# Package ‘analogue’ 

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Maintainer Gavin L. Simpson [ucfagls@gmail.com](mailto:ucfagls@gmail.com)
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Author Gavin L. Simpson [aut, cre] ([https://orcid.org/0000-0002-9084-8413](https://orcid.org/0000-0002-9084-8413)),Jari Oksanen [aut],Martin Maechler [ctb]
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analogue-package Analogue and weighted averaging methods for palaeoecology

## Description

analogue is a package for quantitative palaeoecology with a focus on analogue methods, transfer functions, and data handling and display.

## Analogue methods

analogue provides functions for analogue matching and the modern analogue technique (MAT) via analog and mat. A wide range of dissimilarity coefficients are available via the distance function.

Additional analysis of modern and no-analogue problems is facilitated via a range of functions implementing many methods from the literature. In particular, the receiver operating characteric (ROC) curves method of Gavin et al (2003) is available in roc and a related method employing direct logistic regression modelling (Simpson \& Birks, 2012) is available in logitreg.

## Transfer function methods

Several approaches to fitting transfer function models are provided by analogue:
wa: Simple and tolerance-downweighted weighted averaging with classical, inverse, and monotonic spline deshrinking.
mat: The modern analogue technique (MAT).
pcr : Principal components regression with ecologically meaningful transformations
A range of functions for working with and exploring training sets and palaeoenvironmental reconstructions is also included in analogue. These include
crossval leave-one-out, repeated k-fold, and bootstrap cross-validation methods.
compare compare properties of taxa or other proxies across modern and fossil data sets.
evenSample are training set samples evenly distributed along the gradient of interest?
splitSample splits a gradient into a set of bins or chunks and samples evenly from within each chunk to create a representative test set for cross-validation.
timetrack overlays a fossil or secondary data set on to an (constrained) ordination of a modern or reference data set.
weightedCor implements the weighted correlation test of a Weighted Averaging reconstruction as proposed by Telford \& Birks (2011).

## Utilities

analogue provides a range of utilities for working with palaeo data.
tran a range of transformations applicable to or commonly used with palaeo data.
Stratiplot draws stratigraphic diagrams using the Lattice package.
join merging of modern/training set and fossil data sets.
chooseTaxa selects taxa that meet certain abundance and occurrence criteria.

## Documentation

A full tutorial and worked example for the main features of analogue matching and MAT is avilable in the vignette
analogue_methods Analogue Methods in Palaeoecology

## Author(s)

Gavin L. Simpson, Jari Oksanen
Maintainer: Gavin L. Simpson [ucfagls@gmail.com](mailto:ucfagls@gmail.com)

## References

Gavin, D.G., Oswald, W.W., Wahl, E.R. and Williams, J.W. (2003) A statistical approach to evaluating distance metrics and analog assignments for pollen records. Quaternary Research 60, 356-367.
Simpson, G.L. \& Birks H.J.B. (2012) Statistical Learning in Palaeolimnology. In Birks, H.J.B, Lotter, A.F. Juggins S., and Smol, J.P. (Eds) Tracking Environmental Change Using Lake Sediments, Volume 5: Data Handling and Numerical Techniques. Springer, Dordrecht.
Telford R.J. and Birks, H.J.B. (2011) A novel method for assessing the statistical significance of quantitative reconstructions inferred from biotic assemblages. Quanternary Science Reviews 30:1272-1278.
abernethy Abernethy Forest Pollen Sequence

## Description

The classic pollen data set from Abernethy Forest in the Scottish highlands, UK. The data originate from the work of Hilary Birks and Rolf Mathewes (1978) and have been analysed in several texts on quantitative numerical palaeoecology.
The data set consists of 36 pollen taxa from 49 levels, with two additional variables; Age, the age of each sample, and Depth the depth (in cm ) below the surface of the peat sequence from which the core was taken.

## Usage

data(abernethy)

## Format

abernethy is a data frame with 49 samples on 36 species plus sample Age and Depth (in cm).

## Source

These data were provided in electronic format by Prof. H. John B. Birks. The original source is Birks and Mathewes (1978).

## References

Birks, H.H. and Mathewes, R.W. (1978) Studies in the vegetational history of Scotland. New Phytologist 80, 455-484.

## Examples

```
data(abernethy)
head(abernethy)
(plt <- Stratiplot(Age ~ . - Depth,
    data = chooseTaxa(abernethy, n.occ = 5, max.abun = 10),
    type = "poly"))
```

analog Analogue matching

## Description

Analogue matching is a more general implementation of the modern analogue methodology than MAT, where we are only interested in identifying sufficiently similar samples from a modern training as being suitable modern analogues for one or more fossil samples.

## Usage

```
analog(x, ...)
    ## Default S3 method:
    analog(x, y,
            method = c("euclidean", "SQeuclidean", "chord", "SQchord",
                            "bray", "chi.square", "SQchi.square",
                            "information", "chi.distance", "manhattan",
                            "kendall", "gower", "alt.gower", "mixed"),
        keep.train = TRUE, ...)
    ## S3 method for class 'distance'
    analog(x, train = NULL, keep.train = TRUE, ...)
```


## Arguments

$x, y \quad$ data frames with same columns. $x$ is training data and $y$, the test data.
method character string naming the dissimilarity methods to be used. See Details below.
keep.train logical; should the dissimilarity matrix for the training set be stored?
train a pre-computed dissimilarity matrix for the training set samples. Objects of classes "dist", "vegdist", and "distance" are currently accepted.
... arguments passed to or from other methods.

## Details

analog implements analogue matching sensu Flower et al (1997) and Simpson et al (2005), where the aim is to identify suitable close analogues of fossil samples from a modern training set. These results are generally used within ecological restoration, but the identification of close modern analogues for fossil samples is also used as a technique for assessing transfer function reconstructions.
analog is a simple and very general function that generates a pairwise dissimilarity matrix for the modern training set, and a second matrix containing the pairwise dissimilarities between each fossil sample and each sample in the training set. These results can then be assessed using other functions and to extract the close modern analogues using function cma. See the See Also section below.

Analysis of the pairwise dissimilarity matrix for the modern training set can be used to decide on a suitable dissimilarity threshold for defining close modern analogues. By default this matrix is returned as part of the output from the analog function.

## Value

A list of class "analog" with the following components:

| analogs | matrix of pairwise dissimilarities between each fossil sample $(\mathrm{y})$ and each sam- <br> ple in the modern training set $(\mathrm{x})$. |
| :--- | :--- |
| train | if argument keep. train is TRUE then a pairwise dissimilarity matrix for the <br> modern training set. |
| call | the matched function call. |
| method | character; the dissimilarity coefficient used. |

## Author(s)

Gavin L. Simpson

## References

Flower, R.J., Juggins, S. and Battarbee, R.W. (1997) Matching diatom assemblages in lake sediment cores and modern surface sediment samples: the implications for lake conservation and restoration with special reference to acidified systems. Hydrobiologia 344; 27-40.
Simpson, G.L., Shilland, E.M., Winterbottom, J. M. and Keay, J. (2005) Defining reference conditions for acidified waters using a modern analogue approach. Environmental Pollution 137; 119133.

## See Also

distance for the function that calculates the dissimilarity matrices. cma for extraction of close modern analogues. dissimilarities and plot.dissimilarities for analysis of distribution of pairwise dissimilarity matrix for modern training set.

## Examples

```
## Imbrie and Kipp example
## load the example data
data(ImbrieKipp)
data(SumSST)
data(V12.122)
## merge training and test set on columns
dat <- join(ImbrieKipp, V12.122, verbose = TRUE)
## extract the merged data sets and convert to proportions
ImbrieKipp <- dat[[1]] / 100
V12.122 <- dat[[2]] / 100
## Imbrie and Kipp foraminfera sea-surface temperature
## analog matching between SWAP and RLGH core
ik.analog <- analog(ImbrieKipp, V12.122, method = "chord")
ik.analog
summary(ik.analog)
```

```
## Can take pre-computed dissimilarity objects
d1 <- distance(ImbrieKipp, V12.122)
d2 <- distance(ImbrieKipp)
ik <- analog(d1, d2, keep.train = TRUE)
ik
```

bayesF Bayes factors

## Description

Calculates Bayes factors or likelihood ratios of analogue and no-analogue results.

## Usage

```
bayesF(x, prior = rep(0.5, 2))
## S3 method for class 'bayesF'
plot(x, group = "all", xlab = NULL, ylab = "Pr (A+ | d)",
        col = "red", abline.col = "lightgrey", abline.lty = "dashed", ...)
```


## Arguments

x
prior numeric; the prior probabilities of analogue and no-analogue, provided as a vector of length 2 whose elements sum to 1 . If not provided, the function will use the relative occurences of analogue and no analogue situations used to evaluate the ROC curve.
group character vector of length 1 giving the name of the group to plot, or "all" to plot all groups in x .
$x l a b, y l a b \quad$ the $x$ - and $y$-axis labels for the plot.
col colour of the line used to draw the posterior probability.
abline.col colour of the vertical line drawn to indicate the optimal dissimilarity determined from the ROC curve.
abline.lty Line type for indicator of optimal ROC dissimilarity threshold. See par for the allowed line types.
... other plot arguments passed to plotting functions. Currently ignored.

## Details

$\mathrm{LR}(+)$, is the likelihood ratio of a positive test result, that the value of $d$ assigns the sample to the group it belongs to. LR(-) is the likelihood ratio of a negative test result, that the value of $d$ assigns the sample to the wrong group.
$\operatorname{LR}(+)$ is defined as $L R(+)=T P F / F P F$ (or sensitivity / ( $1-$ specificity)), and $\operatorname{LR}(-)$ is defined as $L R(-)=F P F / T N F$ (or (1- sensitivity) / specificity), in Henderson (1993).
The posterior probability of analogue given a dissimilarity is the $\operatorname{LR}(+)$ likelihood ratio values multiplied by the prior odds of analogue, for given values of the dissimilarity, and is then converted to a probability.
The plotting function currently only draws the posterior probability of analogue based on the Bayes factor or likelihood ratio of a positive event (analogue).

## Value

For plot.bayesF a plot on the currently active device.
For bayesF, a list containing the results of computing Bayes factors for each group in x. Each component of this list is itself a list with the following components:
bayesF, posterior.odds, posterior.probs, prior.prob
Bayes factors, posterior odds and probabilities and prior probabilities of true analogue and true non-analogue events. Each components is a list with two components; pos (for true analogue events) and neg (for true non-analogue events). The components of prior. prob are vectors of length 1 , whilst components of the other lists are numeric vectors.
roc.points numeric; the points at which the ROC curve was evaluated.
optimal numeric; the optimal dissimilarity as assessed by the ROC curve.
max. roc numeric; the position along the ROC curve at which the slope of the ROC curve is maximal. This is the index of this point on the curve, and can be used to extract the element of bayesF, posterior.odds and posterior.probs for the optimal dissimilarity.

## Author(s)

Gavin L. Simpson

## References

Brown, C.D., and Davis, H.T. (2006) Receiver operating characteristics curves and related decision measures: A tutorial. Chemometrics and Intelligent Laboratory Systems 80, 24-38.
Gavin, D.G., Oswald, W.W., Wahl, E.R. and Williams, J.W. (2003) A statistical approach to evaluating distance metrics and analog assignments for pollen records. Quaternary Research 60, 356-367.
Henderson, A.R. (1993) Assessing test accuracy and its clinical consequences: a primer for receiver operating characteristic curve analysis. Annals of Clinical Biochemistry 30, 834-846.

## See Also

roc and plot. bayesF.

## Examples

```
## load the example data
data(swapdiat, swappH, rlgh)
## merge training and test set on columns
dat <- join(swapdiat, rlgh, verbose = TRUE)
## extract the merged data sets and convert to proportions
swapdiat <- dat[[1]] / 100
rlgh <- dat[[2]] / 100
## fit an analogue matching (AM) model using the squared chord distance
## measure - need to keep the training set dissimilarities
swap.ana <- analog(swapdiat, rlgh, method = "SQchord",
    keep.train = TRUE)
## fit the ROC curve to the SWAP diatom data using the AM results
## Generate a grouping for the SWAP lakes
METHOD <- if (getRversion() < "3.1.0") {"ward"} else {"ward.D"}
clust <- hclust(as.dist(swap.ana$train), method = METHOD)
grps <- cutree(clust, 12)
## fit the ROC curve
swap.roc <- roc(swap.ana, groups = grps)
swap.roc
## calculate the Bayes factors of analogue and no-analogue
## (uses observed probabilities of analogue/no-analogue
swap.bayes <- bayesF(swap.roc)
swap.bayes
## plot the probability of analogue
plot(swap.bayes)
## Not run:
## calculate the Bayes factors of analogue and no-analogue
## with prior probabilities c(0.5, 0.05)
swap.bayes2 <- bayesF(swap.roc, prior = c(0.5, 0.05))
swap.bayes
## plot the probability of analogue
plot(swap.bayes2)
## End(Not run)
```


## Description

Function to calculate bootstrap statistics for transfer function models such as bootstrap estimates, model RMSEP, sample specific errors for predictions and summary statistics such as bias and $R^{2}$ between oberved and estimated environment.

```
residuals method for objects of class "bootstrap.mat".
```


## Usage

```
bootstrap(object, ...)
## Default S3 method:
bootstrap(object, ...)
## S3 method for class 'mat'
bootstrap(object, newdata, newenv, k,
        weighted = FALSE, n.boot = 1000, ...)
## S3 method for class 'bootstrap.mat'
fitted(object, k, ...)
## S3 method for class 'bootstrap.mat'
residuals(object, which = c("model", "bootstrap"), ...)
```


## Arguments

object
newdata a data frame containing samples for which bootstrap predictions and sample specific errors are to be generated. May be missing - See Details. "newdata" must have the same number of columns as the training set data.
newenv a vector containing environmental data for samples in "newdata". Used to calculate full suite of errors for new data such as a test set with known environmental values. May be missing - See Details. "newenv" must have the same number of rows as "newdata".
k
weighted logical; should the weighted mean of the environment for the " $k$ " modern analogues be used instead of the mean?
n. boot Number of bootstrap samples to take.
which character; which set of residuals to return, the model residuals or the residuals of the bootstrap-derived estimates?
... arguments passed to other methods.

## Details

bootstrap is a fairly flexible function, and can be called with or without arguments newdata and newenv.

If called with only object specified, then bootstrap estimates for the training set data are returned. In this case, the returned object will not include component predictions.
If called with both object and newdata, then in addition to the above, bootstrap estimates for the new samples are also calculated and returned. In this case, component predictions will contain the apparent and bootstrap derived predictions and sample-specific errors for the new samples.

If called with object, newdata and newenv, then the full bootstrap object is returned (as described in the Value section below). With environmental data now available for the new samples, residuals, RMSE(P) and $R^{2}$ and bias statistics can be calculated.
The individual components of predictions are the same as those described in the components relating to the training set data. For example, returned. object\$predictions\$bootstrap contains the components as returned. object\$bootstrap.

It is not usual for environmental data to be available for the new samples for which predictions are required. In normal palaeolimnological studies, it is more likely that newenv will not be available as we are dealing with sediment core samples from the past for which environmental data are not available. However, if sufficient training set samples are available to justify producing a training and a test set, then newenv will be available, and bootstrap can accomodate this extra information and calculate apparent and bootstrap estimates for the test set, allowing an independent assessment of the RMSEP of the model to be performed.
Typical usage of residuals is

```
resid(object, which = c("model", "bootstrap"), \dots)
```


## Value

For bootstrap.mat an object of class "bootstrap.mat" is returned. This is a complex object with many components and is described in bootstrapObject.

For residuals, a list containg the requested residuals and metadata, with the following components:
model Leave one out residuals for the MAT-estimated model.
bootstrap residuals for the bootstrapped MAT model.
k
n. boot numeric; the number of bootstrap samples taken.
auto logical; whether " $k$ " was choosen automatically or user-selected.
weighted logical; whether the weighted mean was used instead of the mean of the environment for $k$-closest analogues.

## Author(s)

Gavin L. Simpson

## References

Birks, H.J.B., Line, J.M., Juggins, S., Stevenson, A.C. and ter Braak, C.J.F. (1990). Diatoms and pH reconstruction. Philosophical Transactions of the Royal Society of London; Series B, 327; 263-278.

## See Also

```
mat, plot.mat, summary.bootstrap.mat, residuals
```


## Examples

```
## Imbrie and Kipp example
## load the example data
data(ImbrieKipp)
data(SumSST)
data(V12.122)
## merge training and test set on columns
dat <- join(ImbrieKipp, V12.122, verbose = TRUE)
## extract the merged data sets and convert to proportions
ImbrieKipp <- dat[[1]] / 100
V12.122 <- dat[[2]] / 100
## Imbrie and Kipp foraminfera sea-surface temperature
## fit the MAT model using the squared chord distance measure
ik.mat <- mat(ImbrieKipp, SumSST, method = "SQchord")
## bootstrap training set
## IGNORE_RDIFF_BEGIN
ik.boot <- bootstrap(ik.mat, n.boot = 100)
ik.boot
summary(ik.boot)
## IGNORE_RDIFF_END
## Bootstrap fitted values for training set
## IGNORE_RDIFF_BEGIN
fitted(ik.boot)
## IGNORE_RDIFF_END
## residuals
resid(ik.boot) # uses abbreviated form
```


## Description

Function to calculate bootstrap statistics for transfer function models such as bootstrap estimates, model RMSEP, sample specific errors for predictions and summary statistics such as bias and $R^{2}$ between oberved and estimated environment.

## Usage

\#\# S3 method for class 'wa'
bootstrap(object, n.boot $=1000$, verbose $=$ TRUE,.. )

## Arguments

object an R object of class "wa" for which bootstrap statistics are to be generated.
n. boot numeric; the number of bootstrap samples to draw.
verbose logical; should bootstrap progress be printed to the console?
... arguments passed to other methods.

## Details

See bootstrap.mat for further details. This method is not as feature packed as bootstrap.mat but can be used to evaluate the model performance of WA transfer function models.

Value
An object with the same components as predict.wa.

## Author(s)

Gavin L. Simpson

## References

Birks, H.J.B., Line, J.M., Juggins, S., Stevenson, A.C. and ter Braak, C.J.F. (1990). Diatoms and pH reconstruction. Philosophical Transactions of the Royal Society of London; Series B, 327; 263-278.

## See Also

wa, plot.wa.

## Examples

```
## Imbrie and Kipp
data(ImbrieKipp)
data(SumSST)
ik.wa <- wa(SumSST ~ ., data = ImbrieKipp, tol.dw = TRUE,
                    min.tol = 2, small.tol = "min")
ik.wa
## compare actual tolerances to working values
```

```
with(ik.wa, rbind(tolerances, model.tol))
## bootstrap the WA model
ik.boot <- bootstrap(ik.wa, n.boot = 100)
## performance statistics
performance(ik.boot)
```

bootstrapObject Bootstrap object description

## Description

Objects of class bootstrap.mat are a complex containing many sub-components. This object is described here in more detail.

## Details

A large object is returned with some or all of the following depending on whether newdata and newenv are supplied or not.
observed: vector of observed environmental values.
model: a list containing the apparent or non-bootstrapped estimates for the training set. With the following components:
estimated: estimated values for the response
residuals: model residuals
r.squared: Apparent $R^{2}$ between observed and estimated values of response
avg.bias: Average bias of the model residuals
max.bias: Maximum bias of the model residuals
rmse: Apparent error (RMSE) for the model.
k : numeric; indicating the size of model used in estimates and predictions
bootstrap: a list containing the bootstrap estimates for the training set. With the following components:
estimated: Bootstrap estimates for the response
residuals: Bootstrap residuals for the response
r.squared: Bootstrap derived $R^{2}$ between observed and estimated values of the response
avg.bias: Average bias of the bootstrap derived model residuals
max.bias: Maximum bias of the bootstrap derived model residuals
rmsep: Bootstrap derived RMSEP for the model
s1: Bootstrap derived S1 error component for the model
s2: Bootstrap derived S2 error component for the model
k : numeric; indicating the size of model used in estimates and predictions
sample.errors: a list containing the bootstrap-derived sample specific errors for the training set. With the following components:
rmsep: Bootstrap derived RMSEP for the training set samples
s1: Bootstrap derived S1 error component for training set samples
s2: Bootstrap derived S2 error component for training set samples
weighted: logical; whether the weighted mean was used instead of the mean of the environment for $k$-closest analogues
auto: logical; whether " $k$ " was choosen automatically or user-selected
n . boot: numeric; the number of bootstrap samples taken
call: the matched call
type: model type
predictions: a list containing the apparent and bootstrap-derived estimates for the new data, with the following components:
observed: the observed values for the new samples - only if newenv is provided
model: a list containing the apparent or non-bootstrapped estimates for the new samples. A list with the same components as model, above
bootstrap: a list containing the bootstrap estimates for the new samples, with some or all of the same components as bootstrap, above
sample.errors: a list containing the bootstrap-derived sample specific errors for the new samples, with some or all of the same components as sample.errors, above

## Author(s)

Gavin L. Simpson

## See Also

mat, plot.mat, summary.bootstrap.mat, residuals

$$
\text { caterpillarPlot } \quad \text { Caterpillar plot of species' WA optima and tolerance range. }
$$

## Description

Draws a caterpillar plot of the weighted average optima and tolerance range for each of the species in a data set.

## Usage

```
## Default S3 method:
caterpillarPlot(x, tol, mult = 1, decreasing = TRUE,
    labels, xlab = NULL, pch = 21, bg = "white",
    col = "black", lcol = col, lwd = 2, frame.plot = FALSE, ...)
    ## S3 method for class 'data.frame'
    caterpillarPlot(x, env, useN2 = TRUE, xlab, ...)
    ## S3 method for class 'wa'
    caterpillarPlot(x, type = c("observed","model"), ...)
```


## Arguments

X
tol numeric; vector of species tolerances.
env numeric; variable for which optima and tolerances are required.
useN2
decreasing
mult
labels
xlab
$\mathrm{pch}, \mathrm{bg}, \mathrm{col}$ The plotting character to use and its background and foreground colour. See par.
lcol, lwd The colour and line width to use for the tolerance range.
type character; "observed" uses the actual tolerances observed from the data. "model" uses the tolerances used in the WA model where very small tolerances have been reset for some definition of small.
frame.plot logical; should a box be drawn round the plot?
Additional graphical arguments to be passed on to plotting functions.

## Details

The function may also be called using the short form name caterpillar:

```
caterpillar(x, ...)
```


## Value

The function results in a plot on the currently active device. A data frame with components Optima and Tolerance is returned invisibly.

## Author(s)

Gavin L. Simpson

## See Also

For the underlying computations optima and tolerance.

## Examples

```
data(ImbrieKipp)
data(SumSST)
## default plot
caterpillar(ImbrieKipp, SumSST)
## customisation
opttol <-
        caterpillar(ImbrieKipp, SumSST, col = "red2",
            bg = "yellow", lcol = "blue",
            xlab = expression(Summer ~ Sea ~ Surface ~
                                    Temperature~(degree*C)))
## invisibly returns the optima and tolerances
head(opttol)
```

chooseTaxa Select taxa (variables) on basis of maximum abundance attained and number of occurrences.

## Description

Select taxa (variables) from an object on the basis of one or both of maximum abundance and number of occurrences greater than user-specified values. This is a simple utility function to encapsulate this common task in filtering palaeoecological data sets.

## Usage

```
chooseTaxa(object, ...)
## Default S3 method:
chooseTaxa(object, n.occ = 1, max.abun = 0,
    type = c("AND","OR"), value = TRUE, na.rm = FALSE,
    ...)
```


## Arguments

object an R object for which a suitable method exists. The default method assumes a matrix-like object such as a data frame or a numeric matrix.
n. occ numeric; number of occurrences representing the lower limit for selection. A taxon is included in the returned subset if it is present a total of n.occ times or more. See argument type for a modifier which might exclude the taxon even if it would be included on the basis of $n$.occ.
max.abun numeric; maximum abundance representing the lower limit for selection. A taxon is included in the returned subset if it attains abundance equal to or greater than max. abun in one or more sample. See argument type for a modifier which might exclude the taxon even if it would be included on the basis of max. abun.

| type | character; one of "AND" or "OR", controlling how the criteria n. occ and max. abun <br> are combined to generate a subset of the variables in object. |
| :--- | :--- |
| value | logical; should the data for the selected taxa be returned? If TRUE, the default, <br> the data for the chosen taxa are returned. If FALSE, a logical vector is returned, <br> indicating which taxa met the selection criteria. |
| na.rm | logical; should missing values NAs be excluded from the calculation of abun- <br> dances and occurrence? |
| $\ldots$ | arguments passed on to subsequent methods. |

## Value

If value $=$ TRUE, returns the supplied data frame or matrix with a subset of columns (taxa) that meet the criteria chosen. If value $=$ FALSE, a logical vector is returned.

