=FALSE)
CASmargin(CAS, margin=1, verbose=FALSE)
CAS2matrix(CAS, type="cumulative", classWeights=NULL)
CAScenters(CAS, y=NULL)
CASquantile(CAS, q = 0.5, y = NULL)
```

**Arguments**

- `x` An object of class 'doublestratifiedvegdata' (see function `stratifyvegdata`).
transform A function or the name of a function to be applied to each cumulative abundance value.
verbose A logical flag to indicate extra output.
CAS An object of class 'CAS'.
margin Indicates whether marginalization should be done in primary (margin = 1) or secondary (margin = 2) size classes.
type The type of information that the resulting matrix should contain (either "cumulative" or "total").
classWeights A numerical matrix containing the weight for each combination of size classes. If NULL, then all classes are assumed to have the same weight.
y A vector used as a factor to calculate average or quantile surfaces per each level. Alternatively, an object of class vegclust for which CAS centroids or medoids are desired.
q Probability value for which the quantile is desired. By default the median is given.

Details

Function CAS replaces the abundance value of each combination of size classes by the sum of abundances in this and larger size classes. This creates a cumulative abundance surface (similar to a bivariate cumulative distribution function). Function CASmargin takes an object of class 'CAS' and returns an object of class 'CAP' that corresponds marginal profile in either the primary or the secondary size classes. Function CAS2matrix takes an object of class 'CAS' and returns a data matrix, where values differ depending on parameter type: (1) type="cumulative" simply reshapes the 'CAS' object (a list) into a matrix with as many rows as plot records and where columns are organized in blocks (there are as many blocks as species and each block has as many columns as combinations of size classes); (2) type="total" returns a plot-by-species matrix where each value is the total abundance of the species in the plot (i.e. the CAS value at the ground level). When provided, classWeights are used to weight size classes of the cumulative abundance surfaces (for (1) only). Function CAScenters calculates the average abundance surface for a set of plot records. If y is a factor, it is used to specify groups of samples for which average profiles are to be calculated. If y is an object of class 'vegclust' then the function returns the CAS centroids or medoids corresponding to the clustering result. Function CASquantile calculates a quantile surface for a set of CASs. The usage of y is the same as for CAScenters.

Value

Function CAS returns an object of class 'CAS', similar to objects of class 'doublestratifiedvegdata' but where abundance values of upper size classes have been added to those of lower size classes. Function CAS2matrix returns a matrix with species as rows (columns depend on the value of type). Functions CAScenters and CASquantile return an object of class 'CAS'.

Author(s)
Miquel De Cáceres, Forest Science Center of Catalonia.
References


De Cáceres, M., Coll, L., Martín-Alcón, S., González-Olabarria, J.R. (submitted) A general method for the classification of forest stands using structure and composition.

See Also

stratifyvegdata, plot.CAS, vegdiststruct

Examples

```r
## Load tree data
data(treeData)

## Define stratum thresholds (4 strata)
heights = seq(0, 4, by=0.5)
diameters = seq(0, 2, by=0.5)

## Stratify tree data using heights and diameters as structural variables
X = stratifyvegdata(treeData, sizes1=heights, sizes2=diameters, plotColumn="plotID",
                   speciesColumn="species", size1Column="height", size2Column="diam",
                   counts=TRUE)

## Build cumulative abundance surface
Y = CAS(X)

## Extracts the first and second marginal (i.e. CAP on heights or diameters respectively)
Y.M1 = CASmargin(Y, margin = 1)
Y.M1[[2]]

Y.M2 = CASmargin(Y, margin = 2)
Y.M2[[2]]

## For comparison we calculate the same profiles using the stratifyvegdata and CAP functions
Y1 = CAP(stratifyvegdata(treeData, sizes1=heights, plotColumn="plotID",
                         speciesColumn="species", size1Column="height",
                         counts=TRUE))

Y1[[2]]

Y2 = CAP(stratifyvegdata(treeData, sizes1=diameters, plotColumn="plotID",
                         speciesColumn="species", size1Column="diam",
                         counts=TRUE))

Y2[[2]]

## Compare Y.M1[[2]] with Y1[[2]] and Y.M2[[2]] with Y2[[2]]
```
clustcentroid

Cluster centers of a classification

Description

Function clustcentroid calculates the centroid (multivariate average) coordinates of a classification. Function clustmedoid determines the medoid (object whose average dissimilarity to all the other objects is minimal) for each cluster in the classification.

Usage

clustcentroid(x, y, m = 1)
clustmedoid(x, y, m = 1)

Arguments

x Community data, a site-by-species data frame. In function clustmedoid, x can alternatively be an object of class dist (otherwise, the dissimilarity measure is assumed to be the Euclidean distance).

y It can be (a) A vector indicating the cluster that each object in x belongs to; (b) a fuzzy/hard site-by-group matrix of membership values; (c) an object of class vegclust or vegclass

m Fuzziness exponent, only effective when y is a fuzzy membership matrix.

Value

Function clustcentroid returns a group-by-species matrix containing species average abundance values (i.e. the coordinates of each cluster centroid). Function clustmedoid returns a vector of indices (medoids).

Note

In order to assign new plot record data into a predefined set of classes, one should use functions as.vegclust and vegclass instead.

Author(s)

Miquel De Cáceres, Forest Science Center of Catalonia

See Also

as.vegclust, vegclass, vegclust, kmeans
Examples

## Loads stats
library(stats)

## Loads data
data(wetland)

## This equals the chord transformation
## (see also \code{\link{decostand}} in package 'vegan')
wetland.chord = as.data.frame(sweep(as.matrix(wetland), 1,
    sqrt(rowSums(as.matrix(wetland)^2)), "/"))

## Performs a K-means clustering
wetland.km = kmeans(wetland.chord, centers=3, nstart=10)

## Gets the coordinates corresponding to the centroids of KM clusters
clustcentroid(wetland.chord, y=wetland.km$cluster)

## Gets the object indices corresponding to the medoids of KM clusters
clustmedoid(wetland.chord, y=wetland.km$cluster)

---

clustconst  

\textit{Constancy table of a classification}

Description

Allows studying the constancy table (i.e. the frequency of species in each class) of a classification represented in the form of a membership data matrix.

Usage

clustconst(x, memb)

## S3 method for class 'clustconst'
summary(object, mode="all", name=NULL, sort=TRUE, minconst=0.5, digits=3, ...)

Arguments

- \texttt{x} \hspace{1cm} Community data, a site by species data frame.
- \texttt{memb} \hspace{1cm} An site-by-group matrix indicating the (hard or fuzzy) membership of each object in \texttt{x} to a set of groups.
- \texttt{object} \hspace{1cm} An object of class 'clustconst'.
- \texttt{mode} \hspace{1cm} Use \texttt{mode="all"} to print the constancy table, \texttt{mode="cluster"} to print constancy values for one cluster, and \texttt{mode="species"}, to print constancy values for one species.
name
sort
minconst
digits
...
clustvar

Description

Computes the variation in community composition (i.e., beta diversity) found within the sites of a set of hard or fuzzy clusters.

Usage

clustvar(x, cluster = NULL, defuzzify=FALSE,...)

Arguments

x Community data. Either a site-by-species matrix or a site-by-site matrix of compositional distances between sites (i.e., an object of class `dist`). Alternatively, this can be an object of class `vegclust` or `vegclass`, and in this case it is unnecessary to provide `cluster`.

cluster A vector indicating the hard membership of each object in `x` to a set of groups.

defuzzify A flag indicating whether fuzzy memberships should be defuzzified (see function `defuzzify`). Only applies to the case where an object of class `vegclust` or `vegclass` is supplied in `x`.

... Additional parameters for function `defuzzify`.

Details

This function can be used in two ways:

- if `x` is a data matrix (site by species or distances among sites) and `cluster` is null, the function assumes a single cluster of all points in `x`. When `cluster` is provided, the function computes cluster variance for each (hard) group and this computation implies setting the centroid of the group. Cluster variance is defined as the average squared distance to the centroid.

- If `x` is an object of class `vegclust` or `vegclass`, the function uses the information contained there (distances to cluster centers, memberships and exponent of fuzziness) in order to compute cluster variances. Cluster centers do not need to be recomputed, and the distances to cluster centers are used directly. For centroid-based cluster models (KM, FCM, NC, HNC and PCM) the variance is defined as the average squared distance to the centroid. For medoid-based cluster models (KMdd, FCMdd, NCdd, HNCdd and PCMdd) the variance is defined as average distance to the medoid. The variance for both mobile and fixed clusters is returned. Additionally, membership matrices may be defuzzified if `defuzzify=TRUE`. 

```r
d=summary(c, mode="all")

## Prints the most frequent species in the first cluster
summary(c, mode="cluster ", name=names(c)[1])
```
**Value**

A double value (for one cluster) or a vector of values, one per each cluster.

**Author(s)**

Miquel De Cáceres, Forest Science Center of Catalonia

**See Also**

`vegclust`, `kmeans`, `defuzzify`

**Examples**

```r
## Loads data
data(wetland)

## This equals the chord transformation
## (see also \code{\link{decostand}} in package 'vegan')
wetland.chord = as.data.frame(sweep(as.matrix(wetland), 1,
                                sqrt(rowSums(as.matrix(wetland)^2)), "/")

## Create noise clustering with 3 clusters. Perform 10 starts from random seeds
## and keep the best solution
wetland.nc = vegclust(wetland.chord, mobileCenters=3, m = 1.2, dnoise=0.75,
                       method="NC", nstart=10)

## Gets cluster variance of fuzzy clusters
clustvar(wetland.nc)

## Gets cluster variance of fuzzy clusters after defuzzification
clustvar(wetland.nc, defuzzify=TRUE)

## Similar to the previous, this gets cluster variance of defuzzified (i.e. hard) clusters
clustvar(wetland.chord, cluster=defuzzify(wetland.nc)$cluster)

## Gets cluster variance of K-means (i.e. hard) clusters
clustvar(wetland.chord, cluster=kmeans(wetland.chord, centers=3, nstart=10)$cluster)
```

---

**Description**

Computes an index to compare two classifications.

**Usage**

`concordance(x, y, method="adjustedRand",...)`
Arguments

\(x, y\) Classification vector or membership matrix. Alternatively, objects of type \texttt{vegclust} or \texttt{vegclass}.

\texttt{method} A string vector to indicate the desired indices (see details).

\ldots Additional parameters for function \texttt{defuzzify}, which will be called if \(x\) or \(y\) are of type \texttt{matrix}, \texttt{vegclust} or \texttt{vegclass}.

Details

Several indices for comparison of partitions are available:

- \texttt{method="Rand"}: Rand (1971) index.
- \texttt{method="adjustedRand"}: Rand index adjusted for random effects (Hubert & Arabie 1985).
- \texttt{method="Wallace"}: Wallace (1983) index (for asymmetrical comparisons). This index (and its adjusted version) is useful to quantify how much \(x\) is nested into \(y\).
- \texttt{method="adjustedWallace"}: Wallace index adjusted for random effects (Pinto et al. 2008).

Value

A numeric vector with the desired index values.

Author(s)

Miquel De Cáceres, Forest Science Center of Catalonia

References


See Also

\texttt{vegclust}, \texttt{vegclass}, \texttt{defuzzify}, \texttt{adjustedRandIndex}
conformveg

Conform two community data tables

Description
Conforms two community data tables to have the same set of columns (species)

Usage
conformveg(x, y, fillvalue = 0, verbose = FALSE)

Arguments
x Community data, a site-by-species matrix.
y Community data, a site-by-species matrix.
fillvalue The value to be used to fill new entries in inflated matrices.
verbose Displays information about the number of species shared between x and y, as well as the number of species that are in one of the data tables but not in the other.

Details
This function adds to x as many new columns as columns of y that are not in x. The same is done for y, so the two tables have the same set of columns when they are returned.

Value
A list with the two inflated matrices x and y.

Author(s)
Miquel De Cáceres, Centre Tecnologic Forestal de Catalunya.

See Also
vegclust, vegclass

Examples
## Loads data (38 columns and 33 species)
data(wetland)
dim(wetland)

## Splits wetland data into two matrices of 30x27 and 11x22
wetland.30 = wetland[1:30,]
wetland.30 = wetland.30[,colSums(wetland.30)>0]
dim(wetland.30)
wetland.11 = wetland[31:41,]
wetland.11 = wetland.11[, colSums(wetland.11) > 0]
dim(wetland.11)

## Conforms the two matrices so they can eventually be merged
wetland.cf = conformveg(wetland.30, wetland.11)
dim(wetland.cf$x)
dim(wetland.cf$y)
names(wetland.cf$x) == names(wetland.cf$y)

crossmemb

### Cross-table of two fuzzy classifications

**Description**

Calculates a cross-tabulated matrix relating two fuzzy membership matrices

**Usage**

crossmemb(x, y, relativize = TRUE)

**Arguments**

- x: A site-by-group fuzzy membership matrix. Alternatively, an object of class 'vegclust' or 'vegclass'.
- y: A site-by-group fuzzy membership matrix. Alternatively, an object of class 'vegclust' or 'vegclass'.
- relativize: If TRUE expresses the cross-tabulated values as proportions of cluster size in x.

**Value**

A cross-tabulated matrix comparing the two classifications. In general, each cell’s value is the (fuzzy) number of objects that in x are assigned to the cluster corresponding to the row and in y are assigned to the cluster corresponding to the column. If relativize=TRUE then the values of each row are divided by the (fuzzy) size of the corresponding cluster in x.

**Author(s)**

Miquel De Cáceres, Forest Science Center of Catalonia.

**See Also**

defuzzify, vegclust
**defuzzify**

**Examples**

```r
## Loads data
data(wetland)

## This equals the chord transformation
## (see also \code{\link{decostand}} in package vegan)
wetland.chord = as.data.frame(sweep(as.matrix(wetland), 1,
                                    sqrt(rowSums(as.matrix(wetland)^2)), "/"))