## Author(s)

Gavin L. Simpson

## Examples

```
data(ImbrieKipp)
IK2 <- chooseTaxa(ImbrieKipp, n.occ = 5)
dim(ImbrieKipp)
dim(IK2)
## return a logical vector to select species/columns
chooseTaxa(ImbrieKipp, n.occ = 5, value = FALSE)
```

    cma Close modern analogues
    
## Description

Extracts and formats close modern analogue samples from a modern reference set that are closer than a defined cut off threshold.

## Usage

cma(object, ...)
\#\# Default S3 method:
cma(object, ...)
\#\# S3 method for class 'analog'
cma(object, cutoff, prob = c(0.01, 0.025, 0.05), ...)
\#\# S3 method for class 'mat'

```
cma(object, k, cutoff, prob = c(0.01, 0.025, 0.05), ...)
## S3 method for class 'predict.mat'
cma(object, k, cutoff, prob = c(0.01, 0.025,
0.05), ...)
## S3 method for class 'cma'
plot(x, method = c("overplot", "jitter", "stack"),
    jitter = 0.1, vertical = FALSE,
    draw.quant = TRUE, xlab = NULL, ylab = "",
    main = "", cex.axis = NULL, ...,
    col.quant = "red", lty.quant= "dashed")
```


## Arguments

object an object for which close modern analogues are to be returned. Currently only for objects of class analog.

| k | numeric; the number of analogues to return. |
| :---: | :---: |
| cutoff | numeric; critical value determining level below which samples from the modern reference set are defined as close modern analogues. May be missing, in which case the $2.5 \%$ quantile of the training set dissimilarities is used unless object\$train is NULL, in which case "cutoff" must be supplied. |
| prob | numeric vector of probabilities with values in [0,1], for which quantiles of the distribution of training set dissimilarities will be calculated. See quantile. |
|  | arguments to be passed to other cma methods or additional arguments passed to stripchart. |
| x | an object of class "cma". |
| method | the method to be used to separate coincident points. The default method "overplot" causes such points to be overplotted, but it is also possible to specify "jitter" to jitter the points, or "stack" have coincident points stacked. The last method only makes sense for very granular data. |
| jitter | when method="jitter" is used, jitter gives the amount of jittering applied. |
| vertical | when vertical is TRUE the plots are drawn vertically rather than the default horizontal. |
| draw.quant | logical; should the quantiles be drawn on the stripchart? |
| xlab,ylab, main | Graphical parameters |
| cex.axis | The magnification to be used for axis annotation relative to the current setting of cex. See par. |
| col.quant,lty.quant |  |
|  | lour and line type in which |

## Details

The plot method is simply a wrapper to stripchart.
The methods for mat and predict.mat objects allow the user to select the k-closest analogues (argument k) or those samples as close or closer than a stated threshold of dissimilarity (argument
cutoff). Only one of $k$ and cutoff may be specified. If neither is specified, getK is used to extract the value for $k$ stored within object. As such, the default is to return the automatically selected set of k closest samples, behaviour that is consistent with other functions in the package.

## Value

For the plot method, a plot on the current device. Invisibly the plotted data are returned; see Note for further details.
A list of class "cma" with the following components:

$$
\begin{array}{ll}
\text { close } & \begin{array}{l}
\text { a named list of named vectors of close modern analogues and their dissimilari- } \\
\text { ties. The names of the list components are the names of the fossil samples. The } \\
\text { named vector in each component of close is the distances for the close modern } \\
\text { analogues from the training set that are as close as cutoff, or closer, to the fossil } \\
\text { sample. }
\end{array} \\
\text { call } & \text { the matched call. } \\
\text { cutoff } & \text { the cutoff threshold used to define close modern analogues. } \\
\text { quant } & \begin{array}{l}
\text { numeric vector of the requested quantiles. Note returned by the predict.mat } \\
\text { method. }
\end{array} \\
\text { probs } & \text { the probabilities of the requested quantiles. } \\
\text { method } & \text { character; the dissimilarity coefficient used } \\
\text { n. analogs } & \text { numeric vector of the number of analogues per fossil sample. }
\end{array}
$$

## Note

Only objects of classes analog, mat, and predict.mat are supported.
The plot method invisibly returns a list with the following components:

- distancesa vector of stacked distances extracted from object.
- groupsa factor listing the fossil sample for which the distances are the distances to the close modern analogues for the training set.


## Author(s)

Gavin L. Simpson

## References

Flower, R.J., Juggins, S. and Battarbee, R.W. (1997) Matching diatom assemblages in lake sediment cores and modern surface sediment samples: the implications for lake conservation and restoration with special reference to acidified systems. Hydrobiologia 344; 27-40.
Simpson, G.L., Shilland, E.M., Winterbottom, J. M. and Keay, J. (2005) Defining reference conditions for acidified waters using a modern analogue approach. Environmental Pollution 137; 119133.

## See Also

analog, stripchart, or boxplot for an alternative representation.

## Examples

```
## Imbrie and Kipp example
## load the example data
data(ImbrieKipp)
data(SumSST)
data(V12.122)
## merge training and test set on columns
dat <- join(ImbrieKipp, V12.122, verbose = TRUE)
## extract the merged data sets and convert to proportions
ImbrieKipp <- dat[[1]] / 100
V12.122 <- dat[[2]] / 100
## analog matching between SWAP and RLGH reference samples
(ik.ana <- analog(ImbrieKipp, V12.122, method = "chord"))
## close modern analogues
(ik.cma <- cma(ik.ana, cutoff = 0.4))
summary(ik.cma)
## plot the results
plot(ik.cma)
```

```
compare Compare proxies across two data sets
```


## Description

compare() compares a proxy dataset with a training set or other data set that is considered to be the master. A range of metrics is returned, currently for samples only.

## Usage

```
compare(x, ...)
## Default S3 method:
compare(x, y, env,
        by = c("sites", "species"),
        ordination = "rda",
        method = "chord",
        transform = NULL,
        n2limit = 5L,
        ...)
```


## Arguments

x
y
env
by character; compare data sets by sites or species (proxies).
data frame; training set samples to compare against
data frame; passive or core samples
numeric vector of environmental or contraint data for residual length ordination. Ignored if by = "species".
ordination
method
transform
n2limit
character; which constrained ordination method to use character; which dissimilarity method to use. See distance. character: should a transformation be applied to the data. Ignored. integer; the criterion for indicating species with potentially poorly estimated optima. The default value of 5 L is one suggested by R. Telford.
... arguments passed to other methods.

## Details

ToDo

## Value

If by = "species" a data frame of diagnostics for each species (proxy) in y relative to x . If by $=$ "sites", the diagnostics are for each sample (row) in y. Depending on the value of by some of the following columns will be returned

| sumMissing | numeric; abundance sum for species missing from the training set x . |
| :---: | :---: |
| sumPoorOpt | numeric; abundance sum for species with potentially poorly estimated optima. |
| closestSamp | numeric; minimum dissimilarity to a sample in the training data x . |
| residLen | numeric; the squared residual length for each sample in $y$. A measure of how well the sample fits into the species-environment relationship of a constrained ordination. See residLen for details. Not given if env is not provided. |
| inTrain | logical; simple indicator of whether a species in $y$ is present in the training data x. |
| n2 | numeric; Hill's N2 for each species in y . |
| n2Train | numeric; as for n 2 but computed from the training data x . |
| max | numeric; the maximum abundance of each species computed using $y$. |
| maxTrain | numeric; as for max but computed using the training data x . |

## Author(s)

Gavin L. Simpson

## Examples

```
data(ImbrieKipp, V12.122, SumSST)
compare(ImbrieKipp, V12.122, env = SumSST, ordination = "rda",
    method = "chord")
```


## Description

Performs leave-one-out, $k$-fold, $n k$-fold and bootstrap cross-validation of palaeoecological transfer function models.

## Usage

```
    crossval(obj, ...)
    ## S3 method for class 'wa'
    crossval(obj, method = c("LOO","kfold","bootstrap"),
        nboot = 100, nfold = 10, folds = 5,
        verbose = getOption("verbose"), ...)
    ## S3 method for class 'pcr'
    crossval(obj, method = c("LOO","kfold","bootstrap"),
        ncomp, nboot = 100, nfold = 10, folds = 5,
        verbose = getOption("verbose"), ...)
```


## Arguments

obj A fitted transfer function model. Currently, only objects of class wa and per are supported.
method character; type of cross-validation.
ncomp numeric; number of components to fit, as in models with 1 : ncomp components.
nboot numeric; number of bootstrap samples.
nfold numeric; number of chunks into which the training data are split. The $k$ in $k$-fold.
folds numeric; the number of times $k$-fold CV is performed.
verbose logical; should progress of the CV be displayed?
... Arguments passed to other methods.

## Value

Returns an object of class "crossval", a list with the following components:
fitted.values numeric vector; the cross-validated estimates of the response.
residuals numeric vector; residuals computed from the cross-validated estimates of the response.
performance data frame; cross-validation performance statistics for the model.
CVparams list; parameters holding details of the cross-validation process.
call the matched call.
densityplot.residLen

## Author(s)

Gavin L. Simpson

## See Also

## wa

## Examples

```
## Load the Imbrie & Kipp data and
## summer sea-surface temperatures
data(ImbrieKipp)
data(SumSST)
## fit the WA model
mod <- wa(SumSST ~., data = ImbrieKipp)
mod
## Leave one out CV
cv.loo <- crossval(mod)
cv.loo
## k-fold CV (k == 10)
cv.kfold <- crossval(mod, method = "kfold", kfold = 10, folds = 1)
cv.kfold
## n k-fold CV (k == 10, n = 10)
cv.nkfold <- crossval(mod, method = "kfold", kfold = 10, folds = 10)
cv.nkfold
## bootstrap with 100 bootstrap samples
cv.boot <- crossval(mod, method = "bootstrap", nboot = 100)
cv.boot
## extract fitted values and residuals
fitted(cv.boot)
resid(cv.boot)
## Principal Components Regression
mpcr <- pcr(SumSST ~., data = ImbrieKipp, ncomp = 10)
crossval(mpcr, method = "kfold", kfold = 10, folds = 2, ncomp = 10)
crossval(mpcr, method = "bootstrap", nboot = 100, ncomp = 10)
```

densityplot.residLen Lattice density plot for residual lengths

## Description

Lattice densityplot method for residLen objects.

## Usage

```
## S3 method for class 'residLen'
densityplot(x, ..., xlab = NULL, ylab = NULL)
```


## Arguments

$x \quad$ Object of class "residLen", the result of a call to residLen.
$x l a b, y l a b \quad$ Axis labels. If not supplied, suitable defaults are generated, depending on whether RDA or CCA was used as the underlying ordination model.
... Additional arguments passed to densityplot.

## Value

Returns an object of class "trellis". See densityplot for details.

## Author(s)

Gavin L. Simpson

## See Also

residLen, plot.residLen, hist.residLen, histogram.residLen.

## Examples

```
## load the Imbrie and Kipp example data
data(ImbrieKipp, SumSST, V12.122)
## squared residual lengths for Core V12.122
rlens <- residLen(ImbrieKipp, SumSST, V12.122)
rlens
## plot the density functions of the residual distances
densityplot(rlens)
```

    deshrink Deshrinking techniques for WA transfer functions
    
## Description

In Weighted Averaging models averages are taken twice and thus WA estimates shrink towards the training set mean and need to be deshrunk.deshrink performs this deshrinking using several techniques, whilst deshrinkPred will deshrink WA estimates for new samples given a set of deshrinking coefficients.

## Usage

```
deshrink(env, wa.env,
            type = c("inverse", "classical", "expanded", "none",
                            "monotonic"))
deshrinkPred(x, coef,
            type = c("inverse", "classical", "expanded", "none",
                    "monotonic"))
```


## Arguments

env numeric; original environmental values.
wa.env numeric; initial weighted average estimates.
type character; the type of deshrinking. One of "inverse", "classical", "expand", "none".
$x \quad$ numeric; estimates to be deshrunk.
coef numeric; deshrinking coefficients to use. Currently needs to be a vector of length 2. These should be supplied in the order $\beta_{0}, \beta_{1}$.

## Value

For deshrinkPred a numeric vector of deshrunk estimates.
For an object of class "deshrink", inheriting from class "list", with two components. The type of deshrinking performed is stroed within attribute "type". The componets of the returned object are:
coefficients The deshrinking coefficients used.
env The deshrunk WA estimates.

## Warning

deshrinkPred, does not currently check that the correct coefficients have been supplied in the correct order.

## Author(s)

Gavin L. Simpson <br>\& Jari Oksanen

## References

Birks, H.J.B. (1995) Quantitative environmental reconstructions. In Statistical modelling of Quaternary science data (eds.~D.Maddy <br>\& J.S. Brew). Quaternary Research Association technical guide 5. Quaternary Research Association, Cambridge.

## See Also

## Description

Extracts a vector of dissimilarity coefficients from an object for further analysis.

## Usage

```
dissimilarities(object, ...)
dissim(object, ...)
## S3 method for class 'analog'
dissimilarities(object, which = c("train", "analogs"),
    ...)
    ## S3 method for class 'mat'
    dissimilarities(object, ...)
```


## Arguments

object an R object from which the dissimilarity values are to be extracted. Currently only for objects of class "analog".
which character; which set of dissimilarities should be extracted. One of "train" or "analogs".
$\ldots \quad$ arguments passed to other methods.

## Details

The function can be called using the much shorter name "dissim".

## Value

A vector of dissimilarities.

## Author(s)

Gavin L. Simpson

## See Also

analog, plot.dissimilarities

## Examples

```
## Imbrie and Kipp example
## load the example data
data(ImbrieKipp)
data(SumSST)
data(V12.122)
## merge training and test set on columns
dat <- join(ImbrieKipp, V12.122, verbose = TRUE)
## extract the merged data sets and convert to proportions
ImbrieKipp <- dat[[1]] / 100
V12.122 <- dat[[2]] / 100
## analog matching between SWAPImbrie & Kipp and V12.122 core
ik.analog <- analog(ImbrieKipp, V12.122, method = "chord")
ik.analog
summary(ik.analog)
## compare training set dissimilarities with normals
## and derive cut-offs
ik.dij <- dissim(ik.analog)
plot(ik.dij)
```

```
distance
Flexibly calculate dissimilarity or distance measures
```


## Description

Flexibly calculates distance or dissimilarity measures between a training set $x$ and a fossil or test set $y$. If $y$ is not supplied then the pairwise dissimilarities between samples in the training set, $x$, are calculated.

## Usage

```
distance(x, ...)
## Default S3 method:
distance(x, y, method = "euclidean", weights = NULL,
    R = NULL, dist = FALSE, double.zero = FALSE, ...)
## S3 method for class 'join'
distance(x, ...)
oldDistance(x, ...)
## Default S3 method:
```

```
oldDistance(x, y, method = c("euclidean", "SQeuclidean",
    "chord", "SQchord", "bray", "chi.square",
    "SQchi.square", "information", "chi.distance",
    "manhattan", "kendall", "gower", "alt.gower",
    "mixed"),
    fast = TRUE,
    weights = NULL, R = NULL, ...)
## S3 method for class 'join'
oldDistance(x, ...)
```


## Arguments

| x | data frame or matrix containing the training set samples, or and object of class join. |
| :---: | :---: |
| y | data frame or matrix containing the fossil or test set samples. |
| method | character; which choice of dissimilarity coefficient to use. One of the listed options. See Details below. |
| weights | numeric; vector of weights for each descriptor. |
| R | numeric; vector of ranges for each descriptor. |
| dist | logical; should the dissimilarity matrix be returned as an object of class "dist"? Ignored if y is supplied. |
| double.zero | logical; if FALSE, the default, double zeroes are not counted in the distance calculation. If TRUE, absences of a variable in both samples counts as a similarity between the two samples. Currently this only affects methods "mixed" and "metric.mixed" forms of Gower's general coefficient. |
| fast | logical; should fast versions of the dissimilarities be calculated? See details below. |
|  | arguments passed to other methods |

## Details

A range of dissimilarity coefficients can be used to calculate dissimilarity between samples. The following are currently available:

```
euclidean \(\quad d_{j k}=\sqrt{\sum_{i}\left(x_{i j}-x_{i k}\right)^{2}}\)
SQeuclidean \(\quad d_{j k}=\sum_{i}\left(x_{i j}-x_{i k}\right)^{2}\)
chord \(\quad d_{j k}=\sqrt{\sum_{i}\left(\sqrt{x_{i j}}-\sqrt{x_{i k}}\right)^{2}}\)
SQchord \(\quad d_{j k}=\sum_{i}\left(\sqrt{x_{i j}}-\sqrt{x_{i k}}\right)^{2}\)
bray \(\quad d_{j k}=\frac{\sum_{i}\left|x_{i j}-x_{i k}\right|}{\sum_{i}\left(x_{i j}+x_{i k}\right)}\)
chi.square \(\quad d_{j k}=\sqrt{\sum_{i} \frac{\left(x_{i j}-x_{i k}\right)^{2}}{x_{i j}+x_{i k}}}\)
SQchi.square \(d_{j k}=\sum_{i} \frac{\left(x_{i j}-x_{i k}\right)^{2}}{x_{i j}+x_{i k}}\)
information \(\quad d_{j k}=\sum_{i}\left(p_{i j} \log \left(\frac{2 p_{i j}}{p_{i j}+p_{i k}}\right)+p_{i k} \log \left(\frac{2 p_{i k}}{p_{i j}+p_{i k}}\right)\right)\)
chi.distance \(d_{j k}=\sqrt{\sum_{i}\left(x_{i j}-x_{i k}\right)^{2} /\left(x_{i+} / x_{++}\right)}\)
manhattan \(\quad d_{j k}=\sum_{i}\left(\left|x_{i j}-x_{i k}\right|\right)\)
```

kendall
gower
alt.gower
mixed
$d_{j k}=\sum_{i} M A X_{i}-\operatorname{minimum}\left(x_{i j}, x_{i k}\right)$
$d_{j k}=\sum_{i} \frac{\left|p_{i j}-p_{i k}\right|}{R_{i}}$
$d_{j k}=\sqrt{2 \sum_{i} \frac{\left|p_{i j}-p_{i k}\right|}{R_{i}}}$
where $R_{i}$ is the range of proportions for descriptor (variable) $i$
,
$d_{j k}=\frac{\sum_{i=1}^{p} w_{i} s_{j k i}}{\sum_{i=1}^{p} w_{i}}$
where $w_{i}$ is the weight for descriptor $i$ and $s_{j k i}$ is the similarity
between samples $j$ and $k$ for descriptor (variable) $i$.
metric.mixed as for mixed but with ordinal variables converted to ranks and handled as quantitative variables in Gower's n

Argument fast determines whether fast C versions of some of the dissimilarity coefficients are used. The fast versions make use of dist for methods "euclidean", "SQeuclidean", "chord", "SQchord", and vegdist for method == "bray". These fast versions are used only when x is supplied, not when $y$ is also supplied. Future versions of distance will include fast C versions of all the dissimilary coefficients and for cases where $y$ is supplied.

## Value

A matrix of dissimilarities where columns are the samples in $y$ and the rows the samples in $x$. If $y$ is not provided then a square, symmetric matrix of pairwise sample dissimilarities for the training set $x$ is returned, unless argument dist is TRUE, in which case an object of class "dist" is returned. See dist.

The dissimilarity coefficient used (method) is returned as attribute "method". Attribute "type" indicates whether the object was computed on a single data matrix ("symmetric") or across two matrices (i.e. the dissimilarties between the rows of two matrices; "asymmetric".

## Warning

For method = "mixed" it is essential that a factor in $x$ and $y$ have the same levels in the two data frames. Previous versions of analogue would work even if this was not the case, which will have generated incorrect dissimilarities for method = "mixed" for cases where factors for a given species had different levels in $x$ to $y$.
distance now checks for matching levels for each species (column) recorded as a factor. If the factor for any individual species has different levels in $x$ and $y$, an error will be issued.

## Note

The dissimilarities are calculated in native R code. As such, other implementations (see See Also below) will be quicker. This is done for one main reason - it is hoped to allow a user defined function to be supplied as argument "method" to allow for user-extension of the available coefficients.

The other advantage of distance over other implementations, is the simplicity of calculating only the required pairwise sample dissimilarities between each fossil sample ( y ) and each training set sample (x). To do this in other implementations, you would need to merge the two sets of samples, calculate the full dissimilarity matrix and then subset it to achieve similar results.

## Author(s)

Gavin L. Simpson and Jari Oksanen (improvements leading to method "metric.mixed" and proper handling of ordinal data via Podani's (1999) modification of Gower's general coefficient in method "mixed").

## References

Faith, D.P., Minchin, P.R. and Belbin, L. (1987) Compositional dissimilarity as a robust measure of ecological distance. Vegetatio 69, 57-68.
Gavin, D.G., Oswald, W.W., Wahl, E.R. and Williams, J.W. (2003) A statistical approach to evaluating distance metrics and analog assignments for pollen records. Quaternary Research 60, 356-367.
Kendall, D.G. (1970) A mathematical approach to seriation. Philosophical Transactions of the Royal Society of London - Series B 269, 125-135.

Legendre, P. and Legendre, L. (1998) Numerical Ecology, 2nd English Edition. Elsevier Science BV, The Netherlands.
Overpeck, J.T., Webb III, T. and Prentice I.C. (1985) Quantitative interpretation of fossil pollen spectra: dissimilarity coefficients and the method of modern analogues. Quaternary Research 23, 87-108.

Podani, J. (1999) Extending Gower's General Coefficient of Similarity to Ordinal Characters. Taxon 48, 331-340).
Prentice, I.C. (1980) Multidimensional scaling as a research tool in Quaternary palynology: a review of theory and methods. Review of Palaeobiology and Palynology 31, 71-104.

## See Also

vegdist in package vegan, daisy in package cluster, and dist provide comparable functionality for the case of missing $y$.

## Examples

```
## simple example using dummy data
train <- data.frame(matrix(abs(runif(200)), ncol = 10))
rownames(train) <- LETTERS[1:20]
colnames(train) <- as.character(1:10)
fossil <- data.frame(matrix(abs(runif(100)), ncol = 10))
colnames(fossil) <- as.character(1:10)
rownames(fossil) <- letters[1:10]
## calculate distances/dissimilarities between train and fossil
## samples
test <- distance(train, fossil)
## using a different coefficient, chi-square distance
test <- distance(train, fossil, method = "chi.distance")
## calculate pairwise distances/dissimilarities for training
## set samples
test2 <- distance(train)
```

```
## Using distance on an object of class join
dists <- distance(join(train, fossil))
str(dists)
## calculate Gower's general coefficient for mixed data
## first, make a couple of variables factors
## fossil[,4] <- factor(sample(rep(1:4, length = 10), 10))
## train[,4] <- factor(sample(rep(1:4, length = 20), 20))
## ## now fit the mixed coefficient
## test3 <- distance(train, fossil, "mixed")
## ## Example from page 260 of Legendre & Legendre (1998)
x1 <- t(c(2,2,NA, 2, 2,4,2,6))
x2 <- t(c(1,3,3,1,2,2,2,5))
Rj <- c(1,4,2,4,1,3,2,5) # supplied ranges
## 1 - distance(x1, x2, method = "mixed", R = Rj)
## note this gives ~0.66 as Legendre & Legendre describe the
## coefficient as a similarity coefficient. Hence here we do
## 1 - Dij here to get the same answer.
## Tortula example from Podani (1999)
data(tortula)
Dij <- distance(tortula[, -1], method = "mixed") # col 1 includes Taxon ID
## Only one ordered factor
data(mite.env, package = "vegan")
Dij <- distance(mite.env, method = "mixed")
## Some variables are constant
data(BCI.env, package = "vegan")
Dij <- distance(BCI.env, method = "mixed")
```

evenSample

## Description

The number of samples in sections along the gradient is a useful diagnostic as to the quality of reconstructions at gradient values within those sections.

## Usage

evenSample(grad, $\mathrm{n}=10$ )

## Arguments

grad numeric; vector of gradient values
$\mathrm{n} \quad$ number of segments to partition the gradient into

## Details

The sampling design of a training set, i.e. the number of samples taken at points along the gradient, can influence the uncertainty in the transfer function predictions at those values of the gradient. Poorly sampled sections of the gradient may have far larger RMSEP than the overall model RMSEP.

## Value

Numeric vector of length n containing the numbers of samples per gradient segment.

## Author(s)

Gavin L. Simpson

## Examples

```
data(SumSST)
ev <- evenSample(SumSST) ## not an even sample...
plot(ev)
```


## Description

Extracts fitted values for training set samples from logistic regression models fitted to each group of samples that describe the probability two samples are analogues (from the same group) as a function of dissimilarity between the paired samples.

## Usage

\#\# S3 method for class 'logitreg'
fitted(object, combined = FALSE, ...)

## Arguments

object an object of class "logitreg" resulting from a call to logitreg.
combined logical; should the fitted values for the overall combined analysis be returned.
... arguments passed to other methods.

## Value

If combined $==$ FALSE (the default) then a matrix of fitted probabilities, where the rows are the training set samples and the columns the groupings, is returned. If combined $==$ TRUE, then a list with components "group" and "combined". "group" is a matrix of fitted probabilities as above. "combined" is a vector of fitted values for the entire set of pairwise comparisons considered.

## Author(s)

Gavin L. Simpson

## See Also

See logitreg for example usage.
fuse Fused dissimilarities

## Description

Combines dissimilarities from two or more dissimilarity objects into a single dissimilarity object so that both original dissimilarities contribute equally. Weighted combinations of the original objects can also be created.

## Usage

```
fuse(..., weights = NULL)
    ## S3 method for class 'matrix'
    fuse(..., weights = NULL)
    ## S3 method for class 'dist'
    fuse(..., weights = NULL)
```


## Arguments

... objects to fuse. Methods currently exist for objects of class "matrix" and "dist" objects. Method dispatch is performed on the first object specified in .... A minimum of two objects must be supplied.
weights numeric; vector of weights that sum to 1 , one weight for each object supplied in

## Details

Fuses, or combines, dissimilarity objects in a very flexible way to create a single dissimilarity object that incorporates the separate dissimilarities. In analogue matching, we may wish to combine information from two or more proxies, such as diatoms and cladocera, or from biological and chemical or physical data in the case of matching modern samples.
The function can also be used to fuse dissimilarity objects created from a single data set but using different dissimilarity coefficients. In this way one could create a new dissimilarity object combining dissimilarity based on abundance data and presence absence data into a single measure.
fuse uses the method of Melssen et al. (2006) to combine dissimilarities. The dissimilarities in each dissimilarity object are scaled so that the maximum dissimilarity in each object is 1 . The scaled dissimilarity objects are then weighted according to the supplied weights. If no weights are supplied (the default) the dissimilarity objects are weighted equally; weights $=\operatorname{rep}(1 / \mathrm{N}, \mathrm{N})$, where $N$ is the number of dissimilarity objects fused.

$$
D_{\text {fused }}(j, k)=\sum_{i=1}^{N} w_{i} D_{i j k}
$$

where $D_{\text {fused }}(j, k)$ is the fused dissimilarity between samples $j$ and $k, w_{i}$ is the weight assigned to the $i$ th dissimilarity object and $D_{i j k}$ is the dissimilarity between $j$ and $k$ for the $i$ th dissimilarity object.

## Value

fuse returns an object of class "dist" with the attribute "method" set to "fuse".
This is the case even if the supplied objects are full dissimilarity matrices. If you want a full dissimilarity object, use as matrix. dist on the returned object.
The returned object contains an extra attribute "weights", which records the weights used in the fusing.

## Author(s)

Gavin L. Simpson

## References

Melssen W., Wehrens R. and Buydens L. (2006) Supervised Kohonen networks for classification problems. Chemometrics and intelligent laboratory systems 83, 99-113.

## See Also

dist, vegdist, distance.

## Examples

```
train1 <- data.frame(matrix(abs(runif(100)), ncol = 10))
train2 <- data.frame(matrix(sample(c(0,1), 100, replace = TRUE),
    ncol = 10))
rownames(train1) <- rownames(train2) <- LETTERS[1:10]
```

```
    colnames(train1) <- colnames(train2) <- as.character(1:10)
    d1 <- vegdist(train1, method = "bray")
    d2 <- vegdist(train2, method = "jaccard")
    dd <- fuse(d1, d2, weights = c(0.6, 0.4))
    dd
    str(dd)
```

    getK Extract and set the number of analogues
    
## Description

An extractor function to access the number of analogues used in particular models. The stored value of $k$ can be updated using setK.

## Usage

```
getK(object, ...)
## S3 method for class 'mat'
getK(object, weighted = FALSE, ...)
## S3 method for class 'bootstrap.mat'
getK(object, which = c("bootstrap", "model"),
        prediction = FALSE, ...)
## S3 method for class 'predict.mat'
getK(object, which = c("model", "bootstrap"),
    ...)
setK(object, weighted = FALSE) <- value
## S3 replacement method for class 'mat'
setK(object, weighted = FALSE) <- value
```


## Arguments

| object | an R object; currently only for objects of class mat and class bootstrap.mat. |
| :--- | :--- |
| weighted | logical; extract/set number of analogues for a weighted or un-weighted model? <br> character; which $k$ should be extracted, the one from the model or the one from <br> the bootstrap results? |
| prediction | logical; should the extracted $k$ be the one that is minimum for the test set <br> (newdata) or the model (object). |
| $\ldots$ | further arguments to other methods. <br> integer; replacement value for $k$. |

## Details

getK is a generic accessor function, and setK<- is a generic replacement function.
Objects of class bootstrap.mat contain several different k's. If no predictions are performed, there will be two k's, one for the model and one from bootstrapping the model. Where predictions are performed with newenv supplied, in addition to the k's above, there will be two $k$ ' for the predictions, one for the model-based and one for the bootstrap-based predictions. To select $k$ for the predictions, use prediction = TRUE. Argument which determines whether the model-based or the bootstrap-based k is returned.

## Value

For getK, an integer value that is the number of analogues stored for use. The returned object has attributes "auto" and "weighted". "auto" refers to whether the extracted value of $k$ was set automatically (TRUE) or by the user (FALSE). "weighted" states if the returned value is for a weighted analysis or an un-weighted analysis (FALSE).

For setK<-, the updated object.

## Author(s)

Gavin L. Simpson

## See Also

mat

## Examples

```
## Imbrie and Kipp Sea Surface Temperature
data(ImbrieKipp)
data(SumSST)
data(V12.122)
## merge training set and core samples
dat <- join(ImbrieKipp, V12.122, verbose = TRUE)
## extract the merged data sets and convert to proportions
ImbrieKipp <- dat[[1]] / 100
ImbrieKippCore <- dat[[2]] / 100
## fit a MAT model
ik.mat <- mat(ImbrieKipp, SumSST, method = "chord")
## How many analogues gives lowest RMSE?
getK(ik.mat)
## note that this value was chosen automatically
## Now set k to be 10
setK(ik.mat) <- 10
## check
```

```
getK(ik.mat)
```

```
gradientDist Positions of samples along a unit-length ordination gradient.
```


## Description

Extracts information as to the locations of samples along an ordination gradient. gradientDist() standardises the entire gradient to the interval $0, \ldots, 1$, to allow comparison between methods or data sets.

## Usage

```
gradientDist(object, ...)
## Default S3 method:
gradientDist(object, na.rm = TRUE, ...)
## S3 method for class 'cca'
gradientDist(object, na.rm = TRUE, axis = 1L,
            scaling = 0, ...)
    ## S3 method for class 'prcurve'
    gradientDist(object, na.rm = TRUE, ...)
```


## Arguments

object an R object of an appropriate type. For the default method, any R object that can be coerced to a vector.
na.rm logical; should missing values be removed?
axis numeric, length 1 ; the ordination axis to take as the gradient.
scaling Scaling to apply to the site scores. Default is to do no scaling. See scores.cca for details.
... additional arguments passed to other methods. In the "cca" method, these are also passed to scores.cca.

## Value

A numeric vector of positions along the gradient, scaled to the range $0, \ldots, 1$.

## Author(s)

Gavin L. Simpson

## See Also

See cca and prcurve for functions that produce objects that gradientDist() can work with.

## Examples

```
data(abernethy)
## Remove the Depth and Age variables
abernethy2 <- abernethy[, -(37:38)]
## Fit PCA
aber.pca <- rda(abernethy2)
## Distance along the first PCA axis
gradientDist(aber.pca)
```

```
hist.residLen Histogram plot for residual lengths
```


## Description

Base graphics histogram plot method for residLen objects.

## Usage

```
## S3 method for class 'residLen'
```

hist (x, breaks = "Sturges", freq = TRUE,
probs $=c(0.9,0.95,0.99)$, ncol $=1$, lcol = "red",
llty = "dashed", xlab = NULL, ylab = NULL,
main = "Residual distances", rug = TRUE, ...)

## Arguments

X
breaks How breakpoints for the histogram are determined. See hist for more details.
freq logical; if TRUE, the histogram graphic is a representation of frequencies, the counts component of the result; if FALSE, probability densities, component density, are plotted (so that the histogram has a total area of one). Defaults to TRUE
probs numeric; vector of probability quantiles to compute from the sets of residual distances.
ncol numeric; number of columns for the plot layout. Choices are 1 or 2. Determines whether the histograms are plotted above or beside each other.
lcol, llty colour and line-type for the quantiles.

| xlab, ylab | Axis labels. If not supplied, suitable defaults are generated, depending on whether <br> RDA or CCA was used as the underlying ordination model. |
| :--- | :--- |
| main | character; title for the plot. |
| rug | logical; should rug plots of the actual distances be drawn? |
| $\ldots$ | additional arguments passed to hist. |

## Value

A plot on the current device.
Returns a list with two components (train and passive), each of which is an object returned by hist.

## Author(s)

Gavin L. Simpson

## See Also

```
residLen, plot.residLen, histogram.residLen, densityplot.residLen.
```


## Examples

```
## load the Imbrie and Kipp example data
data(ImbrieKipp, SumSST, V12.122)
## squared residual lengths for Core V12.122
rlens <- residLen(ImbrieKipp, SumSST, V12.122)
rlens
## plot a histogram of the residual distances
hist(rlens)
```

histogram.residLen Lattice histogram plot for residual lengths

## Description

Lattice histogram method for residLen objects.

## Usage

\#\# S3 method for class 'residLen'
histogram(x, ..., xlab = NULL, ylab = NULL,
type = c("percent", "count", "density"))

## Arguments

| x | Object of class "residLen", the result of a call to residLen. |
| :--- | :--- |
| $\mathrm{xlab}, \mathrm{ylab}$ | Axis labels. If not supplied, suitable defaults are generated, depending on whether |
| RDA or CCA was used as the underlying ordination model. |  |

## Value

Returns an object of class "trellis". See histogram for details.

## Author(s)

Gavin L. Simpson

## See Also

residLen, plot.residLen, hist.residLen, densityplot.residLen.

## Examples

```
## load the Imbrie and Kipp example data
data(ImbrieKipp, SumSST, V12.122)
## squared residual lengths for Core V12.122
rlens <- residLen(ImbrieKipp, SumSST, V12.122)
rlens
## plot a histogram of the residual distances
histogram(rlens)
```

ImbrieKipp
Imbrie and Kipp foraminifera training set

## Description

The classic Imbrie and Kipp (1971) training set of counts on 27 species of foraminifera from 61 ocean sediment core surface samples and associated measures of summer and winter sea-surface temperatures and salinity at each location.

110 sediment cores samples from core V12-122 are also supplied in V12. 122.

## Usage

data(SumSST)
data(WinSST)
data(Salinity)
data(V12.122)

## Format

ImbrieKipp is a data frame with 61 observations on the following 27 species:
0 .univ a numeric vector
G.cglob a numeric vector
G. ruber a numeric vector
G.tenel a numeric vector
G. saccu a numeric vector
G. rubes a numeric vector
G.pacL a numeric vector
G. pacR a numeric vector
G.bullo a numeric vector
G.falco a numeric vector
G.calid a numeric vector
G. aequi a numeric vector
G.gluti a numeric vector
G.duter a numeric vector
G.infla a numeric vector
G. trnL a numeric vector
G.trnR a numeric vector
G.crasf a numeric vector
G.scitu a numeric vector
G.mentu a numeric vector
P.obliq a numeric vector
C. nitid a numeric vector
S.dehis a numeric vector
G.digit a numeric vector

Other a numeric vector
G.quin a numeric vector
G.hirsu a numeric vector

Summer and Winter sea-surface temperatures, and salinity values for the 61 sites in the Imbrie and Kipp training set (ImbrieKipp):

SumSST a numeric vector of summer sea-surface water temperatures. Values are in degrees C.
WinSST a numeric vector of winter sea-surface water temperatures. Values are in degrees C.
Salinity a numeric vector of sea water salinity values.
V12. 122 is a data frame with 110 observations from core V12-122 on the following 28 species:
O. univ a numeric vector
G.cglob a numeric vector
G. ruber a numeric vector
G.tenel a numeric vector
G. saccu a numeric vector
G.rubes a numeric vector
G. pacL a numeric vector
G.pacR a numeric vector
G.bullo a numeric vector
G.falco a numeric vector
G.calid a numeric vector
G. aequi a numeric vector
G.gluti a numeric vector
G.duter a numeric vector
G.infla a numeric vector
G. trnL a numeric vector
G. trnR a numeric vector
G.crasf a numeric vector
G.scitu a numeric vector
G.mentu a numeric vector
P.obliq a numeric vector
C. nitid a numeric vector
S.dehis a numeric vector
G.digit a numeric vector
G.hexag a numeric vector
G.cglom a numeric vector
cfH .pel a numeric vector
Other a numeric vector

## Source

Imbrie and Kipp (1971) TODO: Get the full reference.
These data were provided in electronic format by Prof. H. John B. Birks.

## Examples

```
data(ImbrieKipp)
head(ImbrieKipp)
data(SumSST)
data(WinSST)
data(Salinity)
plot(cbind(SumSST, WinSST, Salinity))
data(V12.122)
head(V12.122)
```

join Merge species data sets on common columns (species)

## Description

Merges any number of species matrices on their common columns to create a new data set with number of columns equal to the number of unqiue columns across all data frames. Needed for analysis of fossil data sets with respect to training set samples.

## Usage

```
join(..., verbose = FALSE, na.replace = TRUE, split = TRUE, value = 0,
            type = c("outer", "left", "inner"))
## S3 method for class 'join'
head(x, ...)
## S3 method for class 'join'
tail(x, ...)
```


## Arguments

$$
\begin{array}{ll}
\ldots . & \begin{array}{l}
\text { for join, data frames containing the data sets to be merged. For the head and } \\
\text { tail methods, additional arguments to head and tail, in particular " } n \text { " to con- } \\
\text { trol the number of rows of each joined data set to display. }
\end{array} \\
\text { verbose } & \begin{array}{l}
\text { logical; if TRUE, the function prints out the dimensions of the data frames in } \\
\text { "\dots", as well as those of the returned, merged data frame. }
\end{array} \\
\text { na.replace } & \begin{array}{l}
\text { logical; samples where a column in one data frame that have no matching col- } \\
\text { umn in the other will contain missing values (NA). If na. replace is TRUE, these } \\
\text { missing values are replaced with zeros. This is standard practice in ecology and } \\
\text { palaeoecology. If you want to replace with another value, then set na. replace } \\
\text { to FALSE and do the replacement later. } \\
\text { logical; should the merged data sets samples be split back into individual data } \\
\text { frames, but now with common columns (i.e. species)? }
\end{array}
\end{array}
$$

| value | numeric; value to replace NA with if na. replace is TRUE. |
| :--- | :--- |
| type | logical; type of join to perform. "outer" returns the union of the variables in <br> data frames to be merged, such that the resulting objects have columns for all <br> variables found across all the data frames to be merged. "left" returns the |
| left outer (or the left) join, such that the merged data frames contain the set of |  |
| variables found in the first supplied data frame. "inner" returns the inner join, |  |
| such that the merged data frame contain the intersection of the variables in the |  |
| supplied data frames. See Details. |  |
| $x$ | an object of class "join", usually the result of a call to join. |

## Details

When merging multiple data frames the set of variables in the merged data can be determined via a number of routes. join provides for two (currently) join types; the outer join and the left outer (or simply the left) join. Which type of join is performed is determined by the argument type.
The outer join returns the union of the set of variables found in the data frames to be merged. This means that the resulting data frame(s) contain columns for all the variable observed across all the data frames supplied for merging.
With the left outer join the resulting data frame(s) contain only the set of variables found in the first data frame provided.
The inner join returns the intersection of the set of variables found in the supplied data frames. The resulting data frame(s) contains the variables common to all supplied data frames.

## Value

If split = TRUE, an object of class "join", a list of data frames, with as many components as the number of data frames originally merged.

Otherwise, an object of class c("join", "data.frame"), a data frame containing the merged data sets.
head. join and tail. join return a list, each component of which is the result of a call to head or tail on each data set compont of the joined object.

## Author(s)

Gavin L. Simpson

## See Also

merge

## Examples

```
## load the example data
data(swapdiat, swappH, rlgh)
## merge training and test set on columns
dat <- join(swapdiat, rlgh, verbose = TRUE)
```

```
    ## extract the merged data sets and convert to proportions
    swapdiat <- dat[[1]] / 100
    rlgh <- dat[[2]] / 100
    ## merge training and test set using left join
    head(join(swapdiat, rlgh, verbose = TRUE, type = "left"))
    ## load the example data
    data(ImbrieKipp, SumSST, V12.122)
    ## merge training and test set on columns
    dat <- join(ImbrieKipp, V12.122, verbose = TRUE)
    ## extract the merged data sets and convert to proportions
    ImbrieKipp <- dat[[1]] / 100
    V12.122 <- dat[[2]] / 100
    ## show just the first few lines of each data set
    head(dat, n = 4)
    ## show just the last few lines of each data set
    tail(dat, n = 4)
    ## merge training and test set using inner join
    head(join(ImbrieKipp, V12.122, verbose = TRUE, type = "inner"))
    ## merge training and test set using outer join and replace
    ## NA with -99.9
    head(join(ImbrieKipp, V12.122, verbose = TRUE, value = -99.9))
```

logitreg Logistic regression models for assessing analogues/non-analogues

## Description

Fits logistic regression models to each level of group to model the probability of two samples being analogues conditional upon the dissimilarity between the two samples.

## Usage

```
logitreg(object, groups, k = 1, ...)
## Default S3 method:
logitreg(object, groups, k = 1,
    biasReduced = FALSE, ...)
## S3 method for class 'analog'
logitreg(object, groups, k = 1, ...)
## S3 method for class 'logitreg'
summary(object, p = 0.9, ...)
```


## Arguments

$$
\begin{array}{ll}
\text { object } & \begin{array}{l}
\text { for logitreg; a full dissimilarity matrix. For summary.logitreg an object of } \\
\text { class "logitreg", the result of a call to logitreg. }
\end{array} \\
\text { groups } & \begin{array}{l}
\text { factor (or object that can be coerced to one) containing the group membership } \\
\text { for each sample in object. }
\end{array} \\
\mathrm{k} & \begin{array}{l}
\text { numeric; the k closest analogues to use in the model fitting. }
\end{array} \\
\text { biasReduced } & \begin{array}{l}
\text { logical; should Firth's method for bias reduced logistic regression be used to fit } \\
\text { the models? If TRUE, model fits are performed via brglm. The default, FALSE, } \\
\text { indicates that models will be fitted via the standard glm function. }
\end{array} \\
\text { p } & \begin{array}{l}
\text { probability at which to predict the dose needed. }
\end{array} \\
\ldots & \begin{array}{l}
\text { arguments passed to other methods. These arguments are passed on to glm or } \\
\text { brglm. See their respective helps pages for details. Note that logitreg sets } \\
\text { internally the formula, data, and family arguments and hence can not be spec- } \\
\text { ified by the user. }
\end{array}
\end{array}
$$

## Details

Fits logistic regression models to each level of group to model the probability of two samples being analogues (i.e. in the same group) conditional upon the dissimilarity between the two samples.

This function can be seen as a way of directly modelling the probability that two sites are analogues, conditional upon dissimilarity, that can also be done less directly using roc and bayesF.

Often, the number of true analogues in the training set is small, both in absolute terms and as a proportion of comparisons. Logistic regression is known to suffer from a small-sample bias. Firth's method of bias reduction is a general solution to this problem and is implemented in logitreg through the brglm package of Ioannis Kosmidis.

## Value

logitreg returns an object of class "logitreg"; a list whose components are objects returned by glm. See glm for further details on the returned objects.

The components of this list take their names from group.
For summary.logitreg an object of class "summary.logitreg", a data frame with summary statistics of the model fits. The components of this data frame are:

In, Out The number of analogue and non-analogue dissimilarities analysed in each group, Est.(Dij), Std.Err

Coefficient and its standard error for dissimilarity from the logit model,
Z-value, p-value
Wald statistic and associated p-value for each logit model.
$\operatorname{Dij}(\mathrm{p}=$ ? ) , Std.Err(Dij)
The dissimilarity at which the posterior probability of two samples being analogues is equal to $p$, and its standard error. These are computed using dose.p.

## Note

The function may generate warnings from function glm.fit. These should be investigated and not simply ignored.
If the message is concerns fitted probabilities being numerically 0 or 1 , then check the fitted values of each of the models. These may well be numerically 0 or 1 . Heed the warning in glm and read the reference cited therein which may indicate problems with the fitted models, such as (quasi)complete separation.

## Author(s)

Gavin L. Simpson

## References

Firth, D. (1993). Bias reduction of maximum likelihood estimates. Biometrika 80, 27-38.

## See Also

roc, bayesF, glm, and brglm.

## Examples

```
## load the example data
data(swapdiat, swappH, rlgh)
## merge training and test set on columns
dat <- join(swapdiat, rlgh, verbose = TRUE)
## extract the merged data sets and convert to proportions
swapdiat <- dat[[1]] / 100
rlgh <- dat[[2]] / 100
## fit an analogue matching (AM) model using the squared chord distance
## measure - need to keep the training set dissimilarities
swap.ana <- analog(swapdiat, rlgh, method = "SQchord",
    keep.train = TRUE)
## fit the ROC curve to the SWAP diatom data using the AM results
## Generate a grouping for the SWAP lakes
METHOD <- if (getRversion() < "3.1.0") {"ward"} else {"ward.D"}
clust <- hclust(as.dist(swap.ana$train), method = METHOD)
grps <- cutree(clust, 6)
## fit the logit models to the analog object
swap.lrm <- logitreg(swap.ana, grps)
swap.lrm
## summary statistics
summary(swap.lrm)
## plot the fitted logit curves
```

```
plot(swap.lrm, conf.type = "polygon")
## extract fitted posterior probabilities for training samples
## for the individual groups
fit <- fitted(swap.lrm)
head(fit)
## compute posterior probabilities of analogue-ness for the rlgh
## samples. Here we take the dissimilarities between fossil and
## training samples from the `swap.ana` object rather than re-
## compute them
pred <- predict(swap.lrm, newdata = swap.ana$analogs)
head(pred)
## Bias reduction
## fit the logit models to the analog object
swap.brlrm <- logitreg(swap.ana, grps, biasReduced = TRUE)
summary(swap.brlrm)
```

mat

## Description

Modern Analogue Technique (MAT) transfer function models for palaeoecology. The fitted values are the, possibly weighted, averages of the environment for the $k$-closest modern analogues. MAT is a $k-\mathrm{NN}$ method.