## Create clustering with 3 clusters. Perform 10 starts from random seeds
## and keep the best solution. Try both FCM and NC methods:
wetland.fcm = vegclust(wetland.chord, mobileCenters=3, m = 1.2, method="FCM", nstart=10)
wetland.nc = vegclust(wetland.chord, mobileCenters=3, m = 1.2, dnoise=0.75, method="NC",
                      nstart=10)

## Compare the results
crossmemb(wetland.fcm, wetland.nc, relativize=FALSE)
```

---

**defuzzify**  

*Defuzzifies a fuzzy partition*

**Description**

Transforms a fuzzy classification into a crisp (hard) classification.

**Usage**

```r
defuzzify(object, method = "max", alpha = 0.5, na.rm = FALSE)
```

**Arguments**

- **object**: A site-by-group fuzzy membership matrix. Alternatively, an object of class 'vegclus' or 'vegclass'.
- **method**: Either "max" to choose for the maximum membership value across clusters, or "cut" for an alpha-cut.
- **alpha**: Threshold for the alpha-cut, bounded between 0 and 1.
- **na.rm**: If TRUE removes the objects that do not belong to any cluster when using method="cut".

**Details**

Alpha-cut means that memberships lower than alpha are transformed into 0 while memberships higher than alpha are transformed into 1. This means that if alpha values are low (i.e. close to 0), an object may belong to more than one group after defuzzification. These will generate a concatenation of cluster names in the output cluster vector and a row with sum more than one in the memb matrix. Similarly, if alpha is high (i.e. close to 1) there are objects that may be left unclassified. These will get NA in the cluster vector and zero row in the memb matrix.
Value

A list with the following items:

memb       A data frame with the hard membership partition.
cluster    A vector (factor) with the name of the cluster for each object.

Author(s)

Miquel De Cáceres, Forest Science Center of Catalonia.

References


See Also

cmeans, vegclust

Examples

```r
## Loads data
data(wetland)

## This equals the chord transformation
## (see also \code{\link{decostand}} in package vegan)
wetland.chord = as.data.frame(sweep(as.matrix(wetland), 1,
                           sqrt(rowSums(as.matrix(wetland)^2)), "/"))

## Create noise clustering with 3 clusters. Perform 10 starts from random seeds
## and keep the best solution
wetland.nc = vegclust(wetland.chord, mobileCenters=3, m = 1.2, dnoise=0.75,
                      method="NC", nstart=10)

## Defuzzification using an alpha-cut (alpha=0.5)
wetland.nc.df = defuzzify(wetland.nc$memb, method="cut")

## Cluster vector, with 'N' for objects that are unclassified,
## and 'NA' for objects that are intermediate
print(wetland.nc.df$cluster)

## Hard membership matrix (site 22 does not get any cluster assigned)
print(wetland.nc.df$memb)
```
**hcr**

**Heterogeneity-constrained random resampling (HCR)**

**Description**

Returns a set of indices of the original data set that maximizes the mean and minimizes the variance of the distances between pairs of plot records.

**Usage**

```r
hcr(d, nout, nsampl=1000)
```

**Arguments**

- `d`: An object of class `dist` containing the distance values between pairs of sites (plot records).
- `nout`: The number of sites (plot records) to be chosen among those available in `d`.
- `nsampl`: The number of resampling trials to be compared.

**Details**

Many subsets of the input data are selected randomly. These subsets are sorted by decreasing mean dissimilarity between pairs of plot records, and then sorted again by increasing variance of these dissimilarities. Ranks from both sortings are summed for each subset, and the subset with the lowest summed rank is considered as the most representative.

**Value**

Returns a vector containing the indices of the selected sites (plot records) to be used for sub-setting the original table.

**Author(s)**

Miquel De Cáceres, Forest Science Center of Catalonia

**References**


**Examples**

```r
## Loads data (38 columns and 33 species)
data(wetland)
dim(wetland)

## Constructs the chord distance matrix
## (see also \code{\link{decostand}} in package vegan)
```
hier.vegclust

Clustering with several number of clusters

Description

Performs several runs of function 'vegclust' (or 'vegclustdist') on a community data matrix (or distance matrix) using different number of clusters

Usage

hier.vegclust(x, hclust, cmin=2, cmax=20, min.size=NULL, verbose=TRUE, ...)
hier.vegclustdist(x, hclust, cmin=2, cmax=20, min.size=NULL, verbose=TRUE, ...)
random.vegclust(x, cmin=2, cmax=20, nstart=10, min.size=NULL, verbose=TRUE, ...)
random.vegclustdist(x, cmin=2, cmax=20, nstart=10, min.size=NULL, verbose=TRUE, ...)

Arguments

x For hier.vegclust and random.vegclust, a site (rows) by species (columns) matrix or data frame. For hier.vegclustdist and random.vegclustdist, a square distance matrix.
hclust A hierarchical clustering represented in an object of type hclust.
cmin Number of minimum mobile clusters.
cmax Number of maximum mobile clusters.
nstart A number indicating how many random trials should be performed for each number of groups
min.size If min.size != NULL, it specifies the minimum size of clusters. If some clusters are smaller, the algorithm will return the solutions corresponding to lower numbers of clusters.
verbose Flag used to print which number of clusters is currently running.
... Additional parameters for function vegclust or vegclustdist.
**incr.vegclust**

**Description**

Performs several runs of function `vegclust` on a community data matrix using an increasing number of clusters until some conditions are met.
Usage

```r
incr.vegclust(x, method="NC", ini.fixed.centers = NULL,
min.size = 10, max.var=NULL, alpha = 0.5,
nstart=100, fix.previous = TRUE, dnoise=0.75, m=1.0,...)
```

Arguments

- **x**: Community data table. A site (rows) by species (columns) matrix or data frame.
- **method**: A clustering model. Current accepted models are of the noise clustering family:
  - "NC": Noise clustering (Dave and Krishnapuram 1997)
  - "NCdd": Noise clustering with medoids
  - "HNC": Hard noise clustering
  - "HNCDdd": Hard noise clustering with medoids
- **ini.fixed.centers**: The coordinates of initial fixed cluster centers. These will be used as `fixedCenters` in all calls to `vegclust`. If `method="NCdd"` or `method="HNCDdd"` then `ini.fixed.centers` can be specified as a vector of indices for medoids.
- **min.size**: The minimum size (cardinality) of clusters. If any of the current k clusters does not have enough members the algorithm will stop and return the solution with k-1 clusters.
- **max.var**: The maximum variance allowed for clusters (see function `clustvar`). If specified, the algorithm will stop when any of the clusters is at the same time small and has large variance. If `max.var = NULL` then this criterion is not used.
- **alpha**: Criterion to choose cluster seeds from the noise class. Specifically, an object is considered as cluster seed if the membership to the noise class is larger than alpha.
- **nstart**: A number indicating how many random trials should be performed for number of groups. Each random trial uses the k-1 cluster centers plus the coordinates of the current cluster seed as initial solution for `vegclust`. Thus, if there are less cluster seed candidates than `nstart`, then not all runs are conducted.
- **fix.previous**: Flag used to indicate that the cluster centers found when determining k-1 clusters are fixed when determining k clusters.
- **m**: The fuzziness exponent.
- **dnoise**: The distance to the noise cluster.
- ... Additional parameters for function `vegclust`.

Details

Function `hier.vegclust` takes starting cluster configurations from cuts of a dendrogram given by object `hclust`. Function `random.vegclust` chooses random objects as cluster centroids and for each number of clusters performs `nstart` trials.

Value

Returns an object of class `vegclust`; or `NULL` if the initial cluster does not contain enough members.
interclustdist

Author(s)
Miquel De Cáceres, Forest Science Center of Catalonia

References

See Also
vegclust, hier.vegclust

Examples

```r
## Loads data
data(wetland)

## This equals the chord transformation
## (see also \code{\link{decostand}} in package \code{vegan})
wetland.chord = as.data.frame(sweep(as.matrix(wetland), 1,
sqrt(rowSums(as.matrix(wetland)^2)), ")/\))

## Call incremental noise clustering
wetland.nc = incr.vegclust(wetland.chord, method="NC", m = 1.2, dnoise=0.75,
min.size=5)

## Inspect cluster sizes
print(wetland.nc$size)
```

---

**interclustdist**

*Calculates the distance between pairs of cluster centroids*

Description
Calculates the distance between pairs of cluster centroids, given a distance matrix and a cluster vector.

Usage

```r
interclustdist(x, cluster)
```

Arguments

- **x**: A site-by-site data matrix or an object of class `dist` containing the distance values between pairs of sites (plot records).
- **cluster**: A vector indicating the hard membership of each object in `x` to a set of groups. Can contain `NA` values.
Value

An object of class `dist` containing the distances between cluster centers.

Author(s)

Miquel De Cáceres, Forest Science Center of Catalonia

Examples

```r
##TO BE DONE##
```

<table>
<thead>
<tr>
<th>medreg</th>
<th>Regeneration of Mediterranean vegetation data set</th>
</tr>
</thead>
</table>

Description

A stratified vegetation data set containing with several plot records laid to assess vegetation recovery three years after a wildfire. Collected in 2012 by Miquel De Cáceres and Albert Petit in Horta de Sant Joan (Catalonia, Spain).

Usage

```r
data(medreg)
```

Format

An object of class `stratifiedvegdata` with 96 elements (plots), each of them consisting of a data.frame where rows correspond to species groups and columns correspond to vegetation strata. Abundance values are percentage cover.

See Also

`CAP`, `plot.CAP`

Examples

```r
data(medreg)
```
plot.CAP

Draws cumulative abundance profiles

Description

Create plots used to inspect one or more cumulative abundance profiles.

Usage

## S3 method for class 'CAP'
plot(x, sizes=NULL, species=NULL, plots=NULL, switchAxes=FALSE,
    add=FALSE, drawAxes = TRUE, xlab="", ylab="", type="s", ...)

## S3 method for class 'stratifiedvegdata'
plot(x, sizes=NULL, species=NULL, plots=NULL, switchAxes=FALSE,
    add=FALSE, drawAxes = TRUE, xlab="", ylab="", type="s", ...)

Arguments

- **x**: An object returned from function `CAP` or an object of class `stratifiedvegdata` (see documentation for function `stratifyvegdata`).
- **sizes**: A vector containing the size values associated to each size class. If `NULL` the y-axis will be defined using the size class order in `x`.
- **species**: A vector of strings indicating the species whose profile is to be drawn. If `NULL` all species are plotted.
- **plots**: A vector indicating the plot records whose profile is to be drawn. Can be a character vector (for plot names), a numeric vector (for plot indices) or a logical vector (for TRUE/FALSE selection). If `NULL` all plot records are plotted.
- **switchAxes**: A flag indicating whether ordinate and abscissa axes should be interchanged.
- **add**: A flag indicating whether profiles should be drawn on top of current drawing area. If `add=FALSE` a new plot is created.
- **drawAxes**: A flag indicating whether axes should be drawn.
- **xlab**: String label for the x axis.
- **ylab**: String label for the y axis.
- **type**: Type of plot to be drawn ("p" for points, "l" for lines, "s" for steps, ...).
- **...**: Additional plotting parameters.

Author(s)

Miquel De Cáceres, Forest Science Center of Catalonia

References

See Also

CAP

Examples

```r
## Load stratified data
data(medreg)

## Check that 'medreg' has correct class
class(medreg)

## Create cumulative abundance profile (CAP) for each plot
medreg.CAP = CAP(medreg)

## Draw the stratified data and profile corresponding to the third plot
plot(medreg, plots="3")
plot(medreg.CAP, plots="3")

## Look at the plot and CAP of the same plot
medreg[[3]]
medreg.CAP[[3]]
```

---

**plot.CAS**

Draws a cumulative abundance surface

Description

Create plots used to inspect one or more cumulative abundance profiles.

Usage

```r
## S3 method for class 'CAS'
plot(x, plot=NULL, species=NULL, sizes1=NULL, sizes2 = NULL,
     palette = colorRampPalette(c("light blue","light green","white",
                                "yellow","orange","red")), zlim=NULL,...)
```

Arguments

- `x` An object of class **CAS**.
- `plot` A string indicating the plot record whose surface is to be drawn.
- `species` A string indicating the species whose profile is to be drawn.
- `sizes1` A vector containing the size values associated to each primary size class. If NULL the x-axis will be defined using the primary size class order in `x`.
- `sizes2` A vector containing the size values associated to each secondary size class. If NULL the y-axis will be defined using the secondary size class order in `x`.
- `palette` Color palette for z values.
- `zlim` The limits for the z-axis.
- `...` Additional plotting parameters for function **persp**.
Author(s)
Miquel De Cáceres, Forest Science Center of Catalonia

References

See Also
Cas, persp

Examples

```r
## Create synthetic tree data
pl = rep(1,100) # All trees in the same plot
sp = ifelse(runif(100)>0.5,1,2) # Random species identity (species 1 or 2)
h=rgamma(100,10,2) # Heights (m)
d = rpois(100, lambda=h^2) # Diameters (cm)
m = data.frame(plot=pl,species=sp, height=h,diameter=d)
m$ba = pi*(m$diameter/200)^2
print(head(m))

## Size classes
heights = seq(0,4, by=.25)^2 # Quadratic classes
diams = seq(0,130, by=5) # Linear classes

## Stratify tree data
X<-stratifyvegdata(m, sizes1=heights, sizes2=diams,
plotColumn = "plot", speciesColumn = "species",
size1Column = "height", size2Column = "diameter",
abundanceColumn = "ba")