## Usage

```
    mat (x, ...)
```

    \#\# Default S3 method:
    mat (x, y,
        method = c("euclidean", "SQeuclidean", "chord", "SQchord",
                            "bray", "chi.square", "SQchi.square",
                        "information", "chi.distance", "manhattan",
                        "kendall", "gower", "alt.gower", "mixed"),
        kmax, ...)
    \#\# S3 method for class 'formula'
    mat(formula, data, subset, na.action,
    method = c("euclidean", "SQeuclidean", "chord", "SQchord",
                        "bray", "chi.square", "SQchi.square",
            "information", "chi.distance", "manhattan",
            "kendall", "gower", "alt.gower", "mixed"),
        model = FALSE, ...)
    ```
## S3 method for class 'mat'
fitted(object, k, weighted = FALSE, ...)
## S3 method for class 'mat'
residuals(object, k, weighted = FALSE, ...)
```


## Arguments

\(\left.\begin{array}{ll}\mathrm{x} <br>
\mathrm{y} \& a data frame containing the training set data, usually species data. <br>
a vector containing the response variable, usually environmental data to be pre- <br>

dicted from x.\end{array}\right]\)| a symbolic description of the model to be fit. The details of model specification |
| :--- |
| are given below. |
| an optional data frame, list or environment (or object coercible by as. data.frame |
| to a data frame) containing the variables in the model. If not found in data, |
| the variables are taken from environment (formula), typically the environment |
| from which wa is called. |

## Details

The Modern Analogue Technique (MAT) is perhaps the simplest of the transfer function models used in palaeoecology. An estimate of the environment, $x$, for the response for a fossil sample, $y$, is the, possibly weighted, mean of that variable across the $k$-closest modern analogues selected from a modern training set of samples. If used, weights are the reciprocal of the dissimilarity between the fossil sample and each modern analogue.

A typical model has the form response ~ terms where response is the (numeric) response data frame and terms is a series of terms which specifies a linear predictor for response. A typical form for terms is ., which is shorthand for "all variables" in data. If . is used, data must also be provided. If specific species (variables) are required then terms should take the form spp1 + spp2 + spp3.
Pairwise sample dissimilarity is defined by dissimilarity or distance coefficients. A variety of coefficients are supported - see distance for details of the supported coefficients.
$k$ is chosen by the user. The simplest choice for $k$ is to evaluate the RMSE of the difference between the predicted and observed values of the environmental variable of interest for the training set samples for a sequence of models with increasing $k$. The number of analogues chosen is the value of $k$ that has lowest RMSE. However, it should be noted that this value is biased as the data used to build the model are also used to test the predictive power.
An alternative approach is to employ an optimisation data set on which to evaluate the size of $k$ that provides the lowest RMSEP. This may be impractical with smaller sample sizes.
A third option is to bootstrap re-sample the training set many times. At each bootstrap sample, predictions for samples in the bootstrap test set can be made for $k=1, \ldots, n$, where $n$ is the number of samples in the training set. $k$ can be chosen from the model with the lowest RMSEP. See function bootstrap. mat for further details on choosing $k$.

The output from summary.mat can be used to choose $k$ in the first case above. For predictions on an optimsation or test set see predict.mat. For bootstrap resampling of mat models, see bootstrap.mat.

The fitted values are for the training set and are taken as the, possibly weighted, mean of the environmental variable in question across the $k$-closest analogues. The fitted value for each sample does not include a contribution from itself - it is the closest analogue, having zero dissimilarity. This spurious distance is ignored and analogues are ordered in terms of the non-zero distances to other samples in the training set, with the $k$-closest contributing to the fitted value.
Typical usages for residuals.mat are:

```
resid(object, k, weighted = FALSE, \dots)
```


## Value

mat returns an object of class mat with the following components:
standard list; the model statistics based on simple averages of $k$-closest analogues. See below.
weighted list; the model statistics based on weighted of $k$-closest analogues. See below.
Dij matrix of pairwise sample dissimilarities for the training set $x$.
orig.x the original training set data.
orig.y the original environmental data or response, y .
call the matched function call.
method the dissimilarity coefficient used.

If model = TRUE then additional components "terms" and "model" are returned containing the terms object and model frame used.
fitted.mat returns a list with the following components:

- estimatednumeric; a vector of fitted values.
- knumeric; this is the $k$-closest analogue model with lowest apparent RMSE.
- weightedlogical; are the fitted values the weighted averages of the environment for the $k$ closest analogues. If FALSE, the fitted values are the average of the environment for the $k$ closest analogues.


## Note

The object returned by mat contains lists "standard" and "weighted" both with the following elements:

- esta matrix of estimated values for the training set samples for models using $k$ analogues, where $k=1, \ldots, n . n$ is the number of smaples in the training set. Rows contain the values for each model of size $k$, with colums containing the estimates for each training set sample.
- residmatrix; as for "est", but containing the model residuals.
- rmsepvector; containing the leave-one-out root mean square error or prediction.
- avg. biasvector; contains the average bias (mean of residuals) for models using $k$ analogues, where $k=1, \ldots, n$. $n$ is the number of smaples in the training set.
- max.biasvector; as for "avg.bias", but containing the maximum bias statistics.
- r.squaredvector; as for "avg.bias", but containing the $R^{2}$ statistics.


## Author(s)

Gavin L. Simpson

## References

Gavin, D.G., Oswald, W.W., Wahl, E.R. and Williams, J.W. (2003) A statistical approach to evaluating distance metrics and analog assignments for pollen records. Quaternary Research 60, 356-367.
Overpeck, J.T., Webb III, T. and Prentice I.C. (1985) Quantitative interpretation of fossil pollen spectra: dissimilarity coefficients and the method of modern analogues. Quaternary Research 23, 87-108.

Prell, W.L. (1985) The stability of low-latitude sea-surface temperatures: an evaluation of the CLIMAP reconstruction with emphasis on the positive SST anomalies, Report TR 025. U.S. Department of Energy, Washington, D.C.

Sawada, M., Viau, A.E., Vettoretti, G., Peltier, W.R. and Gajewski, K. (2004) Comparison of NorthAmerican pollen-based temperature and global lake-status with CCCma AGCM2 output at 6 ka . Quaternary Science Reviews 23, 87-108.

## See Also

summary.mat, bootstrap.mat for boostrap resampling of MAT models, predict.mat for making predictions from MAT models, fitted.mat and resid.mat for extraction of fitted values and residuals from MAT models respectively. plot.mat provides a plot. lm-like plotting tool for MAT models.

## Examples

```
## Imbrie and Kipp Sea Surface Temperature
data(ImbrieKipp)
data(SumSST)
data(V12.122)
## merge training set and core samples
dat <- join(ImbrieKipp, V12.122, verbose = TRUE)
## extract the merged data sets and convert to proportions
ImbrieKipp <- dat[[1]] / 100
ImbrieKippCore <- dat[[2]] / 100
## fit the MAT model using the squared chord distance measure
ik.mat <- mat(ImbrieKipp, SumSST, method = "chord")
ik.mat
## model summary
summary(ik.mat)
## fitted values
fitted(ik.mat)
## model residuals
resid(ik.mat)
## draw summary plots of the model
par(mfrow = c(2,2))
plot(ik.mat)
par(mfrow = c(1,1))
## reconstruct for the V12.122 core data
coreV12.mat <- predict(ik.mat, V12.122, k = 3)
coreV12.mat
summary(coreV12.mat)
## draw the reconstruction
reconPlot(coreV12.mat, use.labels = TRUE, display.error = "bars",
    xlab = "Depth", ylab = "SumSST")
## fit the MAT model using the squared chord distance measure
## and restrict the number of analogues we fit models for to 1:20
ik.mat2 <- mat(ImbrieKipp, SumSST, method = "chord", kmax = 20)
ik.mat2
```


## Description

Permutations and Monte Carlo simulations to define critical values for dissimilarity coefficients for use in MAT reconstructions.

```
Usage
    mcarlo(object, ...)
    ## Default S3 method:
    mcarlo(object, nsamp = 10000,
        type = c("paired", "complete", "bootstrap", "permuted"),
        replace = FALSE,
        method = c("euclidean", "SQeuclidean", "chord", "SQchord",
            "bray", "chi.square", "SQchi.square",
            "information", "chi.distance", "manhattan",
                            "kendall", "gower", "alt.gower", "mixed"),
        is.dcmat = FALSE, diag = FALSE, ...)
    ## S3 method for class 'mat'
    mcarlo(object, nsamp = 10000,
        type = c("paired", "complete", "bootstrap", "permuted"),
        replace = FALSE, diag = FALSE, ...)
    ## S3 method for class 'analog'
    mcarlo(object, nsamp = 10000,
        type = c("paired", "complete", "bootstrap", "permuted"),
        replace = FALSE, diag = FALSE, ...)
```


## Arguments

object an R object. Currently only object's of class "mat", "analog" or matrix-like object of species data allowed.
nsamp numeric; number of permutations or simulations to draw.
type character; the type of permutation or simulation to perform. See Details, below.
replace logical; should sampling be done with replacement?
method character; for raw species matrices, the dissimilarity coefficient to use. This is predefined when fitting a MAT model with mat or analogue matching via analogue and is ignored in the "mcarlo" methods for classes "mat" and "analog".
is.dcmat logical; is "object" a dissimilarity matrix. Not meant for general use; used internally by "mat" and "analogue" methods to instruct the "default" method that "object" is already a dissimilarity matrix, so there is no need to recalculate.
diag logical; should the dissimilarities include the diagonal (zero) values of the dissimilarity matrix. See Details.
... arguments passed to or from other methods.

## Details

Only "type" "paired" and "bootstrap" are currently implemented.
distance produces square, symmetric dissimilarity matrices for training sets. The upper triangle of these matrices is a duplicate of the lower triangle, and as such is redundant. mcarlo works on the lower triangle of these dissimilarity matrices, representing all pairwise dissimilarity values for training set samples. The default is not to include the diagonal (zero) values of the dissimilarity matrix. If you feel that these diagonal (zero) values are part of the population of dissimilarities then use "diag = TRUE" to include them in the permutations.

## Value

A vector of simulated dissimilarities of length "nsamp". The "method" used is stored in attribute "method".

## Note

The performance of these permutation and simulation techniques still needs to be studied. This function is provided for pedagogic reasons. Although recommended by Sawada et al (2004), sampling with replacement ("replace = TRUE") and including diagonal (zero) values ("diag = TRUE") simulates too many zero distances. This is because the same training set sample can, on occasion be drawn twice leading to a zero distance. It is impossible to find in nature two samples that will be perfectly similar, and as such sampling with replacement and "diag = TRUE" seems undesirable at best.

## Author(s)

Gavin L. Simpson

## References

Sawada, M., Viau, A.E., Vettoretti, G., Peltier, W.R. and Gajewski, K. (2004) Comparison of NorthAmerican pollen-based temperature and global lake-status with CCCma AGCM2 output at 6 ka . Quaternary Science Reviews 23, 87-108.

## See Also

mat for fitting MAT models and analog for analogue matching. roc as an alternative method for determining critical values for dissimilarity measures when one has grouped data.
plot.mcarlo provides a plotting method to visualise the distribution of simulated dissimilarities.

## Examples

```
## Imbrie and Kipp example
## load the example data
data(ImbrieKipp)
data(SumSST)
data(V12.122)
## merge training and test set on columns
dat <- join(ImbrieKipp, V12.122, verbose = TRUE)
## extract the merged data sets and convert to proportions
ImbrieKipp <- dat[[1]] / 100
V12.122 <- dat[[2]] / 100
## perform the modified method of Sawada (2004) - paired sampling,
## with replacement
ik.mcarlo <- mcarlo(ImbrieKipp, method = "chord", nsamp = 1000,
type = "paired", replace = FALSE)
ik.mcarlo
## plot the simulated distribution
layout(matrix(1:2, ncol = 1))
plot(ik.mcarlo)
layout(1)
```

minDC Extract minimum dissimilarities

## Description

Minimum dissimilarity is a useful indicator of reliability of reconstructions performed via MAT and other methods, and for analogue matching. Minimum dissimilarity for a sample is the smallest dissimilarity between it and the training set samples.

## Usage

```
minDC(x, ...)
## Default S3 method:
minDC(x, ...)
## S3 method for class 'predict.mat'
minDC(x, ...)
## S3 method for class 'analog'
minDC(x, probs = c(0.01, 0.02, 0.05, 0.1), ...)
## S3 method for class 'wa'
```

```
minDC(x, y,
    method = c("euclidean", "SQeuclidean", "chord", "SQchord",
                "bray", "chi.square", "SQchi.square", "information",
                "chi.distance", "manhattan", "kendall", "gower",
                "alt.gower", "mixed"),
    percent = FALSE, probs = c(0.01, 0.025, 0.05, 0.1), ...)
```


## Arguments

x
probs numeric; vector of probabilities with values in $[0,1]$.
y an optional matrix-like object containing fossil samples for which the minimum dissimilarities to training samples are to be calculated.
method character; which choice of dissimilarity coefficient to use. One of the listed options. See distance.
percent logical; Are the data percentages? If TRUE, the data ( $x$ and $y$ ) will be divided by 100 to convert them to the proportions expected by distance.
. . other arguments to be passed to other methods. Currently ignored.

## Value

minDC returns an object of class "minDC".
An object of class minDC is a list with some or all of the following components:

| minDC | numeric; vector of minimum dissimilarities. |
| :--- | :--- |
| method | character; the dissimilarity coefficient used. |
| quantiles | numeric; named vector of probability quantiles for distribution of dissimilarities <br> of modern training set. |

## Note

The "default" method of minDC will attempt to extract the relevant component of the object in question. This may be useful until a specific minDC method is written for a given class.

## Author(s)

Gavin L. Simpson

## See Also

predict.mat, and plot.minDC for a plotting method.

## Examples

```
## Imbrie and Kipp example
## load the example data
data(ImbrieKipp)
data(SumSST)
data(V12.122)
## merge training and test set on columns
dat <- join(ImbrieKipp, V12.122, verbose = TRUE)
## extract the merged data sets and convert to proportions
ImbrieKipp <- dat[[1]] / 100
V12.122 <- dat[[2]] / 100
## fit the MAT model using the squared chord distance measure
ik.mat <- mat(ImbrieKipp, SumSST, method = "SQchord")
ik.mat
## reconstruct for the V12-122 core data
v12.mat <- predict(ik.mat, V12.122)
## extract the minimum DC values
v12.mdc <- minDC(v12.mat)
v12.mdc
## draw a plot of minimum DC by time
plot(v12.mdc, use.labels = TRUE, xlab = "Depth (cm.)")
```

n2

Calculate Hill's N2 diversity measure

## Description

Hills N2 is a measure of species diversity, commonly referred to as "effective" diversity. If computed on the rows (samples) then the "effective" number of species in each row is returned, whereas, if computed on the columns (species) then the "effective" number of occurences of each species in the data set is returned.

## Usage

n2 (x, ...)
\#\# Default S3 method:
n2(x, which = c("species", "sites"), ...)

## Arguments

X
which
...
.

## Value

A numeric vector of N 2 values.

## Author(s)

Gavin L. Simpson

## Examples

```
data(swapdiat)
sppN2 <- n2(swapdiat, "species")
head(sppN2)
sampN2 <- n2(swapdiat, "sites")
head(sampN2)
```

optima Weighted averaging optima and tolerance ranges

## Description

Computes weighted average optima and tolerance ranges from species abundances and values of the environment.

## Usage

optima(x, ...)
\#\# Default S3 method:
optima(x, env, boot $=$ FALSE, nboot $=1000$, alpha $=0.05, \ldots$ )
\#\# Default S3 method:
tolerance(x, env, useN2 = TRUE, ...)

## Arguments

$x$
env Numeric; variable for which optima or tolerances are required.
boot, nboot logical (boot), numeric (nboot); should bootstrap resampling be employed to estimate the optima, and if so how many bootstrap samples to draw?
alpha numeric; 1-alpha gives the coverage for the percentile bootstrap confidence interval.
useN2 logical; should Hill's N2 values be used to produce un-biased tolerances?
Arguments passed to other methods.

## Value

Both functions return a named vector containing the WA optima or tolerances for the environmental gradient specified by env.

## Note

Objects of class "optima" or "tolerance" can be coerced to data frames using methods for as.data.frame.

## Author(s)

Gavin L. Simpson

## See Also

wa

## Examples

```
## Load the Imbrie & Kipp data and
## summer sea-surface temperatures
data(ImbrieKipp)
data(SumSST)
## WA optima
(opt <- optima(ImbrieKipp, SumSST))
## WA tolerances
(tol <- tolerance(ImbrieKipp, SumSST, useN2 = TRUE))
## caterpillar plot
caterpillarPlot(opt, tol)
## convert to data frame
as.data.frame(opt)
as.data.frame(tol)
## bootstrap WA optima - 100 resamples too low for SD & pCI
bopt <- optima(ImbrieKipp, SumSST, boot = TRUE, nboot = 100)
head(bopt)
```

panel.Loess Loess smooths to stratigraphic diagrams

## Description

A modified version of panel.loess, for drawing Loess smooths on stratigraphic diagrams.

## Usage

panel.Loess (x, y,
span $=1 / 3$, degree $=1$,
family = c("symmetric","gaussian"),
evaluation = 50,
lwd = plot.line\$lwd,
lty = plot.line\$lty,
col,
col.line = plot.line\$col,
type,
...)

## Arguments

$\mathrm{x}, \mathrm{y} \quad$ variables defining the contents of the panel.
span, degree, family, evaluation
arguments passed to loess. smooth, for which panel. Loess is a wrapper.
lwd, lty, col, col.line
graphical parameters.
type for compatibility with panel.loess, but is ignored within the function.
... graphical parameters can be supplied. Color can usually be specified by col, col. line, the latter overriding the first for lines.

## Details

The standard panel function panel. loess treats the data as the x -axis acting as the time component. For stratigraphic plots where time flows along the y-axis, we want the smoother to be fitted with the x -axis data as the response and the time component ( y -axis) as the predictor.
This modified version of panel. loess flips the two axes to produce the desired effect. Note also that it does not have argument horizontal as this is not required or supported by Stratiplot. In other respects, panel. Loess is equivalent to the lattice panel function panel.loess.

User should note that warnings can be generated by the fitting function if span is set too small for species with few observations. In such cases, the user is directed to the help page for loess. smooth, but increasing span slightly can often stop the warnings.

## Author(s)

Gavin L. Simpson, slightly modified from the Lattice function panel.loess by Deepayan Sarkar.

## See Also

loess.smooth, panel.loess.

```
panel.Stratiplot
```


## Description

A Lattice panel function for drawing individuals panels on stratigraphic diagrams using a range of plot types commonly used within palaeoecology.

## Usage

```
panel.Stratiplot(x, y,
                type \(=" l "\),
                        col,
                        pch = plot.symbol\$pch,
                        cex = plot.symbol\$cex,
                        col.line = plot.line\$col,
                        col.symbol = plot.symbol\$col,
                        col.refline = ref.line\$col,
                        col.smooth = "red",
                        col.poly = plot.line\$col,
                        lty = plot.line\$lty,
                        lwd = plot.line\$lwd,
                        lty.smooth = plot.line\$lty,
                        lwd.smooth \(=2\),
                        lwd.h = 3,
                        fill = plot.symbol\$fill,
                        zones = NULL,
                        col.zones = plot.line\$col,
                        lty.zones = plot.line\$lty,
                        lwd.zones = plot.line\$lwd,
                        gridh \(=-1\), gridv \(=-1\),
                        ...)
```


## Arguments

$\mathrm{x}, \mathrm{y} \quad$ variables defining the contents of the panel.
type character vector consisting of one or more of the following: "l", "p", "o", "b", "h", "g", "smooth", and "poly". It type has more than one element, the effects of each component are combined, though note that some elements will over-plot, and hence obscure, earlier elements.
For types "l", "p", "o", "b" and "g" the standard Lattice interpretation is observed. See panel. xyplot for further details. Note that type "b" is the same as type "o".
" $g$ " adds a reference grid using panel.grid in the background.
For " h ", histogram-like bars are plotted from the $\mathbf{y}$-axis, not from the x -axis with plot and panel.loess.

For "smooth" a loess fit is added to each panel using panel. Loess.
For "poly", a shaded polygon, or silhouette, is drawn for each panel.
col, col.line, col.symbol, col.poly, col.refline, col.smooth, col.zones
colour parameters. For all but col.smooth, default colours are obtained from plot. symbol and plot.line using trellis.par.get. col.refline controls the colour of the reference line drawn at value 0 on the x -axis, as well as the colour of the grid lines if drawn.
pch, cex, lty, lwd, fill
other graphical parameters, defaults for which are obtained from plot. symbol and plot.line using trellis.par.get.
lty.smooth line type for the loess smoother. The default is obtained from plot.line using trellis.par.get.
lwd.smooth, lwd.h
The line width for the loess smoother and histogram-like bars respectively.
zones numeric; vector of zone boundary positions on scale of the depth/time (y-)axis.
lty.zones, lwd.zones
line type and width for the zone markers. The defaults are obtained from plot. line.
gridh, gridv numeric arguments corresponding to $h$ and $v$ of panel.grid, which control the number of grid lines drawn.
... extra arguments passed on to the underlying panel functions; panel.points, panel.lines, panel.segments, panel. polygon, panel.Loess and panel.refline.

## Details

Creates stratigraphic scatter plots of $x$ and $y$, with various modifications possible via the type argument.
Zones can be drawn on the panels by supplying the numeric vector of zone boundaries as argument zones. The panel function will then draw horizontal lines across the panels at the desired $y$-axis locations. Note that the panel function does not attempt to identify the zone boundaries automatically; these must be determined via a chronological (constrained) cluster analysis function or similar.
Note that all the arguments controlling the display can be supplied directly to a high-level call of the function Stratiplot.

## Note

Histogram-like bars (type = "h") are drawn with lineend = "butt" to improve their appearance. This can not be changed by the user and you can't include that grid parameter in any call to panel. Stratiplot that uses type $=" h "$.

## Author(s)

Gavin L. Simpson

## See Also

Stratiplot, panel.Loess, panel.xyplot.

```
pcr
Prinicpal component regression transfer function models
```


## Description

Fits a palaeoecological transfer function model using principal component regression, using an optional transformation of the matrix of predictor variables when these are species abundance data.

## Usage

```
## Default S3 method:
pcr(x, y, ncomp, tranFun, ...)
## S3 method for class 'formula'
pcr(formula, data, subset, na.action, ..., model = FALSE)
Hellinger(x, ...)
ChiSquare(x, apply = FALSE, parms)
## S3 method for class 'pcr'
performance(object, ...)
## S3 method for class 'pcr'
residuals(object, comps = NULL, ...)
## S3 method for class 'pcr'
fitted(object, comps = NULL, ...)
## S3 method for class 'pcr'
coef(object, comps = NULL, ...)
## S3 method for class 'pcr'
screeplot(x, restrict = NULL,
    display = c("RMSE","avgBias","maxBias", "R2"),
    xlab = NULL, ylab = NULL, main = NULL, sub = NULL, ...)
    ## S3 method for class 'pcr'
    eigenvals(x, ...)
```


## Arguments

X
Matrix or data frame of predictor variables. Usually species composition or abundance data for transfer function models. For screeplot and eigenvals, an object of class "pcr".

Numeric vector; the response variable to be modelled.
\(\left.$$
\begin{array}{ll}\text { ncomp } & \begin{array}{l}\text { numeric; number of principal components to build models for. If not supplied } \\
\text { the largest possible number of components is determined. } \\
\text { function; a function or name of a function that performs a transformation of the } \\
\text { predictor variables x. The function must be self-contained as no arguments are } \\
\text { passed to the function when it is applied. See Details for more information. } \\
\text { a model formula. }\end{array}
$$ <br>
formula <br>
an optional data frame, list or environment (or object coercible by as. data.frame <br>
to a data frame) containing the variables specified on the RHS of the model for- <br>
mula. If not found in data, the variables are taken from environment (formula), <br>

typically the environment from which pcr is called.\end{array}\right]\)| an optional vector specifying a subset of observations to be used in the fitting |
| :--- |
| process. |

## Details

When applying cross-validation (CV) to transfer function models, any transformation of the predictors must be applied separately during each iteration of the CV procedure to the part of the data used in fitting the model. In the same way, any samples to be predicted from the model must use any meta-parameters derived from the training data only. For examle, centring is appled to the training data only and the variables means used to centre the training data are used to centre the test samples. The variable means should not be computed on a combination of the training and test samples.
When using PCR, we might wish to apply a transformation to the species data predictor variables such that the PCA of those data preserves a dissimilarity coefficient other than the Euclidean distance. This transformation is applied to allow PCA to better describe patterns in the species data (Legendre \& Gallagher 2001).
How this is handled in per is to take a user-supplied function that takes a single argument, the matrix of predictor variables. The function should return a matrix of the same dimension as the input. If
any meta-parameters are required for subsequent use in prediction, these should be returned as attribute "parms", attached to the matrix.
Two example transformation functions are provided implementing the Hellinger and Chi Square transformations of Legendre \& Gallagher (2001). Users can base their transformation functions on these. ChiSquare () illustrates how meta-parameters should be returned as the attribute "parms".

## Value

Returns an object of class "pcr", a list with the following components:
fitted.values matrix; the PCR estimates of the response. The columns contain fitted values using C components, where C is the Cth column of the matrix.
coefficients matrix; regression coefficients for the PCR. Columns as per fitted above.
residuals matrix; residuals, where the Cth column represents a PCR model using C components.
scores
loadings
Yloadings
$x$ Means numeric; means of the predictor variables in the training data.
yMean numeric; mean of the response variable in the training data.
varExpl numeric; variance explained by the PCR model. These are the squares of the singular values.
totvar numeric; total variance in the training data
call the matched call.
tranFun transformation function used. NA if none supplied/used.
tranParms list; meta parameters used to computed the transformed training data.
performance data frame; cross-validation performance statistics for the model.
ncomp numeric; number of principal components computed

## Author(s)

## Gavin L. Simpson

## See Also

wa

## Examples

```
## Load the Imbrie & Kipp data and
## summer sea-surface temperatures
data(ImbrieKipp)
data(SumSST)
## normal interface and apply Hellinger transformation
```

```
mod <- pcr(ImbrieKipp, SumSST, tranFun = Hellinger)
mod
## formula interface, but as above
mod2 <- pcr(SumSST ~ ., data = ImbrieKipp, tranFun = Hellinger)
mod2
## Several standard methods are available
fitted(mod, comps = 1:4)
resid(mod, comps = 1:4)
coef(mod, comps = 1:4)
## Eigenvalues can be extracted
eigenvals(mod)
## screeplot method
screeplot(mod)
```

    performance Transfer function model performance statistics
    
## Description

A simple extractor function to access the model performance statistics of transfer function models.

## Usage

performance(object, ...)

## Arguments

object A transfer function object.
. . Arguments passed to other methods. Currently ignored.

## Details

performance is a generic function for use with a number of fitted models objects in analogue. The available methods are:

- wa Weighted Averaging Models.
- predict. wa Predictions from a Weighted Average Model.
- pcr Principal Component Regression models.
- bootstrap.wa Bootstrapped Weighted Averaging Models.
- crossval Cross-validated models fitted via crossval.


## Value

A named vector containing the extracted model performance statistics.

## Author(s)

Gavin L. Simpson

## See Also

wa, predict.wa, bootstrap.wa.

## Examples

```
data(ImbrieKipp)
data(SumSST)
## fit the WA model
mod <- wa(SumSST ~., data = ImbrieKipp)
mod
## the model performance statistics
performance(mod)
```

plot.dissimilarities Plots the distribution of extracted dissimilarities

## Description

Produces a plot of the distribution of the extracted dissimilarities and a reference normal distribution with comparable mean and sd.

## Usage

```
## S3 method for class 'dissimilarities'
plot(x, prob = 0.05,
    legend = TRUE, n.rnorm = 1e+05, col = "black",
    col.ref = "red", lty = "solid", lty.quant = "dotted",
    xlab = NULL, ylab = NULL, main = NULL, sub = NULL, ...)
```


## Arguments

x
prob numeric; density probability defining the threshold for close modern analogues.
legend logical; draw a legend on the plotted figure?
n.rnorm numeric; number of random normal deviates for reference line.
col, col.ref colours for the dissimilarity and reference density functions drawn.
lty, lty.quant line types for the dissimilarity and reference density functions drawn.
$x l a b, y l a b \quad c h a r a c t e r ; x-a n d y$-axis labels.
main, sub character; main and subtitle for the plot.
... graphical arguments passed to other graphics functions.

## Value

A plot on the currently active device.

## Author(s)

Gavin L. Simpson

## See Also

dissimilarities

## Examples

```
## Imbrie and Kipp example
## load the example data
data(ImbrieKipp)
data(SumSST)
data(V12.122)
## merge training and test set on columns
dat <- join(ImbrieKipp, V12.122, verbose = TRUE)
## extract the merged data sets and convert to proportions
ImbrieKipp <- dat[[1]] / 100
V12.122 <- dat[[2]] / 100
## analog matching between SWAPImbrie & Kipp and V12.122 core
ik.analog <- analog(ImbrieKipp, V12.122, method = "chord")
ik.analog
summary(ik.analog)
## compare training set dissimilarities with normals
## and derive cut-offs
ik.dij <- dissim(ik.analog)
plot(ik.dij)
```

plot.evenSample Plot distribution of samples along gradient

## Description

plot method for objects produced by evenSample. Draws a histogram of the number of samples per gradient segment.

## Usage

\#\# S3 method for class 'evenSample'
plot(x, add = FALSE, xlim = NULL, ylim = NULL, col = "grey",
border = "grey", lty = NULL, ylab, xlab, main = NULL, sub = NULL, ann = TRUE, axes = TRUE, ...)

## Arguments

X
add
xlim, ylim
col, border
lty
ylab, xlab, main, sub character strings used to label the plot
ann logical; should the default annotations be added to the plot. This relates to the plot main and sub titles and the x and y axis labels.
axes logical; should plot axes be drawn?
.. additional arguments passed to/from other methods.

## Value

A plot is draw is drawn on the currently active device.

## Author(s)

Gavin L. Simpson

## See Also

evenSample for the function used to create objects used by this plot method.

## Description

Draws the fitted logistic regression function describing the posterior probability that two sites are analogues conditional upon the dissimilarity between the two samples. Confidence intervals are also computed and displayed if requested.

## Usage

```
## S3 method for class 'logitreg'
plot(x, group = "all", npred = 100,
    conf.int = 0.9, conf.type = c("none", "polygon", "lines"),
        xlab = expression(D[ij]), ylab = "Pr (A+ | d)",
        rug = TRUE, ticksize = 0.02,
        col = "red", ref.col = "lightgrey",
        lwd = 2, conf.lwd = 1, conf.lty = "dashed",
        shade = "lightgrey", ...)
```


## Arguments

x
group The group to plot the logit model for. Can be one of the group labels or "Combined" to draw the individual logit models. Alternatively, and the default, is to use "all", which divides the plotting region into the required number of plotting regions and draws all the fitted curves.
npred number of points at which the fitted curves are evaluated for plotting purposes.
conf.int numeric; the confidence interval required.
conf.type character; how should the confidence interval be drawn. Default is not to draw the confidence interval.
$x l a b, y l a b \quad$ character; the x and y axis labels.
rug logical; should rug plots be drawn?
ticksize The size of the tick marks used in rug plots.
col The colour in which to draw the line representing the fitted values.
ref.col The colour of the reference lines drawn at 0 and 1 .
lwd The line width in which to draw the line representing the fitted values.
conf.lwd, conf.lty
Line width and line type for the confidence interval. Only used if conf. type = "lines" is specified.
shade The colour for the fill and border of the confidence interval if conf.type $=$ "polygon" is specified.
.. arguments passed on to plot.

## Value

A plot on the current device.

## Author(s)

Gavin L. Simpson

## See Also

logitreg for an example, roc

## Description

Five plots (selectable by which) are currently available: a plot of estimated against observed values, a plot of residuals against estimated values, and screeplots of the apparent RMSE, average bias and maximum bias for MAT models of size $k$, where $k=1, \ldots, n$. By default, the first three and ' 5 ' are provided.

## Usage

```
## S3 method for class 'mat'
plot(x,
    which = c(1:3, 5),
    weighted = FALSE,
    k,
    caption = c("Inferred vs Observed", "Residuals vs Fitted",
                            "Leave-one-out errors", "Average bias",
                            "Maximum bias"),
    max.bias = TRUE,
    n.bias = 10,
    restrict = 20,
    sub.caption = NULL,
    main = "",
    ask = prod(par("mfcol")) < length(which) &&
        dev.interactive(),
        panel = if (add.smooth) panel.smooth else points,
        add.smooth = getOption("add.smooth"))
```


## Arguments

x
which
weighted logical; should the analysis use weighted mean of env data of analogues as fitted/estimated values?
$k \quad$ numeric; the number of analogues to use. If missing $k$ is chosen automatically as the k that achieves lowest RMSE.
caption captions to appear above the plots.
max.bias logical, should max bias lines be added to residuals.
n.bias numeric, number of sections to calculate maximum bias for.
restrict logical; restrict comparison of k-closest model to $k \leq$ restrict.

| sub. caption | common title-above figures if there are multiple; used as 'sub' (s. 'title') other- <br> wise. If NULL, as by default, a possibly shortened version of deparse ( $x \$ c a l l$ ) <br> is used. |
| :--- | :--- |
| main | title to each plot-in addition to the above caption. |
| ask | logical; if TRUE, the user is asked before each plot, see par (ask=.). <br> $\ldots$ <br> panel |
| graphical arguments passed to other graphics functions. <br> panel function. The useful alternative to points, panel. smooth, can be chosen <br> by add. smooth = TRUE. |  |
| add. smooth | logical indicating if a smoother should be added to fitted $\backslash \&$ residuals plots; see <br> also panel above. |

## Details

This plotting function is modelled closely on plot. Im and many of the conventions and defaults for that function are replicated here.
sub. caption - by default the function call - is shown as a subtitle (under the x-axis title) on each plot when plots are on separate pages, or as a subtitle in the outer margin (if any) when there are multiple plots per page.

## Value

One or more plots, drawn on the current device.

## Author(s)

Gavin L. Simpson. Code borrows heavily from plot. lm.

## See Also

mat

## Examples

```
## Imbrie and Kipp example
## load the example data
data(ImbrieKipp)
data(SumSST)
data(V12.122)
## merge training and test set on columns
dat <- join(ImbrieKipp, V12.122, verbose = TRUE)
## extract the merged data sets and convert to proportions
ImbrieKipp <- dat[[1]] / 100
V12.122 <- dat[[2]] / 100
## MAT
ik.mat <- mat(ImbrieKipp, SumSST, method = "chord")
```

\#\# summary plot of MAT model
layout(matrix(1:4, ncol = 2, byrow = TRUE))
plot(ik.mat)
layout(1)
plot.mcarlo Plot Monte Carlo simulated dissimilarity distributions

## Description

A plot.lm-like plotting function for objects of class "mcarlo" to visualise the simulated distribution of dissimilarities.

## Usage

```
## S3 method for class 'mcarlo'
plot(x,
    which = c(1:2),
    alpha = 0.05,
    caption = c("Distribution of dissimilarities",
        expression(paste("Simulated probability Pr (Dissim < ",
                alpha, ")"))),
    col.poly = "lightgrey",
    border.poly = "lightgrey",
    ask = prod(par("mfcol")) < length(which) &&
                            dev.interactive(),
    ...)
```


## Arguments

x
which numeric; which of the plots should be produced?
alpha numeric; the Monte Carlo significance level to be marked on the cumulative frequency plot.
caption character, length 2; captions to appear above the plots.
col.poly, border.poly
character; the colour to draw the region and border of the polygon enclosing the Monte Carlo significance on the cummulative frequency plot.
ask logical; should the function wait for user confirmation to draw each plot? If missing, the function makes a reasonable attempt to guess the current situation and act accordingly.
... additional graphical parameters to be passed to the plotting functions. Currently ignored.

## Details

The "Distribution of dissimilarities" plot produces a histogram and kernel density estimate of the distribution of simulated dissimilarity values.
The "Simulated probability" plot shows a cumulative probability function of the simulated dissimlarity values, and highlights the proportion of the curve that is less than alpha.

## Value

One or more plots on the current device.

## Author(s)

Gavin L. Simpson

## References

Sawada, M., Viau, A.E., Vettoretti, G., Peltier, W.R. and Gajewski, K. (2004) Comparison of NorthAmerican pollen-based temperature and global lake-status with CCCma AGCM2 output at 6 ka . Quaternary Science Reviews 23, 87-108.

## See Also

```
mcarlo
```


## Examples

```
## Imbrie and Kipp example
## load the example data
data(ImbrieKipp)
data(SumSST)
data(V12.122)
## merge training and test set on columns
dat <- join(ImbrieKipp, V12.122, verbose = TRUE)
## extract the merged data sets and convert to proportions
ImbrieKipp <- dat[[1]] / 100
V12.122 <- dat[[2]] / 100
## perform the modified method of Sawada (2004) - paired sampling,
## with replacement
ik.mcarlo <- mcarlo(ImbrieKipp, method = "chord", nsamp = 1000,
    type = "paired", replace = FALSE)
ik.mcarlo
## plot the simulated distribution
layout(matrix(1:2, ncol = 1))
plot(ik.mcarlo)
layout(1)
```

```
plot.minDC Plot of minimum dissimilarity per sample
```


## Description

Minimum dissimilarity is a useful indicator of reliability of reconstructions performed via MAT and other methods, and for analogue matching. Minimum dissimilarity for a sample is the smallest dissimilarity between it and the training set samples.

```
Usage
    ## S3 method for class 'minDC'
    plot(x, depths, use.labels = FALSE,
        quantiles = TRUE, rev.x = TRUE, type = "l",
        xlim, ylim, xlab = "", ylab = "Dissimilarity",
        main = "", sub = NULL,
        col.quantile = "red", lty.quantile = "dotted",
        ...)
```

Arguments
$x \quad$ an object of class "minDC".
depths numeric; a vector of depths for which predicted values exist or will be generated.
Can be missing, in which case, if use. labels = TRUE, the function will attempt
to derive suitable values for you. See Details below.
use.labels logical; should reconPlot attempt to derive values for argument depths from
the names of the predicted values? Only use if depths is missing. See Details
below.
quantiles logical; should the probability quantiles be drawn on the plot?
rev.x logical; should the depth/age axis be reversed (drawn from high to low)?
type type of line drawn. See par and argument "type".
$x l a b, y l a b \quad$ character; the $x$ - and $y$-axis labels respectively.
main, sub character; main title and subtitle for the plot.
xlim, ylim numeric, length 2; the $x$ - and y-limits for the plotted axes. If not provided, the
function will calculate appropriate values to cover the range of plotted values
and any quantile lines (if requested via "quantiles".
col.quantile colour in which to draw the quantile lines.
lty.quantile line type in which to draw the quantile lines.
...
arguments to be passed to methods, such as graphical parameters (see par).
Currently ignored.

## Details

Conventionally, these plots are drawn on a depth or an age scale. Argument depths is used to provide the depth or age axis, against which the predicted values are plotted.

If depths is not provided, then the function will try to derive the appropriate values from the labels of the predictions if use.labels = TRUE. You must provide depths or set use.labels = TRUE otherwise an error will result. The derived labels will be coerced to numerics. If your labels are coercible, then you'll either get nonsense on the plot or an error from R. If so, provide suitable values for depths.

## Value

A plot on the currently active device.

## Author(s)

Gavin L. Simpson

## See Also

minDC

## Examples

```
## Imbrie and Kipp example
## load the example data
data(ImbrieKipp)
data(SumSST)
data(V12.122)
## merge training and test set on columns
dat <- join(ImbrieKipp, V12.122, verbose = TRUE)
## extract the merged data sets and convert to proportions
ImbrieKipp <- dat[[1]] / 100
V12.122 <- dat[[2]] / 100
## fit the MAT model using the chord distance measure
(ik.mat <- mat(ImbrieKipp, SumSST, method = "chord"))
## reconstruct for the RLGH core data
v12.mat <- predict(ik.mat, V12.122)
## extract the minimum DC values
v12.mdc <- minDC(v12.mat)
v12.mdc
## draw a plot of minimum DC by time
plot(v12.mdc, use.labels = TRUE, xlab = "Depth (cm.)")
```

```
plot.prcurve
```


## Description

Projects the principal curve into PCA space and draws it and the underlying data in a biplot.

## Usage

```
## S3 method for class 'prcurve'
plot(x, axes = 1:2, scaling = 0, segments = TRUE,
        col = "red", col.seg = "forestgreen", lwd = 2,
        lwd.seg = 1, ...)
    ## S3 method for class 'prcurve'
    lines(x, axes = 1:2, scaling = 0, segments = TRUE,
        col = "red", col.seg = "forestgreen", lwd = 2,
        lwd.seg = 1, ...)
```


## Arguments

x
axes numeric vector of length 2; this is passed to the choices argument of the scores function.
scaling numeric; the scaling to use. See scores.rda for the available options. The default is not to scale the scores, but scaling $=1$ might be a useful alternative.
segments logical; should segments be drawn between the observed points to the location on the principal curve on to which they project.
col The colour to draw the principal curve in.
col.seg The colour to draw the segments in.
lwd, lwd.seg The line thickness used to draw the principal curve and segments respectively.
additional arguments passed on to points when drawing the observations in PCA space.

## Value

A plot on the currently active device. The function does not return anything.

## Author(s)

Gavin L. Simpson

## See Also

prcurve; rda for the code used to perform the PCA.

## Examples

```
## Load the Abernethy Forest data
data(abernethy)
## Remove the Depth and Age variables
abernethy2 <- abernethy[, -(37:38)]
## Fit the principal curve using varying complexity of smoothers
## for each species
aber.pc <- prcurve(abernethy2, method = "ca", trace = TRUE,
    vary = TRUE, penalty = 1.4)
## Plot the curve
plot(aber.pc)
## The lines() method can be used to add the principal curve to an
## existing plot
ord <- rda(abernethy2)
plot(ord, scaling = 1)
lines(aber.pc, scaling = 1)
```

plot.residLen Plot method for residual lengths

## Description

Base graphics plot method for residLen objects.

## Usage

\#\# S3 method for class 'residLen'
plot $(x$, probs $=c(0.9,0.95,0.99)$, ncol $=1$,
lcol = "red", llty = "dashed", xlab = NULL, ylab = NULL,
main = "Residual distances", rug = TRUE, ...)

## Arguments

x
probs
ncol numeric; number of columns for the plot layout. Choices are 1 or 2. Determines whether the histograms are plotted above or beside each other.
lcol, llty
$x l a b, y l a b \quad$ Axis labels. If not supplied, suitable defaults are generated, depending on whether RDA or CCA was used as the underlying ordination model.
main character; title for the plot.
rug logical; should rug plots of the actual distances be drawn?
... additional arguments passed to plot.

## Value

A plot on the current device.
Returns, invisibly, a list with two components (train and passive), each and object of the type returned by density.

## Author(s)

Gavin L. Simpson

## See Also

residLen, plot.residLen, histogram.residLen, densityplot.residLen.

## Examples

```
## load the Imbrie and Kipp example data
data(ImbrieKipp, SumSST, V12.122)
## squared residual lengths for Core V12.122
rlens <- residLen(ImbrieKipp, SumSST, V12.122)
rlens
## plot a histogram of the residual distances
plot(rlens)
```

    plot.roc Plot ROC curves and associated diagnostics
    
## Description

Produces up to four plots (selectable by "which") from the results of a call to roc, including the ROC curve itself.

## Usage

```
## S3 method for class 'roc'
plot(x,
    which = c(1:3,5),
    group = "Combined",
    prior = NULL,
    show.stats = TRUE,
    abline.col = "grey",
    abline.lty = "dashed",
    inGroup.col = "red",
    outGroup.col = "blue",
    lty = "solid",
```

```
caption = c("ROC curve", "Dissimilarity profiles",
    "TPF - FPF vs Dissimilarity",
    "Likelihood ratios"),
legend = "topright",
ask = prod(par("mfcol")) < length(which) && dev.interactive(),
...)
```


## Arguments

| x | an object of class "roc". |
| :--- | :--- |
| which | numeric vector; which aspects of "roc" object to plot if a subset of the plots is <br> required, specify a subset of the numbers $1: 5$. |
| character vector of length 1 giving the name of the group to plot. |  |
| prior | numeric vector of length 2 (e.g. c $(0.5,0.5)$ ) specifiying the prior probabilities <br> of analogue and no-analogue. Used to generate posterior probability of analogue <br> using Bayes factors in plot 5 (which = 5). |
| show.stats | logical; should concise summary statistics of the ROC analysis be displayed on <br> the plot? |
| abline.col | character string or numeric value; the colour used to draw vertical lines at the <br> optimal dissimilarity on the plots. |
| abline.lty | Line type for indicator of optimal ROC dissimilarity threshold. See par for the <br> allowed line types. <br> character string or numeric value; the colour used to draw the density curve for |
| inGroup.col | the dissimilarities between sites in the same group. <br> character string or numeric value; the colour used to draw the density curve for <br> the dissimilarities between sites not in the same group. |
| outGroup.col |  |

## Details

This plotting function is modelled closely on plot. 1m and many of the conventions and defaults for that function are replicated here.
First, some definitions:
TPF True Positive Fraction, also known as sensitivity.
TNF True Negative Fraction, also known as specificity.

FPF False Positive Fraction; the complement of TNF, calculated as 1 - TNF. This is often referred to a 1 - specificity. A false positive is also known as a type I error.

FNF False Negative Fraction; the complement of TPF, calculated as 1 - TPF. A false negative is also known as a type II error.

AUC The Area Under the ROC Curve.

The "ROC curve" plot (which = 1, draws the ROC curve itself as a plot of the False Positive Fraction against the True Positive Fraction. A diagonal 1:1 line represents no ability for the dissimilarity coefficient to differentiate between groups. The AUC statistic may also be displayed (see argument "show.stats" above).
The "Dissimilarity profile" plot (which = 2), draws the density functions of the dissimilarity values (d) for the correctly assigned samples and the incorrectly assigned samples. A dissimilarity coefficient that is able to well distinguish the sample groupings will have density functions for the correctly and incorrectly assigned samples that have little overlap. Conversely, a poorly discriminating dissimilarity coefficient will have density profiles for the two assignments that overlap considerably. The point where the two curves cross is the optimal dissimilarity or critical value, $d^{\prime}$. This represents the point where the difference between TPF and FPF is maximal. The value of $d$ at the point where the difference between TPF and FPF is maximal will not neccesarily be the same as the value of $d^{\prime}$ where the density profiles cross. This is because the ROC curve has been estimated at discrete points $d$, which may not include excatly the optimal $d^{\prime}$, but which should be close to this value if the ROC curve is not sampled on too coarse an interval.
The "TPF - FPF vs Dissimilarity" plot, draws the difference between the ROC curve and the $1: 1$ line. The point where the ROC curve is farthest from the $1: 1$ line is the point at which the ROC curve has maximal slope. This is the optimal value for $d$, as discussed above.

The "Likelihood ratios" plot, draws two definitions of the slope of the ROC curve as the likelihood functions $\operatorname{LR}(+)$, and $\operatorname{LR}(-) . \operatorname{LR}(+)$, is the likelihood ratio of a positive test result, that the value of $d$ assigns the sample to the group it belongs to. LR(-) is the likelihood ratio of a negative test result, that the value of $d$ assigns the sample to the wrong group.
$\mathrm{LR}(+)$ is defined as $L R(+)=T P F / F P F$ (or sensitivity / (1-specificity)), and LR(-) is defined as $L R(-)=F P F / T N F$ (or (1-sensitivity) / specificity), in Henderson (1993).

The "probability of analogue" plot, draws the posterior probability of analogue given a dissimilarity. This is the $\mathrm{LR}(+)$ likelihood ratio values multiplied by the prior odds of analogue, for given values of the dissimilarity, and is then converted to a probability.

## Value

One or more plots, drawn on the current device.

## Author(s)

Gavin L. Simpson. Code borrows heavily from plot. lm.

## References

Brown, C.D., and Davis, H.T. (2006) Receiver operating characteristics curves and related decision measures: A tutorial. Chemometrics and Intelligent Laboratory Systems 80, 24-38.

Gavin, D.G., Oswald, W.W., Wahl, E.R. and Williams, J.W. (2003) A statistical approach to evaluating distance metrics and analog assignments for pollen records. Quaternary Research 60, 356-367.
Henderson, A.R. (1993) Assessing test accuracy and its clinical consequences: a primer for receiver operating characteristic curve analysis. Annals of Clinical Biochemistry 30, 834-846.

## See Also

roc for a complete example

```
plot.sppResponse Plot species responses along gradients or latent variables
```


## Description

Observed abundances and fitted response curves along

## Usage

```
## S3 method for class 'sppResponse'
plot(x,
    which = seq_along(x),
    display = c("observed", "fitted"),
    xlab = "Gradient", ylab = "Abundance",
    main = NULL,
    lcol = "red",
    lwd = 2,
    ...)
```


## Arguments

an object of class "sppResponse".
which numeric or logical vector; which species (components of $x$ ) to plot.
display character; plot the observed data or the fitted species response curve or both?
xlab, ylab, main
titles for the x and y axis label and the main title. If main is NULL, suitable main titles are taken from $x$.
lwd numeric; width of the line used to draw the fitted species response function if drawn.
lcol the colour to use to draw the fitted species response.
... graphical arguments passed plot and lines used internally.

## Details

Currently, no attempt is made to split the device into an appropriate number of panels. This is up to the user to decide. See the Examples section of sppResponse for one way to handle this.

## Value

One or more plots, drawn on the current device.

## Author(s)

Gavin L. Simpson

## See Also

sppResponse for a complete example using fitted responses along principal curves.