## Build cumulative abundance surface
Y = CAS(X)

## Plot the surface of species '1' in plot '1' using heights and diameters
plot(Y, species=1, sizes1=heights[-1], xlab="height (m)",
ylab="diameter (cm)", sizes2=diams[-1], zlab="Basal area (m2)",
xlim = c(0,6), main="Species 1")
```

Description
Create plots used to study vegclust clustering results for an increasing number of clusters.
Usage

## S3 method for class 'mvegclust'
plot(x, type="hnc", excludeFixed=TRUE, verbose=FALSE, ylim=NULL, xlab=NULL, ylab=NULL, maxvar=0.6, minsize=20,...)

Arguments

x An object returned from functions `hier.vegclust` or `random.vegclust`.

type A string indicating the type of plot desired. Current accepted values are "hnc","hmemb","var","hcs" and "valid".

excludeFixed A flag to indicate whether clusters with fixed centroids should be excluded from plots.

verbose A flag to print extra information.

ylim A vector with the limits for the y axis.

xlab String label for the x axis.

ylab String label for the y axis.

maxvar Maximum cluster variance allowed for the type="valid" plot.

minsize Minimum cluster size allowed for the type="valid" plot.

... Additional plotting parameters.

Value

Different information is returned depending on the type of plot chosen.

Author(s)

Miquel De Cáceres, Forest Science Center of Catalonia

Examples

## Loads data
data(wetland)

## This equals the chord transformation
## (see also \code{\link{decostand}} in package 'vegan')
wetland.chord = as.data.frame(sweep(as.matrix(wetland), 1, sqrt(rowSums(as.matrix(wetland)^2)), "/"))

## Create noise clustering from hierarchical clustering at different number of clusters
wetland.hc = hclust(dist(wetland.chord), method="ward")
wetland.nc = hier.vegclust(wetland.chord, wetland.hc, cmin=2, cmax=5, m = 1.2, dnoise=0.75, method="NC")

## Plot changes in the number of objects falling into the noise cluster
plot(wetland.nc, type="hnc")

## Plots the number of objects falling into "true" clusters,
relate.levels

## the number of objects considered intermediate,
## and the number of objects falling into the noise
plot(wetland.nc, type="hmemb")

## Plot minimum, maximum and average cluster size
plot(wetland.nc, type="hcs")

## Plot minimum, maximum and average cluster variance
plot(wetland.nc, type="var")

## Plot number of groups with high variance, low membership or both
plot(wetland.nc, type="valid")

---

**relate.levels**

Relates two clustering level results.

**Description**

Analyzes how lower level clusters are assigned into upper level ones. The analysis is made for several number of clusters.

**Usage**

```r
relate.levels(lower, upper, defuzzify = FALSE, excludeFixed = FALSE, verbose=FALSE, ...)
```

**Arguments**

- `lower` A list of objects of type `vegclust` or `vegclass` that represent classifications at a finer level of resolution.
- `upper` A list of objects of type `vegclust` or `vegclass` that represent classifications at an broader level of resolution.
- `defuzzify` A logical flag used to indicate whether the result of calling `crossmemb` should be defuzzified.
- `excludeFixed` A logical used to indicate whether fixed clusters should be excluded from the comparison of levels.
- `verbose` A flag used to ask for extra screen output.
- `...` Additional parameters for function `defuzzify`.

**Details**

For each pair of `vegclust` (or `vegclass`) objects in `upper` and `lower`, the function calls function `crossmemb` and then, if asked, defuzzifies the resulting memberships (by calling function `defuzzify`) and several quantities are calculated (see 'value' section).
Value

A list with several data frames (see below). In each of them, the rows are items of upper and columns are items of lower. The names of rows and columns are the number of clusters of each `vegclust` (or `vegclass`) object.

- `nnoise` The number of low level clusters that are assigned to the Noise class (for upper objects using Noise clustering).
- `maxnoise` The maximum membership value of low level clusters to the Noise class (for upper objects using Noise clustering).
- `minmaxall` The minimum value (across upper level clusters) of the maximum membership value observed among the lower level clusters.
- `minallsize` The minimum value (across upper level clusters) of the sum of membership values across lower level clusters.
- `empty` The number of upper level clusters (mobile or fixed) that do not have any member among the lower level clusters.

Author(s)

Miquel De Cáceres, Forest Science Center of Catalonia

See Also

`vegclust`, `vegclass`, `defuzzify`

Examples

```r
## Loads data
data(wetland)

## This equals the chord transformation
## (see also \code{\link{decostand}} in package vegan)
wetland.chord = as.data.frame(sweep(as.matrix(wetland), 1, 
                           sqrt(rowSums(as.matrix(wetland)^2)), 
                           Method=2))

## Create noise clustering from hierarchical clustering at different number of cluster
wetland.hc = hclust(dist(wetland.chord), method="ward")
wetland.nc1 = hier.vegclust(wetland.chord, wetland.hc, cmin=2, cmax=6, m = 1.2, 
                           dnoise=0.75, method="NC")
wetland.nc2 = hier.vegclust(wetland.chord, wetland.hc, cmin=2, cmax=4, m = 1.2, 
                           dnoise=0.85, method="NC")

## Studies the assignment of levels
relate.levels(wetland.nc1, wetland.nc2, method="cut")
```
stratifyvegdata

Reshapes community data from individual into stratified form

Description

Function stratifyvegdata reshapes individual abundance values into species abundance values per size class or combination of size classes. Function as.stratifiedvegdata checks if the input list has appropriate properties and turns it into an object of class 'stratifiedvegdata'.

Usage

stratifyvegdata(x, sizes1, sizes2 = NULL, treesel=NULL, spcodes = NULL,
plotColumn="plot", speciesColumn = "species",
abundanceColumn="abundance", size1Column = "size", size2Column = NULL,
cumulative=FALSE, counts=FALSE, mergeSpecies=FALSE, verbose=FALSE)
as.stratifiedvegdata(X)

Arguments

x A data frame containing individual plant data. Individuals are in rows, while measurements are in columns.
sizes1 A numerical vector containing the breaks for primary size classes in ascending order.
sizes2 A numerical vector containing the breaks for secondary size classes in ascending order.
treesel A logical vector specifying which rows in x to be used. By default (treesel = NULL) all rows are taken.
spcodes A character vector indicating the codes of species to be used for stratification (species codes beyond those appearing in x are possible). If spcodes = NULL then all species in x are used.
plotColumn The name of the column in x that contains plot identifiers.
speciesColumn The name of the column in x that contains species names.
abundanceColumn The name of the column in x that contains abundance values.
size1Column The name of the column in x that contains values for primary size classes.
size2Column The name of the column in x that contains values for secondary size classes.
cumulative A flag to indicate that cumulative abundance profiles or surfaces are desired.
counts A flag to indicate that the output should be individual counts instead of added abundance values.
mergeSpecies A flag to indicate that species identity should be ignored. This leads to analyzing the structure of biomass disregarding species identity.
verbose A logical flag to indicate extra output.
stratifyvegdata

X

A list with as many elements as plot records. Each element should be of class 'matrix' or 'data.frame' with species in rows and strata in columns. Furthermore, the number of rows (species) and columns (strata) should be the same for all elements.

Details

For each individual (row) in x, stratifyvegdata assigns it to the size class (stratum) containing its size. The corresponding abundance value (e.g. crown cover) of the individual is added to the abundance of the corresponding species at the size class (stratum). If sizes2 and size2Column are supplied, the function assigns each individual (row) in x to the combination of size classes (e.g. tree height and diameter).

Value

Both functions return an object of class 'stratifiedvegdata', which is a list of matrices, one for each plot record. Each element (matrix) has as many rows as species and as many columns as size classes (i.e., as many as elements in vector sizes1). Columns are named starting with 'S' and continuing with the size class (stratum) number. If mergeSpecies=TRUE then all matrices have a single row (whose name is "all"). If sizes2 and size2Column are supplied to stratifyvegdata, the function returns an object of class 'doublestratifiedvegdata', which is a list of arrays, one for each plot record. Each element (array) has three dimensions corresponding to species, primary sizes (number of elements in in vector sizes1) and secondary sizes (number of elements in in vector sizes2). If cumulative=TRUE then the function returns cumulative abundances (see CAP and CAS).

Author(s)

Miquel De Cáceres, Forest Science Center of Catalonia.

References


See Also

reshape, CAP, CAS

Examples

```R
## Load tree data
data(treedata)

## Inspect tree data
head(treedata)

## Define stratum thresholds (4 strata)
heights = seq(0,4, by=0.5)
diameters = seq(0,2, by=0.5)

## Stratify tree data using heights as structural variable
```
X = stratifyvegdata(treedata, sizes!=heights, plotColumn="plotID",
                   speciesColumn="species", size1Column="height", counts=TRUE)

## Inspect the second plot record
X[[2]]

## Stratify tree data using heights as structural variable and cover as abundance
Y = stratifyvegdata(treedata, sizes!=heights, plotColumn="plotID",
                    speciesColumn="species", size1Column="height",
                    abundanceColumn="cover")
Y[[2]]

## Stratify tree data using heights and diameters as structural variables
Z = stratifyvegdata(treedata, sizes!=heights, sizes2=diameters, plotColumn="plotID",
                    speciesColumn="species", size1Column="height", size2Column="diam",
                    counts=TRUE)
Z[[2]]

---

**trajectories**

**Community trajectory analysis**

**Description**

Given a distance matrix between community states, functions `segmentdistances` and `trajectorydistances` calculate the distance between pairs of directed segments and community trajectories, respectively. Function `trajectorylengths` calculates lengths of directed segments and complete trajectories. Function `trajectorypcoa` performs principal coordinates analysis (`cmdscale`) and draws trajectories in the ordination scatterplot.

**Usage**

```
segmentdistances(d, sites, surveys = NULL,
                  distance.type = "directed-segment", verbose = FALSE)
```

```
trajectorydistances(d, sites, surveys = NULL, distance.type = "DSPD",
                   symmetrization = "mean", verbose = FALSE)
```

```
trajectorylengths(d, sites, surveys = NULL, verbose = FALSE)
```

```
trajectorypcoa(d, sites, surveys = NULL, selection = NULL,
               traj.colors = NULL, axes = c(1, 2), ...)
```

**Arguments**

- `d`: A symmetric matrix or an object of class `dist` containing the distance values between pairs of community states.
sites A vector indicating the site corresponding to each community state.
surveys A vector indicating the survey corresponding to each community state (only necessary when surveys are not in order).
distance.type The type of distance index to be calculated (Besse et al. 2016; De Cáceres et al. submitted). For segmentDistances the available indices are:
  • Hausdorff: Hausdorff distance between two segments.
  • directed-segment: Directed segment distance (default).
  • PPA: Perpendicular-parallel-angle distance.
whereas for trajectoryDistances the available indices are:
  • Hausdorff: Hausdorff distance between two trajectories.
  • SPD: Segment path distance.
  • DSPD: Directed segment path distance (default).
verbose Provides console output informing about process (useful for large dataset).
symmetrization Function used to obtain a symmetric distance, so that DSPD(T1,T2) = DSPD(T2,T1) (e.g., mean or min).
selection A numeric or logical vector of the same length as sites, indicating a subset of site trajectories to be plotted.
traj.colors A vector of colors (one per site). If selection != NULL the length of the color vector should be equal to the number of sites selected.
aves The pair of principal coordinates to be plotted.
... Additional parameters for function arrows.

Details

These functions consider community dynamics as trajectories in a chosen space of community resemblance and takes trajectories as objects to be compared. By adapting concepts and procedures used for the analysis of trajectories in space (i.e. movement data) (Besse et al. 2016), the functions allow assessing the resemblance between trajectories. Details of calculations are given in De Cáceres et al (submitted)

Value

Function trajectoryDistances returns an object of class dist containing the distances between trajectories. Function trajectorySegments returns a list with the following elements:

  • Dseg: Distance matrix between segments.
  • Dini: Distance matrix between initial points of segments.
  • Dfin: Distance matrix between final points of segments.
  • Dini-fin: Distance matrix between initial points of one segment and the final point of the other.
  • Dfinini: Distance matrix between final points of one segment and the initial point of the other.

Function trajectoryLengths returns a data frame with the length of each segment on each trajectory and the total length of all trajectories. Function trajectoryPCoA returns the result of calling cmdscale.
trajectories

Author(s)
Miquel De Cáceres, Forest Sciences Center of Catalonia

References


See Also
cmdscale

Examples

# Description of sites and surveys
sites = c(1,1,1,2,2,2)
surveys=c(1,2,3,1,2,3)

# Raw data table
xy<-matrix(0, nrow=6, ncol=2)
x[2,2]<-1
xy[3,2]<-2
xy[4:6,1] <- .5
xy[4:6,2] <- xy[1:3,2]
x[6,1]<-1

# Distance matrix
d = dist(xy)
d

trajectoryLengths(d, sites, surveys)$dseg
segmentDistances(d, sites, surveys)
trajectoryDistances(d, sites, surveys, distance.type = "Hausdorff")

# Draw trajectories
trajectoryPCoa(d, sites, traj.colors = c("black","red"), lwd = 2)

# Should give the same results if surveys are not in order
# (here we switch surveys for site 2)
temp = xy[5,]
xy[5,] = xy[6,]
xy[6,] = temp
surveys[5] = 3
surveys[6] = 2
trajectoryLengths(dist(xy), sites, surveys)
segmentDistances(dist(xy), sites, surveys)$dseg
trajectoryDistances(dist(xy), sites, surveys, distance.type = "Hausdorff")
**treedata**  
*Synthetic vegetation data set with tree data*

**Description**

A synthetic data set used to illustrate the stratification of data originally collected on an individual basis (e.g. forest inventory).

**Usage**

```r
data(treedata)
```

**Format**

A data frame where each row corresponds to a different tree. Columns are plot code, species identity, tree height, tree diameter and cover value.

**See Also**

`stratifyvegdata`

---

**vegclass**  
*Classifies vegetation communities*

**Description**

Classifies vegetation communities into a previous fuzzy or hard classification.

**Usage**