```
plot.wa
Plot diagnostics for a weighted averaging model
```


## Description

Two plots (selectable by which) are currently available: a plot of estimated against observed values, a plot of residuals against estimated values.

## Usage

```
    ## S3 method for class 'wa'
    plot(x,
        which = 1:2,
        caption = c("Inferred vs Observed", "Residuals vs Fitted"),
        max.bias = TRUE,
        n.bias = 10,
        sub.caption = NULL,
        main = "",
        ask = prod(par("mfcol")) < length(which) &&
        dev.interactive(),
        panel = if (add.smooth) panel.smooth else points,
        add.smooth = getOption("add.smooth"))
```


## Arguments

x
which which aspects of the "wa" object to plot if a subset of the plots is required, specify a subset of the numbers $1: 2$.
caption captions to appear above the plots.
max.bias logical, should max bias lines be added to residuals.
n.bias numeric, number of sections to calculate maximum bias for.
sub.caption common title-above figures if there are multiple; used as 'sub' (s.'title') otherwise. If NULL, as by default, a possibly shortened version of deparse ( $x \$ c a l l$ ) is used.

| main | title to each plot-in addition to the above caption. |
| :--- | :--- |
| ask | logical; if TRUE, the user is asked before each plot, see par (ask=.). |
| $\ldots$ | graphical arguments passed to other graphics functions. |
| panel | panel function. The useful alternative to points, panel. smooth, can be chosen <br> by add. smooth = TRUE. |
| add. smooth | logical indicating if a smoother should be added to fitted $\backslash \&$ residuals plots; see <br> also panel above. |

## Details

This plotting function is modelled closely on plot. 1m and many of the conventions and defaults for that function are replicated here.
sub. caption - by default the function call - is shown as a subtitle (under the x-axis title) on each plot when plots are on separate pages, or as a subtitle in the outer margin (if any) when there are multiple plots per page.

## Value

One or more plots, drawn on the current device.

## Author(s)

Gavin L. Simpson. Code borrows heavily from plot.lm.

## See Also

mat

## Examples

```
## see full example in ?wa
```

North American Modern Pollen Database

## Description

A database of modern pollen samples from a network of sites from North America and Greenland, compiled by Whitmore et al. (2005). Associated climatic and vegetation data are also record. The version of the NAMPD included here is latest version 1-7.3 (February 2013), as of January 2016.

## Usage

data(Pollen)
data(Climate)
data(Biome)
data(Location)

## Format

For Pollen, a data frame of 4833 samples and 135 columns (the unique identifier and 134 pollen taxa).
For Biome, a data frame of 4833 samples on, currently, a single vegetation variable (plus the unique identifier):

ID2 Unique, sequential number assigned by NAMPD.
Fedorova Factor; Vegetation type (Biome) See Whitmore et al. (2005), Figs 3 \& 4. Reclassified biomes from Fedorova et al (1994).

For Location, a data frame of the latitude and longitude locational data for 4833 samples.
ID2 Unique, sequential number assigned by NAMPD.
Latitude Latitude of the sampling location in decimal degrees.
Longitude Longitude of the sampling location in decimal degrees.
For Climate, a data frame with 4833 observations on the following 32 variables.
ID2 Unique, sequential number assigned by NAMPD.
$t[j a n: d e c]$ numeric vectors; Mean monthly temperatures for the indicated month. Degrees C.
$\mathrm{p}[j \mathrm{an}: \mathrm{dec}]$ numeric vectors; Mean total monthly precipitation (mm) for the indicated month.
tave numeric; annual average temperature (Degrees C)
tmax numeric; maximum temperature (in Degrees C) observed over the period of record.
tmin numeric; minimum temperature (in Degrees $C$ ) observed over the period of record.
gdd0 numeric; Growing degree days computed using a base of 0 degrees C .
gdd5 numeric; Growing degree days computed using a base of 5 degrees C.
mtco numeric; mean temperature of the coldest month.
mtwa numeric; mean temperature of the warmest month.
annp numeric; mean annual total precipitation (mm).

## Details

These datasets were extracted from Version 1.7 of the North American Modern Pollen Database.
All pollen species were included, however, only the Vegetation type (Biome) field of the AVHRR data and selected Climatic variables were extracted. Requests for additional variables to be included in the versions of the data included in the package should me sent to the package maintainer.

## Warning

Note that the data for the pollen species are a mixture of types. The DataForm variable contains information on the type of data included for each site. The codes are:
$\mathbf{R C}$ raw counts
$\mathbf{R P}$ raw counts expressed as percentages
DC digitised counts
DP digitised counts expressed as percentages
PM counts in permille
This value is not known for all samples.

## Source

The database is archived electronically at two sites:
http://www.lpc.uottawa.ca/data/modern/index.html
https://williamspaleolab.github.io/datavis/

## References

Whitmore, J., Gajewski, K., Sawada, M., Williams, J.W., Shuman, B., Bartlein, P.J., Minckley, T., Viau, A.E., Webb III, T., Anderson, P.M., and Brubaker L.B., 2005. North American and Greenland modern pollen data for multi-scale paleoecological and paleoclimatic applications. Quaternary Science Reviews 24:1828-1848.
Williams, J.W., Shuman, B., Bartlein, P.J., Whitmore, J., Gajewski, K., Sawada, M., Minckley, T., Shafer, S., Viau, A.E., Webb, III, T., Anderson, P.M., Brubaker, L.B., Whitlock, C. and Davis, O.K., 2006. An Atlas of Pollen-Vegetation-Climate Relationships for the United States and Canada. American Association of Stratigraphic Palynologists Foundation, Dallas, TX, 293p.

Williams, J.W. and Shuman, B., 2008. Obtaining accurate and precise environmental reconstructions from the modern analog technique and North American surface pollen dataset. Quaternary Science Reviews, 27: 669-687.
Fedorova, I.T., Volkova, Y.A., Varlyguin, E., 1994. World vegetation cover. Digital raster data on a 30-minute cartesian orthonormal geodetic (lat/long) $1080 \times 2160$ grid. In: Global Ecosystems Database Version 2.0. USDOC/NOAA National Geophysical Data Center, Bould, CO.

## Examples

```
data(Pollen)
data(Climate)
data(Biome)
data(Location)
```


## Description

A principal curve is a non-parametric generalisation of the principal component and is a curve that passes through the middle of a cloud of data points for a certain definition of 'middle'.

## Usage

```
prcurve(X, method = c("ca", "pca", "random", "user"), start = NULL,
    smoother = smoothSpline, complexity, vary = FALSE,
    maxComp, finalCV = FALSE, axis = 1, rank = FALSE,
        stretch = 2, maxit = 10, trace = FALSE, thresh = 0.001,
        plotit = FALSE, ...)
    initCurve(X, method = c("ca", "pca", "random", "user"), rank = FALSE,
        axis = 1, start)
```


## Arguments

X a matrix-like object containing the variables to which the principal curve is to be fitted.
method character; method to use when initialising the principal curve. "ca" fits a correspondence analysis to $X$ and uses the axis-th axis scores as the initial curve. "pca" does the same but fits a principal components analysis to X . "random" produces a random ordering as the initial curve.
start numeric vector specifying the initial curve when method = "user". Must be of length $\operatorname{nrow}(X)$.
smoother function; the choice of smoother used to fit the principal curve. Currently, the only options are smoothSpline, which is a wrapper to smooth.spline, and smoothGAM, which is a wrapper to gam.
complexity numeric; the complexity of the fitted smooth functions.
The function passed as argument smoother should arrange for this argument to be passed on to relevant aspect of the underlying smoother. In the case of smoothSpline, complexity is the df argument of smooth. spline.
vary logical; should the complexity of the smoother fitted to each variable in $X$ be allowed to vary (i.e. to allow a more or less smooth function for a particular variable. If FALSE the median complexity over all $m$ variables is chosen as the fixed complexity for all $m$ smooths.
maxComp numeric; the upper limt on the allowed complexity.
finalCV logial; should a final fit of the smooth function be performed using cross validation?
axis
numeric; the ordinaion axis to use as the initial curve.

| rank | logical; should rank position on the gradient be used? Not yet implemented. <br> stretch <br> numeric; a factor by which the curve can be extrapolated when points are pro- <br> jected. Default is 2 (times the last segment length). |
| :--- | :--- |
| maxit | numeric; the maximum number of iterations. |
| trace | logical; print progress on the iterations be printed to the console? <br> thresh |
| numeric; convergence threshold on shortest distances to the curve. The algo- |  |
| rithm is considered to have converged when the latest iteration produces a total |  |
| residual distance to the curve that is within thresh of the value obtained during |  |
| the previous iteration. |  |

## Value

An object of class "prcurve" with the following components:

| s | a matrix corresponding to $X$, giving their projections onto the curve. |
| :--- | :--- |
| tag | an index, such that s[tag, ] is smooth. |
| lambda | for each point, its arc-length from the beginning of the curve. |
| dist | the sum-of-squared distances from the points to their projections. |
| converged | logical; did the algorithm converge? |
| iter | numeric; the number of iterations performed. |
| totalDist | numeric; total sum-of-squared distances. |
| complexity | numeric vector; the complexity of the smoother fitted to each variable in X. |
| call | the matched call. <br> ordination |
| an object of class "rda", the result of a call to rda. This is a principal compo- |  |
| nents analysis of the input data $X$. |  |

## Note

The fitting function uses function project_to_curve in package princurve to find the projection of the data on to the fitted curve.

## Author(s)

Gavin L. Simpson

## See Also

smoothGAM and smoothSpline for the wrappers fitting smooth functions to each variable.

## Examples

```
    ## Load Abernethy Forest data set
    data(abernethy)
    ## Remove the Depth and Age variables
    abernethy2 <- abernethy[, -(37:38)]
    ## Fit the principal curve using the median complexity over
    ## all species
    aber.pc <- prcurve(abernethy2, method = "ca", trace = TRUE,
        vary = FALSE, penalty = 1.4)
    ## Extract fitted values
    fit <- fitted(aber.pc) ## locations on curve
    abun <- fitted(aber.pc, type = "smooths") ## fitted response
    ## Fit the principal curve using varying complexity of smoothers
    ## for each species
    aber.pc2 <- prcurve(abernethy2, method = "ca", trace = TRUE,
        vary = TRUE, penalty = 1.4)
    ## Predict new locations
    take <- abernethy2[1:10, ]
    pred <- predict(aber.pc2, take)
    ## Not run:
    ## Fit principal curve using a GAM - currently slow ~10secs
    aber.pc3 <- prcurve(abernethy2 / 100, method = "ca", trace = TRUE,
        vary = TRUE, smoother = smoothGAM, bs = "cr", family = mgcv::betar())
    ## End(Not run)
```

    predict.logitreg Posterior probability of analogue-ness for fossil samples
    
## Description

Predict the posterior probability of analogue-ness between fossil and training set samples based on logistic regression fits.

## Usage

```
## S3 method for class 'logitreg'
predict(object, newdata, group = "all", k = 1, ...)
```


## Arguments

| object | an object of class "logitreg", resulting from a call to logitreg. |
| :--- | :--- |
| newdata |  |
| an object of class "distance" representing the dissimilarity between fossil sam- |  |
| ples and those from the training set. |  |
| character; for which group(s) are predictions sought. "all", the default, indi- |  |
| cates that predictions for all groups and the combined analysis are produced. |  |
| Currently ignored. |  |$\quad$| numeric; the number of close modern analogues to each fossil sample to con- |
| :--- |
| sider. Not currently used and may be removed from the method as varying k |
| should require refitting the logistic regression model with that number of close |
| modern analogues considered. |

## Value

A matrix of posterior probabilities of analogue-ness is returned. The rows represent the fossil samples and the columns the groupings. There are $g+1$ columns where $g$ is the number of groups. The $g+1$ th column represents the Combined analysis which is the overall posterior probability that a fossil sample is an analogue to a training set sample (i.e. to any one of the groups).

## Author(s)

Gavin L. Simpson

## See Also

See logitreg for example usage. cma for extraction of close modern analogue information and analog for the underlying computations.

## Description

Predicted values based on a MAT model object.

## Usage

```
## S3 method for class 'mat'
predict(object, newdata, k, weighted = FALSE,
        bootstrap = FALSE, n.boot = 1000,
        probs = c(0.01, 0.025, 0.05, 0.1), ...)
```


## Arguments

| object | an object of mat. |
| :--- | :--- |
| newdata | data frame; required only if predictions for some new data are required. Mst <br> have the same number of columns, in same order, as x in mat. See example <br> below or join for how to do this. If newdata not provided, the fitted values are <br> returned. |
| k number of analogues to use. If missing, k is chosen automatically as the k that |  |
| achieves lowest RMSE. |  |
| weighted | logical; should the analysis use the weighted mean of environmental data of <br> analogues as predicted values? |
| bootstrap | logical; should bootstrap derived estimates and sample specific errors be calculated- <br> ignored if newdata is missing. |
| n.boot | numeric; the number of bootstrap samples to take. |
| probs | numeric; vector of probabilities with values in $[0,1]$. <br> arguments passed to of from other methods. |

## Details

This function returns one or more of three sets of results depending on the supplied arguments:
Fitted values: the fitted values of the mat model are returned if newdata is missing.
Predicted values: the predicted values for some new samples are returned if newdata is supplied. Summary model statistics and estimated values for the training set are also returned.
Bootstrap predictions and standard errors: if newdata is supplied and bootstrap = TRUE, the predicted values for newdata plus bootstrap estimates and standard errors for the new samples and the training set are returned.

The latter is simply a wrapper for bootstrap(model, newdata, . . .) - see bootstrap.mat.

## Value

A object of class predict.mat is returned if newdata is supplied, otherwise an object of fitted.mat is returned. If bootstrap = FALSE then not all components will be returned.
observed vector of observed environmental values.
model a list containing the model or non-bootstrapped estimates for the training set. With the following components:

- estimatedestimated values for " $y$ ", the environment.
- residualsmodel residuals.
- r.squaredModel $R^{2}$ between observed and estimated values of " $y$ ".
- avg.biasAverage bias of the model residuals.
- max. biasMaximum bias of the model residuals.
- rmsepModel error (RMSEP).
- knumeric; indicating the size of model used in estimates and predictions.

| bootstrap | a list containing the bootstrap estimates for the training set. With the following components: <br> - estimatedBootstrap estimates for " $y$ ". <br> - residualsBootstrap residuals for " $y$ ". <br> - r. squaredBootstrap derived $R^{2}$ between observed and estimated values of "y". <br> - avg.biasAverage bias of the bootstrap derived model residuals. <br> - max. biasMaximum bias of the bootstrap derived model residuals. <br> - rmsepBootstrap derived RMSEP for the model. <br> - s1Bootstrap derived S1 error component for the model. <br> - s2Bootstrap derived S2 error component for the model. <br> - knumeric; indicating the size of model used in estimates and predictions. |
| :---: | :---: |
| sample.errors | a list containing the bootstrap-derived sample specific errors for the training set. With the following components: <br> - rmsepBootstrap derived RMSEP for the training set samples. <br> - s1Bootstrap derived S1 error component for training set samples. <br> - s2Bootstrap derived S2 error component for training set samples. |
| weighted | logical; whether the weighted mean was used instead of the mean of the environment for $k$-closest analogues. |
| auto | lo |
| n. boot | numeric; the number of bootstrap samples taken. |
| predictions | a list containing the model and bootstrap-derived estimates for the new data, with the following components: <br> - observedthe observed values for the new samples - only if newenv is provided. <br> - modela list containing the model or non-bootstrapped estimates for the new samples. A list with the same components as model, above. <br> - bootstrapa list containing the bootstrap estimates for the new samples, with some or all of the same components as bootstrap, above. <br> - sample.errorsa list containing the bootstrap-derived sample specific errors for the new samples, with some or all of the same components as sample.errors, above. |
| method | the dissimilarity measure used to fit the underlying MAT models. |
| quantiles | probability quantiles of the pairwise dissimilarities computed from the training set. |
| minDC | smallest distances between each sample in newdata and the training set samples. |
| Dij | dissimilarities between newdata and training set samples. |

## Author(s)

Gavin L. Simpson

## References

Birks, H.J.B., Line, J.M., Juggins, S., Stevenson, A.C. and ter Braak, C.J.F. (1990). Diatoms and pH reconstruction. Philosophical Transactions of the Royal Society of London; Series B, 327; 263-278.

## See Also

mat, bootstrap.mat

## Examples

```
## Imbrie and Kipp example
## load the example data
data(ImbrieKipp)
data(SumSST)
data(V12.122)
## merge training and test set on columns
dat <- join(ImbrieKipp, V12.122, verbose = TRUE)
## extract the merged data sets and convert to proportions
ImbrieKipp <- dat[[1]] / 100
V12.122 <- dat[[2]] / 100
## fit the MAT model using the chord distance measure
(ik.mat <- mat(ImbrieKipp, SumSST, method = "chord"))
## predict for V12.122 data
predict(ik.mat, V12.122)
```

predict.pcr

Predicted values from a principal components regression

## Description

Calculates predicted values from a fitted principal components regression model. Leave-one-out, bootstrap or n k -fold crossvalidated predictions are also implemented.

## Usage

```
## S3 method for class 'pcr'
predict(object, newdata, ncomp = object$ncomp,
    CV = c("none", "LOO", "bootstrap", "kfold"),
    verbose = FALSE, nboot = 100, kfold = 10, folds = 5,
    ...)
```


## Arguments

| object | a fitted model of class "pcr", the result of a call to pcr. |
| :--- | :--- |
| newdata | data frame of new observations for which predictions are sought. |
| ncomp | numeric; the PCR components for which predictions are sought. If ncomp = c, <br> predictions for components 1:c are produced. <br> character; the type of crossvalidation required. Currently, no crossvalidation <br> methods are implemented. |
| CV | logical; should progress on crossvalidation be printed to the console? |
| verbose | numeric; the number of bootstrap samples to draw. |
| nboot | numeric; the number of folds to split data into. |
| kfold | numeric; the number of repetitions of k-fold CV. |
| folds | arguments passed to other methods. |

## Details

predict.por arranges for any transformation applied to the training data to be applied to the newdata prior to prediction.

## Value

A matrix of predicted values with rows representing samples in newdata and columns, the PCR components requested via ncomp.

## Author(s)

Gavin L. Simpson

## See Also

pcr

## Examples

```
## Load the Imbrie & Kipp data and
## summer sea-surface temperatures
data(ImbrieKipp)
data(SumSST)
## choose 10 samples to act as a test set, for illustration
take <- c(5,58,31,51,42,28,30,57,8,50)
## normal interface and apply Hellinger transformation
mod <- pcr(ImbrieKipp[-take, ], SumSST[-take], tranFun = Hellinger)
## predictions
predict(mod, ImbrieKipp[take, ], ncomp = 4)
## predictions
```

```
set.seed(123)
predict(mod, ImbrieKipp[take, ], ncomp = 4, CV = "bootstrap",
    nboot = 100)
```

predict.prcurve Predict news locations $\backslash \&$ fitted values on a principal curve

## Description

Locations on a fitted principal curve are predicted by projecting the new observations in $m$ dimensions on to the corresponding closest point on the curve. Fitted values for data used to fit the curve are available with respect to the principal curve or to the individual smooth functions.

## Usage

```
## S3 method for class 'prcurve'
predict(object, newdata, ...)
## S3 method for class 'prcurve'
fitted(object, type = c("curve","smooths"), ...)
```


## Arguments

object an object of class prcurve.
newdata a matrix or data frame of new observations within the space of the orginal data. Variables are matched against those of the original data via their names or colnames. If a data frame is supplied, it is converted to a matrix via data. matrix.
type character; the type of fitted values to return.
. . . other arguments passed to other methods. Not currently used.

## Details

Fitting a principal curve involves two procedures. In one, the current curve is bent towards the data via the fitting of spline functions with distance along the curve as the predictor variable and each variable in turn as the response. The second procedure, a projection step, involves projecting the observed points in $m$ dimensions on to locations along the current curve to which they are closest in the hyperspace.
Given a fitted curve, the projection step can be used to find new points on the fitted curve by projecting the new points located in the hyperspace on to points on the curve to which they are closest.

Fitted values are available for the data used to the fit the principal curve. There are two types of fitted value available. For type = "curve", the fitted locations on the principal curve. For type $=$ "smooths", the fitted values of the variables from the individual smooth functions with respect to distance along the principal curve.

## Value

A matrix of points in the space of the original data. Rows correspond to the new samples and columns to the variables (ordered as per the original data used to fit the curve).
How these points are ordered along the fitted curve is contained in attributed tag.

## Author(s)

Gavin L. Simpson

## See Also

See prcurve for details on fitting principal curves and an example.