```r
vegclass(y, x)
```

**Arguments**

- `y`  
  An object of class `vegclust` that represents a previous knowledge.

- `x`  
  Community data to be classified, in form of a site by species matrix (if the `vegclust` object is in `raw` mode) or a data frame containing the distances between the new sites in rows and the old sites in columns (if the `vegclust` object is in `distance` mode).
Details

This function uses the classification model specified in \(y\) to classify the communities (rows) in \(x\). When vegclust is in `raw` mode, the function calls first to `conformveg` in order to cope with different sets of species. See the help of `as.vegclust` to see an example of vegclass with distance matrices.

Value

Returns an object of type `vegclass` with the following items:

- **method**: The clustering model used in \(y\)
- **m**: The fuzziness exponent in \(y\)
- **dnoise**: The distance to the noise cluster used for noise clustering (models NC, NCdd, HNC, HNCdd). This is set to NULL for other models.
- **eta**: The reference distance vector used for possibilistic clustering (models PCM and PCMdd). This is set to NULL for other models.
- **memb**: The fuzzy membership matrix.
- **dist2clusters**: The matrix of object distances to cluster centers.

Author(s)

Miquel De Cáceres, Forest Science Center of Catalonia.

References


See Also

`vegclust, as.vegclust, kmeans, cmeans, conformveg`

Examples

```r
## Loads data (38 columns and 33 species)
data(wetland)
dim(wetland)

## This equals the chord transformation
## (see also \code{\link{decostand}} in package vegan)
wetland.chord = as.data.frame(sweep(as.matrix(wetland), 1,
                                 sqrt(rowSums(as.matrix(wetland)^2)), "/"))

## Splits wetland data into two matrices of 30x27 and 11x22
```
### vegclust

**Vegetation clustering methods**

#### Description

Performs hard or fuzzy clustering of vegetation data

#### Usage

```r
vegclust(x, mobileCenters, fixedCenters = NULL, method="NC", m = 2, dnoise = NULL, eta = NULL, alpha=0.001, iter.max=100, nstart=1, maxminJ = 10, seeds=NULL, verbose=FALSE)
vegclustdist(x, mobileMemb, fixedDistToCenters = NULL, method="NC", m = 2, dnoise = NULL, eta = NULL, alpha=0.001, iter.max=100, nstart=1, seeds=NULL, verbose=FALSE)
```

#### Arguments

- **x**
  Community data. A site-by-species matrix or data frame (for `vegclust`) or a site-by-site dissimilarity matrix or `dist` object (for `vegclustdist`).

- **mobileCenters**
  A number, a vector of seeds, or coordinates for mobile clusters.

- **fixedCenters**
  A matrix or data frame with coordinates for fixed (non-mobile) clusters.

- **mobileMemb**
  A number, a vector of seeds, or starting memberships for mobile clusters.

- **fixedDistToCenters**
  A matrix or data frame with the distances to fixed cluster centers.
method A clustering model. Current accepted models are:

- "KM": K-means or hard c-means (MacQueen 1967)
- "KMdd": Hard c-medoids (Krishnapuram et al. 1999)
- "FCM": Fuzzy c-means (Bezdek 1981)
- "FCMdd": Fuzzy c-medoids (Krishnapuram et al. 1999)
- "NC": Noise clustering (Dave and Krishnapuram 1997)
- "NCdd": Noise clustering with medoids
- "HNC": Hard noise clustering
- "HNCdd": Hard noise clustering with medoids
- "PCM": Possibilistic c-means (Krishnapuram and Keller 1993)
- "PCMdd": Possibilistic c-medoids

m The fuzziness exponent to be used (this is relevant for all models except for kmeans)

dnoise The distance to the noise cluster, relevant for noise clustering (NC).

eta A vector of reference distances, relevant for possibilistic C-means (PCM).

alpha Threshold used to stop iterations. The maximum difference in the membership matrix of the current vs. the previous iteration will be compared to this value.

iter.max The maximum number of iterations allowed.

nstart If mobileCenters or mobileMemb is a number, how many random sets should be chosen?

maxminj When random starts are used, these will stop if at least maxminj runs ended up in the same functional value.

seeds If mobileCenters or mobileMemb is a number, a vector indicating which objects are potential initial centers. If NULL all objects are valid seeds.

verbose Flag to print extra output.

Details

Functions vegclust and vegclustdist try to generalize the kmeans function in stats in three ways.

Firstly, they allow different clustering models. Clustering models can be divided in (a) fuzzy or hard; (b) centroid-based or medoid-based; (c) Partitioning (KM and FCM family), noise clustering (NC family), and possibilistic clustering (PCM and PCMdd). The reader should refer to the original publications to better understand the differences between models.

Secondly, users can specify fixed clusters (that is, centroids that do not change their positions during iterations). Fixed clusters are intended to be used when some clusters were previously defined and new data has been collected. One may allow some of these new data points to form new clusters, while some other points will be assigned to the original clusters. In the case of models with cluster repulsion (such as KM, FCM or NC) the new (mobile) clusters are not allowed to ‘push’ the fixed ones. As a result, mobile clusters will occupy new regions of the reference space.

Thirdly, vegclustdist implements the distance-based equivalent of vegclust. The results of vegclust and vegclustdist will be the same (if seeds are equal) if the distance matrix is calculated using the Euclidean distance (see function dist). Otherwise, the equivalence holds by resorting on principal coordinates analysis.
Note that all data frames or matrices used as input of `vegclust` should be defined on the same space of species (see `conformveg`). Unlike `kmeans`, which allows different specific algorithms, here updates of prototypes (centroids or medoids) are done after all objects have been reassigned (Forgy 1965). In order to obtain hard cluster definitions, users can apply the function `defuzzify` to the `vegclust` object.

**Value**

Returns an object of type `vegclust` with the following items:

- **mode**  
  raw for function `vegclust` and dist for function `vegclustdist`.

- **method**  
  The clustering model used

- **m**  
  The fuzziness exponent used ($m=1$ in case of kmeans)

- **dnoise**  
  The distance to the noise cluster used for noise clustering (NC, HNC, NCdd or HNCdd). This is set to NULL for other models.

- **eta**  
  The reference distance vector used for possibilistic clustering (PCM or PCMdd). This is set to NULL for other models.

- **memb**  
  The fuzzy membership matrix. Columns starting with "M" indicate mobile clusters, whereas columns starting with "F" indicate fixed clusters.

- **mobileCenters**  
  If `vegclust` is used, this contains a data frame with the coordinates of the mobile centers (centroids or medoids). If `vegclustdist` is used, it will contain the indices of mobile medoids for models KMdd, FCMdd, HNCdd, NCdd and PCMdd; or NULL otherwise.

- **fixedCenters**  
  If `vegclust` is used, this contains a data frame with the coordinates of the fixed centers (centroids or medoids). If `vegclustdist` is used, it will contain the indices of fixed medoids for models KMdd, FCMdd, HNCdd, NCdd and PCMdd; or NULL otherwise.

- **dist2clusters**  
  The matrix of object distances to cluster centers. Columns starting with "M" indicate mobile clusters, whereas columns starting with "F" indicate fixed clusters.

- **withinss**  
  In the case of methods KM, FCM, NC, PCM and HNC it contains the within-cluster sum of squares for each cluster (squared distances to cluster center weighted by membership). In the case of methods KMdd, FCMdd, NCdd, HNCdd and PCMdd it contains the sum of distances to each cluster (weighted by membership).

- **size**  
  The number of objects belonging to each cluster. In case of fuzzy clusters the sum of memberships is given.

- **functional**  
  The objective function value (the minimum value attained after all iterations).

**Author(s)**

Miquel De Cáceres, Forest Science Center of Catalonia
References


See Also

hierNvegclust, incrNvegclust, kmeans, cmeans, vegclass, defuzzify, clustvar

Examples

```r
### Loads data
data(wetland)

### This equals the chord transformation
### (see also 'decostand' in package vegan)
wetland.chord = as.data.frame(sweep(as.matrix(wetland), 1,
  sqrt(rowSums(as.matrix(wetland)^2)), "/"))

### Create noise clustering with 3 clusters. Perform 10 starts from random seeds
### and keep the best solution
wetland.nc = vegclust(wetland.chord, mobileCenters=3, m = 1.2, dnoise=0.75,
  method="NC", nstart=10)

### Fuzzy membership matrix
wetland.nc$memb

### Cardinality of fuzzy clusters (i.e., the number of objects belonging to each cluster)
wetland.nc$size

### Obtains hard membership vector, with 'N' for objects that are unclassified
defuzzify(wetland.nc$memb)$cluster

### The same result is obtained with a matrix of chord distances
wetland.d = dist(wetland.chord)
wetland.d.nc = vegclustdist(wetland.d, mobileMem=3, m = 1.2, dnoise=0.75,
  method="NC", nstart=10)
```
Description

This function casts an object of class `vegclust` into an object of class `kmeans`.

Usage

vegclust2kmeans(x)

Arguments

x An object of class `vegclust` to be casted, where `method="KM"` and `mode="raw"`.

Author(s)

Miquel De Cáceres, Forest Science Center of Catalonia

See Also

`vegclust`, `kmeans`

Examples

```r
## Loads data
data(wetland)

## This equals the chord transformation
## (see also \code{\link{decostand}} in package vegan)
wetland.chord = as.data.frame(sweep(as.matrix(wetland), 1,
  sqrt(rowSums(as.matrix(wetland)^2)), "/"))

## Create noise clustering with 3 clusters. Perform 10 starts from random seeds
wetland vc = vegclust(wetland.chord, mobileCenters=3,
  method="KM", nstart=10)

## Reshapes as kmeans object
wetland.km = vegclust2kmeans(wetland vc)
wetland.km
```
**vegclustIndex**

*Compute fuzzy evaluation statistics*

**Description**

Computes several evaluation statistics on the fuzzy clustering results on objects of class `vegclust`.

**Usage**

`vegclustIndex(y)`

**Arguments**

- `y` An object of class `vegclust` or a membership matrix.

**Details**

These statistics were conceived to be computed on fuzzy partitions, such as the ones coming from Fuzzy C-means (Bezdek 1981). Maximum values of PCN or minimum values of PEN can be used as criteria to choose the number of clusters.

**Value**

Returns a vector of four values: partition coefficient (PC), normalized partition coefficient (PCN), partition entropy (PE) and normalized partition entropy (PEN).

**Author(s)**

Miquel De Cáceres, Forest Science Center of Catalonia

**References**


**See Also**

cmeans, vegclust

**Examples**

```r
## Loads data
data(wetland)

## This equals the chord transformation
## (see also \code{\link{decostand}} in package vegan)
wetland.chord = as.data.frame(sweep(as.matrix(wetland), 1, sqrt(rowSums(as.matrix(wetland)^2)), "/"))
```
## vegdiststruct

### Structural and compositional dissimilarity

#### Description

Function to calculate the dissimilarity between ecological communities taking into account both their composition and the size of organisms.

#### Usage

```r
type = "cumulative", method = "bray",
transform = NULL, classWeights = NULL)
```

#### Arguments

- **x**: A stratified vegetation data set (see function `stratifyvegdata`), a set of cumulative abundance profiles (see function `CAP`) or a set of cumulative abundance surfaces (see function `CAS`).
- **y**: A second stratified vegetation data set (see function `stratifyvegdata`), a second set of cumulative abundance profiles (see function `CAP`) or a second set of cumulative abundance surfaces (see function `CAS`) against which object `x` should be compared.
- **paired**: Only relevant when `y` != NULL. If `paired = TRUE` pairwise comparisons are calculated between elements in `x` and `y` (and `x` and `y` need to be of the same length). If `paired = FALSE` then all objects in `x` are compared to all objects in `y`.
- **type**: Whether dissimilarities between pairs of sites should be calculated from differences in cumulative abundance ("cumulative"), in total abundance ("total") or in volumes of cumulative abundance profiles ("volume").
- **method**: The dissimilarity coefficient to calculate (see details).
- **transform**: A function or the name of a function to be applied to each cumulative abundance value.
- **classWeights**: A numerical vector or a matrix containing the weight of each size class or combination of size classes (see functions `CAP2matrix` and `CAS2matrix`). If NULL, then the function assumes classes of equal weight.
Details

The six different coefficients available are described in De Cáceres et al. (2013): (1) method="bray" for percentage difference (alias Bray-Curtis dissimilarity); (2) method="ruzicka" for Ruzicka index (a generalization of Jaccard); (3) method="kulczynski" for the Kulczynski dissimilarity index; (4) method="ochiai" for the complement of a quantitative generalization of Ochiai index of similarity; (5) method="canberra" for the Canberra index (Adkins form); (6) method="relman" for the relativized Manhattan coefficient (Whittaker's index of association). Currently, the function also supports (7) method="manhattan" for the city block metric.

Value

Returns an object of class 'dist'.

Author(s)

Miquel De Cáceres, Forest Science Center of Catalonia.

References


See Also

stratifyvegdata, vegdist

Examples

```r
## Load stratified data
data(medreg)

## Check that 'medreg' has correct class
class(medreg)

## Create cumulative abundance profile (CAP) for each plot
medreg.CAP = CAP(medreg)

## Create dissimilarity (percentage difference) matrix using profiles
medreg.D = vegdiststruct(medreg, method="bray")

## Create dissimilarity (percentage difference) matrix using abundances
medreg.D2 = vegdiststruct(medreg, method="bray", type="total")

## Calculate correlation
cor(as.vector(medreg.D), as.vector(medreg.D2))
```
wetland 

Wetland vegetation data set

Description

Vegetation of the Adelaide river alluvial plain (Australia). This data set was published by Bowman & Wilson (1987) and used in Dale (1988) to compare fuzzy classification approaches.

Usage

data(wetland)

Format

A data frame with 41 sites (rows) and 33 species (columns). Abundance values are represented in abundance classes.

Source


References


Examples

data(wetland)
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