```
predict.wa Predict from a weighted average model
```


## Description

Model predictions and cross-validation predictions for weighted averaging transfer function models.

## Usage

```
## S3 method for class 'wa'
    predict(object, newdata,
            CV = c("none", "LOO", "bootstrap", "nfold"),
            verbose = FALSE, n.boot = 100, nfold = 5, ...)
```


## Arguments

object an object of class "wa", usually the result of a call to wa
newdata An optional data frame in which to look for variables with which to predict. If omitted, the fitted values are used.
CV Should cross-validation be performed? Leave-one-out ("LOO"), bootstrap ("bootstrap") and $k$-fold ("nfold") CV are currently available.
verbose $\quad$ Should CV progress be printed to the console?
n.boot The number of bootstrap samples or $k$-fold steps.
nfold $\quad$ Number of subsets in $k$-fold CV.
... further arguments passed to or from other methods.

## Details

Not all CV methods produce the same output. CV = "bootstrap" and CV = "nfold" produce sample specific errors.

## Value

An object of class "predict.wa", a list with the following components:
pred A list with components pred and rmsep containing the predicted values and the sample specific errors if available.
performance A list with model performance statistics.
model.pred A list with components pred and rmsep containing the predicted values for the training set samples and the sample specific errors if available.
call the matched function call.
CV.method The CV method used.

## Author(s)

Gavin L. Simpson and Jari Oksanen ( $k$-fold CV)

## References

Birks, H.J.B., Line, J.M., Juggins, S., Stevenson, A.C. and ter Braak, C.J.F. (1990). Diatoms and pH reconstruction. Philosophical Transactions of the Royal Society of London; Series B, 327; 263-278.

## See Also

wa, predict.mat, performance, reconPlot.

## Examples

```
## Imbrie and Kipp
data(ImbrieKipp)
ImbrieKipp <- ImbrieKipp / 100
data(SumSST)
ik.wa <- wa(SumSST ~ ., data = ImbrieKipp, tol.dw = TRUE,
    min.tol = 2, small.tol = "min")
ik.wa
## load V12.122 core data
data(V12.122)
V12.122 <- V12.122 / 100
## predict summer sea-surface temperature for V12.122 core
set.seed(2)
v12.pred <- predict(ik.wa, V12.122, CV = "bootstrap", n.boot = 100)
## draw the fitted reconstruction
reconPlot(v12.pred, use.labels = TRUE, display = "bars")
## extract the model performance stats
performance(v12.pred)
```


## Description

Computes the rank correlation between distances between sites in terms of gradient variables, such as environmental ones, and species composition.

## Usage

```
rankDC(g, y, dc = c("chord", "bray", "euclidean", "chi.square", "gower"),
        method = "spearman")
## S3 method for class 'rankDC'
plot(x, sort = TRUE, decreasing = FALSE,
        xlab = "Rank correlation", color = "blue",
        pch = 21, bg = "blue", lcolor = "grey",
        lty = "solid", ...)
    ## S3 method for class 'rankDC'
    dotplot(x, data = NULL, sort = TRUE, decreasing = FALSE,
        xlab = "Rank correlation", ...)
```


## Arguments

| g | the gradient values; usually a data frame of environmental data. |
| :---: | :---: |
| y | the species data; usually a data frame. |
| dc | character; the set of dissimilarity coefficients for which rank correlations with gradient distance are to be computed. |
| method | character; the correlation method to use. See the method argument of cor. |
| x | an object of class "rankDC". |
| data | NULL; ignored, only present for purposes of conformance to generic definition. |
| sort, decreasing |  |
|  | logical; should observations be sorted prior to plotting? If so, should they be sorted in order of decreasing size? |
| xlab | The x-axis label for the plot. |
| color, | lcolor, lty arguments passed to dotchart. |
|  | arguments passed to other methods, including dotchart and dotplot. |

## Value

A named vector of rank correlations is returned.

## Author(s)

Gavin L. Simpson, based on rankindex from the vegan package.

## See Also

rankindex from package vegan. For the dotplot method, see dotplot.

## Examples

```
data(swappH)
data(swapdiat)
rc <- rankDC(swappH, swapdiat, dc = c("chord","euclidean","gower"))
## base plot uses dotchart()
plot(rc)
## lattice version of the base plot
dotplot(rc)
```

    reconPlot Stratigraphic plots of palaeoenvironmental reconstructions
    
## Description

Draws a palaeoenvironmental reconstruction of predicted environmental values for sub-fossil assemblages.

## Usage

```
reconPlot(x, ...)
## Default S3 method:
reconPlot(x, depths, errors,
    display.error = c("none", "bars", "lines"),
    rev.x = TRUE,
    col.error = "grey", lty.error = "dashed",
    type = "l",
    xlim, ylim,
    xlab = "", ylab = "", main = "",
    ...)
## S3 method for class 'predict.mat'
reconPlot(x, depths, use.labels = FALSE,
    predictions = c("model", "bootstrap"),
    display.error = c("none", "bars", "lines"),
```

```
    sample.specific = TRUE, ...)
## S3 method for class 'predict.wa'
reconPlot(x, depths, use.labels = FALSE,
    display.error = c("none", "bars", "lines"),
    sample.specific = TRUE, ...)
```


## Arguments

```
x An R object.
depths numeric; a vector of depths for which predicted values exist or will be generated.
    Can be missing, in which case, if use.labels = TRUE, the function will attempt
    to derive suitable values for you. See Details below.
errors numeric; a vector of errors for plotting error bars or lines.
display.error character; hown should error bars be drawn on the plot? One of "none", "bars",
    or "lines". If "bars", error bars are drawn for each sample. If "lines", lines
    are drawn enclosing the region prediction +/- RMSEP.
rev.x logical; should the depth/age axis be reversed (drawn from high to low)?
col.error, lty.error
    the colour and type of line drawn. See par and arguments "col" and "lty".
type type of line drawn. See par and argument "type".
xlab,ylab character; the x- and y-axis labels respectively.
main character; main title for the plot.
xlim, ylim numeric, length 2; the x- and y-limits for the plotted axes. If not provided, the
    function will calculate appropriate values to cover the range of plotted values
    and any error bars (if requested via "error.bars".
use.labels logical; should reconPlot attempt to derive values for argument depths from
    the names of the predicted values? Only use if depths is missing. See Details
    below.
predictions character; one of "model" or "bootstrap". Which type of predicted values
    should be plotted? The model predictions ("model") or the bootstrap-derived
    predictions ("bootstrap").
sample.specific
    logical; should sample specific errors be used? Only for predictions = "bootstrap".
... arguments passed to other graphics functions.
```


## Details

Conventionally, these plots are drawn on a depth or an age scale. Argument depths is used to provide the depth or age axis, against which the predicted values are plotted.

If depths is not provided, then the function will try to derive the appropriate values from the labels of the predictions if use.labels = TRUE. You must provide depths or set use.labels = TRUE otherwise an error will result. The derived labels will be coerced to numerics. If your labels are not coercible, then you'll either get nonsense on the plot or an error from R. If so, provide suitable values for depths.

## Value

A plot on the currently active device.

## Author(s)

Gavin L. Simpson

## See Also

mat, and predict.mat for MAT transfer functions and wa and predict. wa for WA models.

## Examples

```
## Imbrie and Kipp example
## load the example data
data(ImbrieKipp)
data(SumSST)
data(V12.122)
## merge training and test set on columns
dat <- join(ImbrieKipp, V12.122, verbose = TRUE)
## extract the merged data sets and convert to proportions
ImbrieKipp <- dat[[1]] / 100
V12.122 <- dat[[2]] / 100
## Fit a MAT model
(ik.mat <- mat(ImbrieKipp, SumSST, method = "chord"))
## Reconstruct pH for the RLGH core
v12.pH <- predict(ik.mat, V12.122)
## draw the reconstruction
reconPlot(v12.pH, use.labels = TRUE, display.error = "bars",
    xlab = "Depth", ylab = "Summer Seas-surface Temperature")
```

residLen Squared residual length diagnostics

## Description

The squared residual length between the fitted values of a constrained ordination and the original species data is one diagnostic for transfer function models.

## Usage

```
    residLen(X, env, passive, method = c("cca","rda"))
    fittedY(ord, newdata, colsum)
    sqrlLinear(Y, fitted)
    sqrlUnimodal(Y, colsum, fitted)
```


## Arguments

$X \quad$ data frame; the training set species data.
env vector; the training set environmental data.
passive data frame; the passive samples species data.
method the ordination technique to use. One of "rda" or "cca", with the latter the default.
ord an ordination object, the result of a call to cca or rda.
newdata Species data matrix for passive samples. Must have same columns as data used to fit ord.
colsum column (species) sums for training set data used to fit ord.
Y Original species data matrix, the response for which squared residual lengths are to be computed.
fitted The fitted values of the response derived from the constrained ordination model.

## Details

The squared residual lengths are computed for the training set samples and the passive samples separately. Passive samples that are poorly fitted in the transfer function model will have large squared residual distances between the observed species data and the fitted values from the constrained ordination.
residLen is the main user-interface function and can be called with either the training data and passive samples.
fittedY returns the fitted approximation of the passive sample response data (i.e. species data). sqrlLinear and sqrlUnimodal return the squared residual distances between the observed species data and the fitted values from the constrained ordination model.

## Value

fittedY returns a matrix of fitted species abundances for passive samples.
sqrlLinear and sqrlUnimodal return a vector of residual distances.
residLen returns an object of class "residLen" with the attribute "method" set to "method". This object is a list with the following components:
train, passive The squared residual lengths for the training set and the passive samples.
ordination The fitted ordination.
call The matched call.

## Author(s)

Gavin L. Simpson

## References

Ter Braak C.J.F. and Smilauer P. (2002) CANOCO Reference manual and CanoDraw for Windows User's guide: Software for Canonical Ordination (version 4.5). Microcomputer Power (Ithaca, NY, USA), 500pp.

## See Also

cca and predict.cca for some of the underlying computations.

## Examples

```
## load the Imbrie and Kipp example data
data(ImbrieKipp, SumSST, V12.122)
## squared residual lengths for Core V12.122
rlens <- residLen(ImbrieKipp, SumSST, V12.122)
rlens
## as before but using linear RDA
residLen(ImbrieKipp, SumSST, V12.122, method = "rda")
```

residuals.prcurve Residuals of a principal curve fit.

## Description

Returns various representations of the residuals of a principal curve fit.

## Usage

```
## S3 method for class 'prcurve'
residuals(object, which = c("distance", "raw", "smooths", "pca"),
    ...)
```


## Arguments

object an object of class "prcurve", the result of a call to prcurve.
which character; the type of residuals to return. See Details.
... arguments passed to other methods. See Details.

## Details

Various types of residual are available for the principal curve. In a departure from the usual convention, which residuals are returned is controlled via the which argument. This is to allow users to pass a type argument to the residuals method for the function used to fit the individual smooth functions when which = "smooths".
The types of residuals available are
"distance" the default residual for a principal curve. This residual is taken as the Euclidean distance between each observations and the point on the principal curve to which it projects, in full multivariate space.
"raw" raw residuals are the basis for "distance" residuals, and are the difference between the observed and fitted values (position on the curve) for each observation in terms of each variable in the data set. These residuals are in the form of a matrix with number of observation rows and number of variables cols.
"smooths" these residuals are the result of calling residuals() on each of the smooth models fitted to the individual variables. See below for further details. A matrix of the same dimensions as for which = "raw" is returned.
"pca" similar to the raw residuals, but expressed in terms of the principal components of the input data. Hence these residuals are the difference between each observation's location in PCA space and their corresponding location on the curve.

For "smooths" residuals, what is returned is governed by the residuals method available for the smooth model fitted to the individual variables. For principal curves fitted using the smoothSpline plugin, see smooth.spline. For principal curves fitted via the smoothGAM plugin, see residuals.gam.
...can be used to pass additional arguments to these residuals methods. In particular, the type argument is commonly used to choose which type of residual is returned by the specific methods.
In the case of principal curves fitted using the plugin smoothSpline, residuals for which = "smooths" are only available if the the additional argument keep. data was specified during fitting via prcurve. See the examples for an illustration of this usage.

## Value

A vector of residual distances (which = "distance") or a matrix of residuals (for the other options).

## Author(s)

Gavin L. Simpson

## See Also

prcurve for fitting a principal curve.

## Examples

```
## Load Abernethy Forest data set
data(abernethy)
```

```
## Remove the Depth and Age variables
abernethy2 <- abernethy[, -(37:38)]
## Fit the principal curve, preserving the data in the smooth.spline
## smooth functions fitted via keep.data = TRUE
aber.pc <- prcurve(abernethy2, method = "ca", keep.data = TRUE)
## default "distance" residuals
res <- resid(aber.pc)
head(res)
## residuals from the underlying smooth models, also illustrates
## how to select specific types of residual from the individual
## method using argument 'type'
res <- resid(aber.pc, which = "smooths", type = "deviance")
dim(res)
head(res[, 1:5]) # just show a few species residuals
```

```
rlgh Round Loch of Glenhead Diatoms
```


## Description

Diatom sub-fossil assemblage counts from the Round Loch of Glenhead.

## Usage

data(rlgh)

## Format

A data frame with 101 observations on 139 diatom species.

## Details

The samples are taken from various depths down a sediment core retrieved from the Round Loch of Glenhead, Galloway, Scotland.

## References

Jones, V.J., Stevenson, A.C. and Battarbee, R.W. (1989) Acidification of lakes in Galloway, south west Scotland - A diatom and pollen study of the post-glacial history of the Round Loch of Glenhead. Journal of Ecology 77(1), 1-23.

## Examples

```
data(rlgh)
```


## Description

Calculates or extracts the RMSEP from transfer function models.

## Usage

```
    RMSEP(object, ...)
```

    \#\# S3 method for class 'mat'
    RMSEP (object, \(k\), weighted \(=\) FALSE,
            ...)
    \#\# S3 method for class 'bootstrap.mat'
    RMSEP (object, type = c("birks1990", "standard"),
            ...)
    \#\# S3 method for class 'bootstrap.wa'
    RMSEP(object, type = c("birks1990", "standard"),
        ...)
    
## Arguments

object An R object.
k
numeric; the number of analogues to use in calculating the RMSEP. May be missing. If missing, $k$ is extracted from the model using getK.
weighted logical; Return the RMSEP for the weighted or unweighted model? The default is for an unweighted model.
type The type of RMSEP to return/calculate. See Details, below.
... Arguments passed to other methods.

## Details

There are two forms of RMSEP in common usage. Within palaeoecology, the RMSEP of Birks et al. (1990) is most familiar:

$$
\mathrm{RMSEP}=\sqrt{s_{1}^{2}+s_{2}^{2}}
$$

where where $s_{1}$ is the standard deviation of the out-of-bag ( OOB ) residuals and $s_{2}$ is the mean bias or the mean of the OOB residuals.
In the wider statistical literature, the following form of RMSEP is more commonly used:

$$
\mathrm{RMSEP}=\sqrt{\frac{\sum_{i=1}^{n}\left(y_{i}-\hat{y}_{i}\right)^{2}}{n}}
$$

where $y_{i}$ are the observed values and $\hat{y}_{i}$ the transfer function predictions/fitted values.
The first form of RMSEP is returned by default or if type = "birks1990" is supplied. The latter form is returned if type = "standard" is supplied.
The RMSEP for objects of class "mat" is a leave-one-out cross-validated RMSEP, and is calculated as for type = "standard".

## Value

A numeric vector of length 1 that is the RMSEP of object.

## Author(s)

Gavin L. Simpson

## References

Birks, H.J.B., Line, J.M., Juggins, S., Stevenson, A.C. and ter Braak, C.J.F. (1990). Diatoms and pH reconstruction. Philosophical Transactions of the Royal Society of London; Series B, 327; 263-278.

## See Also

mat, bootstrap, wa, bootstrap.wa.

## Examples

```
## Imbrie and Kipp example
## load the example data
data(ImbrieKipp)
data(SumSST)
data(V12.122)
## merge training and test set on columns
dat <- join(ImbrieKipp, V12.122, verbose = TRUE)
## extract the merged data sets and convert to proportions
ImbrieKipp <- dat[[1]] / 100
V12.122 <- dat[[2]] / 100
## fit the MAT model using the squared chord distance measure
(ik.mat <- mat(ImbrieKipp, SumSST, method = "chord"))
## Leave-one-out RMSEP for the MAT model
RMSEP(ik.mat)
## bootstrap training set
(ik.boot <- bootstrap(ik.mat, n.boot = 100))
## extract the Birks et al (1990) RMSEP
RMSEP(ik.boot)
```

```
## Calculate the alternative formulation
RMSEP(ik.boot, type = "standard")
```

roc ROC curve analysis

## Description

Fits Receiver Operator Characteristic (ROC) curves to training set data. Used to determine the critical value of a dissimilarity coefficient that best descriminate between assemblage-types in palaeoecological data sets, whilst minimising the false positive error rate (FPF).

## Usage

```
roc(object, groups, k = 1, ...)
## Default S3 method:
roc(object, groups, k = 1, thin = FALSE,
    max.len = 10000, ...)
## S3 method for class 'mat'
roc(object, groups, k = 1, ...)
    ## S3 method for class 'analog'
    roc(object, groups, k = 1, ...)
```


## Arguments

object an R object.
groups a vector of group memberships, one entry per sample in the training set data. Can be a factor, and will be coerced to one if supplied vecvtor is not a factor.
$k \quad$ numeric; the $k$ closest analogues to use to calculate ROC curves.
thin logical; should the points on the ROC curve be thinned? See Details, below.
max.len numeric; length of analolgue and non-analogue vectors. Used as limit to thin points on ROC curve to.
... arguments passed to/from other methods.

## Details

A ROC curve is generated from the within-group and between-group dissimilarities.
For each level of the grouping vector (groups) the dissimilarity between each group member and it's k closest analogues within that group are compared with the k closest dissimilarities between the non-group member and group member samples.

If one is able to discriminate between members of different group on the basis of assemblage dissimilarity, then the dissimilarities between samples within a group will be small compared to the dissimilarities between group members and non group members.
thin is useful for large problems, where the number of analogue and non-analogue distances can conceivably be large and thus overflow the largest number R can work with. This option is also useful to speed up computations for large problems. If thin $==$ TRUE, then the larger of the analogue or non-analogue distances is thinned to a maximum length of max. len. The smaller set of distances is scaled proportionally. In thinning, we approximate the distribution of distances by taking max. len (or a fraction of max. len for the smaller set of distances) equally-spaced probability quantiles of the distribution as a new set of distances.

## Value

A list with two components; i, statistics, a summary of ROC statistics for each level of groups and a combined ROC analysis, and ii, roc, a list of ROC objects, one per level of groups. For the latter, each ROC object is a list, with the following components:

| TPF | The true positive fraction. |
| :--- | :--- |
| FPE | The false positive error |
| optimal | The optimal dissimilarity value, asessed where the slope of the ROC curve is <br> maximal. |
| AUC | The area under the ROC curve. |
| se.fit | Standard error of the AUC estimate. |
| n. in | numeric; the number of samples within the current group. |
| n. out | numeric; the number of samples not in the current group. <br> p.value p-value of a Wilcoxon rank sum test on the two sets of dissimilarities. This <br> is also known as a Mann-Whitney test. |
| roc.points | The unique dissimilarities at which the ROC curve was evaluated <br> mameric; the position along the ROC curve at which the slope of the ROC curve |
| prior | numaximal. This is the index of this point on the curve. <br> is mamer <br> numeric, length 2. Vector of observed prior probabilities of true analogue and <br> true non-analogues in the group. |
| analogue | a list with components yes and no containing the dissimilarities for true ana- <br> logue and true non-analogues in the group. |

## Author(s)

Gavin L. Simpson, based on code from Thomas Lumley to optimise the calculation of the ROC curve.

## References

Brown, C.D., and Davis, H.T. (2006) Receiver operating characteristics curves and related decision measures: A tutorial. Chemometrics and Intelligent Laboratory Systems 80, 24-38.

Gavin, D.G., Oswald, W.W., Wahl, E.R. and Williams, J.W. (2003) A statistical approach to evaluating distance metrics and analog assignments for pollen records. Quaternary Research 60, 356-367.

Henderson, A.R. (1993) Assessing test accuracy and its clinical consequences: a primer for receiver operating characteristic curve analysis. Annals of Clinical Biochemistry 30, 834-846.

## See Also

mat for fitting of MAT models. bootstrap.mat and mcarlo for alternative methods for selecting critical values of analogue-ness for dissimilarity coefficients.

## Examples

```
## load the example data
data(swapdiat, swappH, rlgh)
## merge training and test set on columns
dat <- join(swapdiat, rlgh, verbose = TRUE)
## extract the merged data sets and convert to proportions
swapdiat <- dat[[1]] / 100
rlgh <- dat[[2]] / 100
## fit an analogue matching (AM) model using the squared chord distance
## measure - need to keep the training set dissimilarities
swap.ana <- analog(swapdiat, rlgh, method = "SQchord",
    keep.train = TRUE)
## fit the ROC curve to the SWAP diatom data using the AM results
## Generate a grouping for the SWAP lakes
METHOD <- if (getRversion() < "3.1.0") {"ward"} else {"ward.D"}
clust <- hclust(as.dist(swap.ana$train), method = METHOD)
grps <- cutree(clust, 12)
## fit the ROC curve
swap.roc <- roc(swap.ana, groups = grps)
swap.roc
## draw the ROC curve
plot(swap.roc, 1)
## draw the four default diagnostic plots
layout(matrix(1:4, ncol = 2))
plot(swap.roc)
layout(1)
```

scores.prcurve scores method for principal curve objects of class "prcurve".

## Description

A scores method to extract the position on the curve to which each observation projects (display = "curve") or the coordinates of the curve in the dimensions of the input data (display = "dimensions").

## Usage

```
## S3 method for class 'prcurve'
scores(x, display = c("curve", "dimensions"), ...)
```


## Arguments

x
an object of class "prcurve", usually from a call to prcurve.
display character; which type of scores to extract. display = "curve" returns the position along the curve onto which each observation projects; this can be used like a PCA axis score. display = "dimensions" returns the coordinates of the curve in the dimensions of the original data.
... Arguments passed to other methods. Not used.

## Value

If display = "curve" a 1-column matrix is returned with a row for each observation in the input data. If display = "dimensions", a matrix of coordinates for the principal curve is returned. The dimensions of this matrix relate to the dimensions of the input data; if there were $n$ samples (rows) and $m$ variables columns then the matrix returned by scores.prcurve will have $n$ rows and $m$ columns.

## Author(s)

Gavin L. Simpson

## See Also

prcurve for fitting principal curves to data.

## Examples

```
## Load the Abernethy Forest data set
data(abernethy)
## Remove the Depth and Age variables
abernethy2 <- abernethy[, -(37:38)]
## Fit the principal curve using varying complexity of smoothers
## for each species
aber.pc <- prcurve(abernethy2, method = "ca", trace = TRUE,
    vary = TRUE, penalty = 1.4)
## Extract position on the curve
pos <- scores(aber.pc, display = "curve")
head(pos)
```

```
## Extract the coordinates of the curve
coord <- scores(aber.pc, display = "dimensions")
dim(coord)
all.equal(dim(coord), dim(abernethy2))
```

screeplot Screeplots of model results

## Description

Draws screeplots of performance statistics for models of varying complexity.

## Usage

```
## S3 method for class 'mat'
screeplot(x, k, restrict = 20,
    display = c("rmsep", "avg.bias",
            "max.bias", "r.squared"),
    weighted = FALSE, col = "red", xlab = NULL,
    ylab = NULL, main = NULL, sub = NULL, ...)
## S3 method for class 'bootstrap.mat'
screeplot(x, k, restrict = 20,
    display = c("rmsep","avg.bias","max.bias",
            "r.squared"),
    legend = TRUE, loc.legend = "topright",
    col = c("red", "blue"),
    xlab = NULL, ylab = NULL,
    main = NULL, sub = NULL,
    ...,
    lty = c("solid","dashed"))
```


## Arguments

x
$k$ number of analogues to use. If missing ' $k$ ' is chosen automatically as the ' $k$ ' that achieves lowest RMSE.
restrict logical; restrict comparison of k-closest model to $\mathrm{k}<=$ restrict.
display which aspect of $x$ to plot? Partial match.
weighted logical; should the analysis use weighted mean of env data of analogues as fitted/estimated values?
$x l a b, y l a b \quad x-$ and $y$-axis labels respectively.
main, sub main and subtitle for the plot.
\(\left.\begin{array}{ll}legend \& logical; should a legend be displayed on the figure? <br>
loc.legend <br>
character; a keyword for the location of the legend. See legend for details of <br>

allowed keywords.\end{array}\right]\)| Colours for lines drawn on the screeplot. Method for class "bootstrap.mat" |
| :--- |
| takes a vector of two colours. |

## Details

Screeplots are often used to graphically show the results of cross-validation or other estimate of model performance across a range of model complexity.

Four measures of model performance are currently available: i) root mean square error of prediction (RMSEP); ii) average bias - the mean of the model residuals; iii) maximum bias - the maximum average bias calculated for each of $n$ sections of the gradient of the environmental variable; and $v$ ) model $R^{2}$.
For the maximum bias statistic, the response (environmental) gradient is split into $n=10$ sections.
For the bootstrap method, apparent and bootstrap versions of these statistics are available and plotted.

## Note

Currently only models of class mat and bootstrap.mat are supported.

## Author(s)

Gavin Simpson

## See Also

screeplot

## Examples

```
## Imbrie and Kipp example
## load the example data
data(ImbrieKipp)
data(SumSST)
data(V12.122)
## merge training and test set on columns
dat <- join(ImbrieKipp, V12.122, verbose = TRUE)
## extract the merged data sets and convert to proportions
ImbrieKipp <- dat[[1]] / 100
V12.122 <- dat[[2]] / 100
```

```
## fit the MAT model using the chord distance measure
(ik.mat <- mat(ImbrieKipp, SumSST, method = "chord"))
screeplot(ik.mat)
```

smoothers

Smoother plugin function for use in fitting a principal curve

## Description

Functions to be used as plugins to prcurve that fit smooth functions to each variable that, when combined, give the principal curve. The functions act as wrappers to the main fitting functions, which currently include smooth. spline and gam.

## Usage

```
smoothSpline(lambda, x, choose = TRUE, complexity, ...,
                        penalty = 1, cv = FALSE, keep.data = FALSE,
                        control.spar = list(low = 0))
smoothGAM(lambda, x, choose = TRUE, complexity, bs = "tp", ...,
        family = gaussian(), method = "REML", select = FALSE,
        control = gam.control())
```


## Arguments

lambda the current projection function; the position that each sample projects to on the current principal curve. This is the predictor variable or covariate in the smooth function.
x
numeric vector; used as the response variable in the smooth function. The principal curve algorithm fits a separate scatterplot smoother (or similar smoother) to each variable in X in turn as the response.
choose logical; should the underlying smoother function be allowed to choose the degree of smooth complexity for each variable?
complexity numeric; the complexity of the fitted smooth functions.
penalty, cv, keep.data, control.spar
arguments to smooth. spline.
bs, family arguments to s.
method, select, control
arguments to gam.
... arguments passed on the the underlying function smooth. spline and users should read that function's help page for further details.

## Value

An object of class "prcurveSmoother" with the following components:
lambda for each observations, its arc-length from the beginning of the curve.
x numeric vector of response values.
fitted.values numeric vector of fitted values for the observations generated from the fitted smooth function.
complexity numeric; the degrees of freedom used for the smooth function. The exact details of what these pertain to are in the help for the respective fitting functions smooth. spline and gam.
model the object fitted by the wrapped fitting function.

## Author(s)

Gavin L. Simpson

## See Also

prcurve for how these functions are used.
splitSample Select samples from along an environmental gradient

## Description

Select samples from along an environmental gradient by splitting the gradient into discrete chunks and sample within each chunk. This allows a test set to be selected which covers the environmental gradient of the training set, for example.

## Usage

```
splitSample(env, chunk = 10, take, nchunk,
    fill = c("head", "tail", "random"),
    maxit = 1000)
```


## Arguments

env numeric; vector of samples representing the gradient values.
chunk numeric; number of chunks to split the gradient into.
take numeric; how many samples to take from the gradient. Can not be missing.
nchunk numeric; number of samples per chunk. Must be a vector of length chunk and sum(chunk) must equal take. Can be missing (the default), in which case some simple heuristics are used to determine the number of samples chosen per chunk. See Details.
fill character; the type of filling of chunks to perform. See Details.
maxit numeric; maximum number of iterations in which to try to sample take observations. Basically here to stop the loop going on forever.

## Details

The gradient is split into chunk sections and samples are selected from each chunk to result in a sample of length take. If take is divisible by chunk without remainder then there will an equal number of samples selected from each chunk. Where chunk is not a multiple of take and nchunk is not specified then extra samples have to be allocated to some of the chunks to reach the required number of samples selected.
An additional complication is that some chunks of the gradient may have fewer than nchunk samples and therefore more samples need to be selected from the remaining chunks until take samples are chosen.
If nchunk is supplied, it must be a vector stating exactly how many samples to select from each chunk. If chunk is not supplied, then the number of samples per chunk is determined as follows:

1. An intial allocation of floor (take / chunk) is assigned to each chunk
2. If any chunks have fewer samples than this initial allocation, these elements of nchunk are reset to the number of samples in those chunks
3. Sequentially an extra sample is allocated to each chunk with sufficient available samples until take samples are selected.

Argument fill controls the order in which the chunks are filled. fill = "head" fills from the low to the high end of the gradient, whilst fill = "tail" fills in the opposite direction. Chunks are filled in random order if fill = "random". In all cases no chunk is filled by more than one extra sample until all chunks that can supply one extra sample are filled. In the case of fill = "head" or fill = "tail" this entails moving along the gradient from one end to the other allocating an extra sample to available chunks before starting along the gradient again. For fill = "random", a random order of chunks to fill is determined, if an extra sample is allocated to each chunk in the random order and take samples are still not selected, filling begins again using the same random ordering. In other words, the random order of chunks to fill is chosen only once.

## Value

A numeric vector of indices of selected samples. This vector has attribute lengths which indicates how many samples were actually chosen from each chunk.

## Author(s)

Gavin L. Simpson

## Examples

```
data(swappH)
## take a test set of 20 samples along the pH gradient
test1 <- splitSample(swappH, chunk = 10, take = 20)
test1
swappH[test1]
## take a larger sample where some chunks don't have many samples
## do random filling
set.seed(3)
```

```
test2 <- splitSample(swappH, chunk = 10, take = 70, fill = "random")
test2
swappH[test2]
```

sppResponse

Species responses along gradients.

## Description

The fitted responses of species along gradients are estimated or extracted from appropriate objects.

## Usage

```
sppResponse(x, ...)
## S3 method for class 'prcurve'
sppResponse(x, n = 100, ...)
```


## Arguments

x
an R object.
$\mathrm{n} \quad$ numeric; the number of locations on the gradient to evaluate the response curve. ... additional arguments passed to other methods.

## Details

sppResponse estimates species responses along indicated gradients.
There is currently no "default" method and the only specified method supplied is for objects fitted by prcurve. This method extracts the fitted responses of species along the principal curve and is a useful diagnostic for identifying overly-complex curves.

## Value

A list is returned with components observed and fitted.values containing the observed and fitted values of the species response and gradient respectively. Each is a list with two components, gradient and response, containing the gradient and response values.

## Author(s)

Gavin L. Simpson

## See Also

prcurve for one function that can be used with sppResponse. A plot method is available; see plot.sppResponse for details.

## Examples

```
## Load the Abernethy Forest data set
data(abernethy)
## Remove the Depth and Age variables
abernethy2 <- abernethy[, -(37:38)]
## Fit the principal curve using varying complexity of smoothers
## for each species
aber.pc <- prcurve(abernethy2, method = "ca", trace = TRUE,
    vary = TRUE, penalty = 1.4)
## Extract the fitted species response curves
resp <- sppResponse(aber.pc)
## Look at only the most abundant/frequently occurring taxa
take <- chooseTaxa(abernethy2, max.abun = 25, n.occ = 10, value = FALSE)
layout(matrix(1:12, ncol = 3)) # split device into panels
plot(resp, which = take)
layout(1) # reset device
```

stdError Standard error of MAT fitted and predicted values

## Description

Computes the (weighted) standard deviation of the environment for the $k$-closest analogues for each sample. This was proposed as one measure of reconstruction uncertainty for MAT models (ter Braak, 1995).

## Usage

```
stdError(object, ...)
## S3 method for class 'mat'
stdError(object, k, weighted = FALSE, ...)
## S3 method for class 'predict.mat'
stdError(object, k, weighted = FALSE, ...)
```


## Arguments

object Object for which the uncertainty measure is to be computed. Currently methods for mat and predict.mat.
$\mathrm{k} \quad$ numeric; how many analogues to take? If missing, the default, k is chosen using getK.
weighted logical; use a weighted computation?
... Additional arguments passed to other methods. Currently not used.
stdError

## Details

Two types of standard error can be produced depending upon whether the mean or weighted mean of $y$ for the $k$ closest analogues is used for the MAT predictions. If weighted = FALSE then the usual standard deviation of the response for the $k$ closest analogues is returned, whereas for weighted $=$ TRUE a weighted standard deviation is used. The weights are the inverse of the dissimilarity between the target observation and each of the $k$ closest analogues.

## Value

A named numeric vector of weighted standard deviations of the environment for the $k$ closest analogues used to compute the MAT predicted values.
The returned vector has attributes " $k$ " and "auto", indicating the number of analogues used and whether this was determined from object or supplied by the user.

## Author(s)

Gavin L. Simpson

## References

Simpson, G.L. (2012) Analogue methods in palaeolimnology. In Birks, H.J.B, Lotter, A.F. Juggins S., and Smol, J.P. (Eds) Tracking Environmental Change Using Lake Sediments, Volume 5: Data Handling and Numerical Techniques. Springer, Dordrecht.
ter Braak, C.J.F. (1995) Non-linear methods for multivariate statistical calibration and their use in palaeoecology: a comparison of inverse ( $k$-nearest neighbours, partial least squares, and weighted averaging partial least squares) and classical approaches. Chemometrics and Intelligent Laboratory Systems 28:165-180.

## See Also

minDC, mat, predict.mat.

## Examples

```
## Imbrie and Kipp Sea Surface Temperature
data(ImbrieKipp)
data(SumSST)
data(V12.122)
## merge training set and core samples
dat <- join(ImbrieKipp, V12.122, verbose = TRUE)
## extract the merged data sets and convert to proportions
ImbrieKipp <- dat[[1]] / 100
ImbrieKippCore <- dat[[2]] / 100
## fit the MAT model using the squared chord distance measure
ik.mat <- mat(ImbrieKipp, SumSST, method = "SQchord")
## standard errors - unweighted
```

```
stdError(ik.mat)
## standard errors - weighted version for above
stdError(ik.mat, k = getK(ik.mat), weighted = TRUE)
## standard errors - weighted; note this uses more (7) analogues
## than the above as this model had lowest LOO error
stdError(ik.mat, weighted = TRUE)
## reconstruct for the V12-122 core data
coreV12.mat <- predict(ik.mat, V12.122, k = 3)
## standard errors
stdError(coreV12.mat)
```

Stratiplot Palaeoecological stratigraphic diagrams

## Description

Draws palaeoecological stratigraphic diagrams of one or more variables as a function of depth/age, with the time dimension flowing from the bottom to the top of the $y$-axis, using the Lattice graphics package.

## Usage

```
Stratiplot(x, ...)
## Default S3 method:
Stratiplot(x, y, type = "l", ylab = NULL, xlab = "",
        pages = 1, rev = TRUE, ylim, sort = c("none", "wa", "var"),
        svar = NULL, rev.sort = FALSE, strip = FALSE, topPad =6,
        varTypes = "relative", absoluteSize = 0.5,
        zoneNames = NULL, drawLegend = TRUE, na.action = "na.omit",
        labelValues = NULL, labelAt = NULL, labelRot = 60, yticks,
        ...)
## S3 method for class 'formula'
Stratiplot(formula, data, subset,
    na.action = "na.pass", type = "l",
    ylab = NULL, xlab = "", pages = 1, ...)
```


## Arguments

x
$y$ numeric vector of depths/ages corresponding to rows in $x$. Length of $y$ must be the same as $\operatorname{nrow}(x)$ or exactly equal to $\operatorname{nrow}(x) / n \operatorname{col}(x)$. See Details.
formula an object of class "formula" (or one that can be coerced to that class): a symbolic description of the model to be fitted. The details of plot specification are given under 'Details'.

| type | character; The type of plotting. Can be a vector. Note that not all Lattice 'type's are supported and some new types are allowed. See panel.Stratiplot for further details. |
| :---: | :---: |
| data | an optional data frame, list or environment (or object coercible by as. data.frame to a data frame) containing the variables to plot. If not found in data, the variables are taken from environment(formula), typically the environment from which Stratiplot is called. |
| subset | an optional vector specifying a subset of observations to be used in the fitting process. |
| na.action | a function which indicates what should happen when the data contain NAs. The default is "na.omit" for the default method, which strips NAs from the stacked data, whereas the default for the formula method is "na. pass" which results in NA being passed on to the plotting function. See Details for further information. |
| ylab, xlab | the x - and y -axis labels |
| pages | numeric; the number of pages to draw the plot over. May be useful for data sets with many species. |
| rev | logical; should the y-axis limits be reversed |
| ylim | user supplied limits for the $y$-axis (time/depth). If not supplied, suitable limits will be determined from the data. As such, in general use ylim need not be supplied. If you choose to supply your own ylim values, note the default for argument rev; the default will reverse the values you supply to ylim. |
| sort | character; how should the variables (columns) of $x$ be sorted on the plot. "wa" sorts by weighted averages of variable svar if not NULL or of $y$ otherwise. The default when "wa" is specified is to order by wiehgted average of the depth/time axis - y. If "var", then ordering is done as per the order of svar. |
| svar | vector; optional variable to sort columns of x by. |
| rev.sort | logical; should the sorting order be reversed. |
| strip | logical; Should panels have strips containing variable labels drawn on them? Default is FALSE, which labels each panel with a label resulting in a more familiar plot style. |
| topPad | numeric; additional padding for the top axis to accomodate long variable names. This is a temporary fudge until the actual space required can be automagically calculated from the variable names themselves. The currently gets most of the way there, but topPad is used to add some extra space if required. |
| varTypes | a character vector of length 1 or equal in length to the number of variables plotted. If length 1 , the vector is expanded to the required length. Two values are allowed; i. "relative", and ii. "absolute". "relative" treats the indicated variable as a percentage type variable and the panel width is scaled relative to the percentage type variables plotted. "absolute" treats the indicated variable as an absolute variable whose panel width should be independent of the other panels. Use "absolute" for variables that are not species compositional data or other percentage type data. |
| absoluteSize | numeric, length 1. This controls the width of panels for variables marked as "absolute", and is the proportion of the largest non-"absolute" panel. |


| zoneNames | character vector of labels, one per zone, with which to label the zone legend, if <br> drawn (see argument drawLegend). See Details. |
| :--- | :--- |
| drawLegend | logical; should a legend for the zones <br> a vector of labels for the variables plotted. Should be equal in length to the <br> number or variables in the resulting plot. The main use for labelValues is to <br> provide non-standard labels for the variables, including a vector of expressions. <br> See Examples for an illustration of this. |
| labelAt, labelRot |  |
| these control the placement and rotation, respectively, of the variable labels. |  |
| labelAt is the coordinate at which the label is drawn; currently only one value |  |
| is used so you can't place labels in different locations depending on which panel |  |
| is drawn. This will be fixed in a future version. The default location for the label |  |
| is the panel mid-point. labelAt controls the rotation of the label; it is a numeric |  |
| value in degree. |  |

## Details

The function now includes preliminary code to handle both relative (proportional or percentage data) and absolute data types, and mixtures thereof. Mixtures can be specified by supplying a vector of types to varTypes, in the same order as the variables are drawn on the plot.
Plots can be specified symbolically using a formula. A typical model has the form $Y \sim$ variables, where $Y$ is either the core depths or sample ages/dates (to be plotted on the $y$-axis) and variables is a series of terms which specifies the variables to plot against $Y$. Terms should be specified with the form var1 + var2 + var3 to plot only those variables. Other, standard, notation for formulae apply, such as model formulae used in lm.

For the formula method the default for argument na.action is "na.pass", which results in any NA values being passed on to the plotting code. This allows for plotting of proxies that been measured on different levels of the stratigraphy. Should you wish to have NA removed from the data before plotting, use na.action = "na.omit", though do note this will remove all rows where any column/variable takes the value NA. The default Stratiplot method, which is used by the formula method for plotting, will strip any NA values from the data provided to it. This allows the function to correctly handle the situation where proxies are measured on different levels of the core and you are displaying the data using lines of polygons. If the NA were not dropped by Stratiplot.default, any NA values would show up as breaks in the line or polygon drawn for each panel.

In summary, the two methods have different defaults for na. action to allow them to handle proxies measured on different levels of the same core. This does mean that you can not use the formula interface and strip NA's at the Stratiplot. default level. If you need that level of control use Stratiplot. default directly by not providing a formula as argument x and by supplying data for the $y$-axis via argument $y$. See Examples for an illustration of these features.

Note that formula is not passed on to xyplot. Instead, the formula is parsed and evaluated within Stratiplot and an appropriate data structure formed to facilitate plotting via xyplot. As such, the special features of Lattice formulae cannot be used.
If zones are drawn on the stratigraphic plot, the zoneNames argument can be used to supply a set of names with which to label the zones using a legend. This legend is drawn on the right-hand side of the the straigraphic diagram if drawLegend = TRUE is supplied. The zoneNames must be supplied in stratigraphic order, as that is the order in which they are drawn in the legend. Whether this ordering is reversed or not will depend on the value of argument rev. It is up to the user to provide the labels in the correct order. Zones are specified by the zone boundaries (excluding the core sequence top and bottom), and as a result 1 more label is required than the number of zone boundaries supplied. If no zoneNames is not supplied, but a legend is requested, suitable names will be produced. If you do not wish to have any labelling at all, then set zoneNames $=" "$ as this will get recycled to the correct length. See the Example section for an illustration of how this drawing zones works.

## Value

A side effect of calling Stratiplot is that a plot is drawn on the currently active device. A Lattice plot object of class "trellis" is returned invisibly. This is a change from pre $0.17-0$ version of the package.

## Note

The function currently doesn't know about ages/dates and will interpret these as 'depths' instead. This will be fixed in a future version.

## Author(s)

Gavin L. Simpson.

## See Also

xyplot, panel.Stratiplot, panel.Loess.

## Examples

```
data(V12.122)
Depths <- as.numeric(rownames(V12.122))
(plt <- Stratiplot(Depths ~ O.univ + G.ruber + G.tenel + G.pacR,
    data = V12.122, type = c("h","l","g","smooth")))
## Order taxa by WA in depth --- ephasises change over time
(plt <- Stratiplot(Depths ~ O.univ + G.ruber + G.tenel + G.pacR,
    data = V12.122, type = c("h"), sort = "wa"))
## Using the default interface
spp.want <- c("O.univ","G.ruber","G.tenel","G.pacR")
(plt <- Stratiplot(V12.122[, spp.want], y = Depths,
    type = c("poly", "g")))
## Adding zones to a Stratigraphic plot
```

```
## Default labelling and draw zone legend
## Here we choose 4 arbitrary Depths as the zone boundaries
set.seed(123)
Zones <-sample(Depths, 4)
Stratiplot(Depths ~ O.univ + G.ruber + G.tenel + G.pacR,
    data = V12.122, type = c("poly","g"),
    zones = Zones)
## As before, but supplying your own zone labels
zone.labs <- c("A","B", "C", "D","E")
Stratiplot(Depths ~ O.univ + G.ruber + G.tenel + G.pacR,
    data = V12.122, type = c("poly","g"),
    zones = Zones, zoneNames = zone.labs)
## Suppress the drawing of the zone legend
Stratiplot(Depths ~ O.univ + G.ruber + G.tenel + G.pacR,
    data = V12.122, type = c("poly","g"),
    zones = Zones, drawLegend = FALSE)
## Add zones and draw a legend, but do not label the zones
Stratiplot(Depths ~ O.univ + G.ruber + G.tenel + G.pacR,
    data = V12.122, type = c("poly","g"),
    zones = Zones, zoneNames = "")
## Show illustration of NA handling
set.seed(42)
dat <- data.frame(Depth = 1:20, LOI = runif(20), TC = NA)
dat <- within(dat, TC[sample(20, 10)] <- runif(10))
## default is 'na.action = "na.pass"'
Stratiplot(Depth ~ LOI + TC, data = dat, type = c("l","p"))
## to remove rows with NA, use 'na.action = "na.omit"'
Stratiplot(Depth ~ LOI + TC, data = dat, type = c("l","p"),
    na.action = "na.omit")
## Example of two proxies measured on different levels of core
## (Here measurements on alternate levels)
set.seed(5)
dat2a <- data.frame(Depth = seq(1, by = 2, length = 20), LOI = runif(20))
dat2b <- data.frame(Depth = seq(0, by = 2, length = 20), TC = runif(20))
dat2 <- join(dat2a, dat2b, na.replace = FALSE, split = FALSE)
dat2 <- dat2[order(dat2$Depth), ]
head(dat2)
## Default is to allow NA through formula, but drop them when plotting
Stratiplot(Depth ~ LOI + TC, data = dat2, type = c("l","p"))
## compare with this if we didn't suppress NA in default Stratiplot
## method (can't use formula interface for this yet
Stratiplot(dat2[,-1], dat2[,1], type = c("l","p"),
    na.action = "na.pass")
## Notice no lines are draw as there a no "sections" ithout missing
## levels. If you want/desire this behaviour then you can't use formula
## interface yet as there is no way to specify the na.action separately
```

```
## Works with matrices
M <- as.matrix(V12.122)
Stratiplot(M, Depths, type = c("h"))
## Custom variable labels using expressions
df <- data.frame(Age = 1:10, Var1 = rnorm(10), Var2 = rnorm(10),
    Var3 = rnorm(10))
## Use a vector of expressions to label variables on plot
## See ?plotmath for syntax of expressions
exprs <- expression(delta^{15}*N, # label for Var1
    delta^{18}*0, # label for Var2
    delta^{13}*C) # label for Var3
Stratiplot(Age ~ ., data = df, labelValues = exprs, varTypes = "absolute")
```

summary. analog Summarise analogue matching results

## Description

summary method for class "analog".

## Usage

\#\# S3 method for class 'analog'
summary(object, display = c("dist", "names", "quantiles"),

$$
\mathrm{k}=10, \text { probs }=\mathrm{c}(0.01,0.02,0.05,0.1,0.2), \ldots)
$$

## Arguments

object an object of class "analog", usually as a result of a call to analog.
display character; one or more of the listed choices. Determines which aspects of the analog results are summarised.
k number of analogues to use. If missing, $k$ is chosen automatically as the $k$ that achieves lowest RMSE.
probs numeric; giving the probabilities of the distribution to return quantiles for. See quantile.
... arguments passed to or from other methods.

## Value

A list with one or more of the components listed below. Attributes "method", "train", "call" and " $k$ " contain the dissimilarity coefficient used, whether the training set dissimilarities were saved, the matched function call and the number of close analogues to return respectively.
dists a matrix of dissimilarities between training set samples and fossil samples. The number of rows is given by argument $k$. There is a column for each fossil sample.

# names a matrix of names of samples from the training set that are analogues for each fossil sample. The number of rows is given by argument $k$. There is a column for each fossil sample. <br> quantiles numeric; the quantiles of the distribution of the pairwise dissimilarities for the training set for probabilities prob. 

## Author(s)

Gavin L. Simpson

## See Also

```
analog.
```


## Examples

```
## Not run:
## continue the RLGH example from ?join
example(join)
## analog matching between SWAP and RLGH core
swap.analog <- analog(swapdiat, rlgh, method = "chord")
swap.analog
summary(swap.analog)
## End(Not run)
```


## summary.bootstrap.mat Summarise bootstrap resampling for MAT models

## Description

summary method for class "bootstrap.mat".

## Usage

\#\# S3 method for class 'bootstrap.mat'
summary (object, ...)

## Arguments

object an object of class "bootstrap.mat", usually the result of a call to bootstrap.mat.
... arguments passed to or from other methods.

## Value

A data frame with the following components:

| observed | vector of observed environmental values. |
| :--- | :--- |
| model | a list containing the apparent or non-bootstrapped estimates for the training set. |
| With the following components: |  |
| estimated: estimated values for the response |  |
| residuals: model residuals |  |
| r.squared: Apparent $R^{2}$ between observed and estimated values of y |  |
| avg.bias: Average bias of the model residuals |  |
| max.bias: Maximum bias of the model residuals |  |
| rmse: Apparent error (RMSE) for the model |  |
| k: numeric; indicating the size of model used in estimates and predictions |  |
| a list containing the bootstrap estimates for the training set. With the following |  |
| components: |  |
| estimated: Bootstrap estimates for the response |  |
| residuals: Bootstrap residuals for the response |  |
| r.squared: Bootstrap derived $R^{2}$ between observed and estimated values of |  |
| the response |  |
| avg.bias: Average bias of the bootstrap derived model residuals |  |
| max.bias: Maximum bias of the bootstrap derived model residuals |  |
| rmsep: Bootstrap derived RMSEP for the model |  |

bootstrap: a list containing the bootstrap estimates for the new samples, with some or all of the same components as bootstrap, above
sample.errors: a list containing the bootstrap-derived sample specific errors for the new samples, with some or all of the same components as sample.errors, above

## Author(s)

Gavin L. Simpson

## See Also

bootstrap.mat, mat, summary.

## Examples

```
## Not run:
## continue the RLGH example from ?join
example(join)
## fit the MAT model using the squared chord distance measure
swap.mat <- mat(swapdiat, swappH, method = "SQchord")
## bootstrap training set
swap.boot <- bootstrap(swap.mat, k = 10, n.boot = 100)
swap.boot
summary(swap.boot)
## End(Not run)
```

summary.cma

Summarise the extraction of close modern analogues

## Description

summary method for class "cma".

## Usage

\#\# S3 method for class 'cma'
summary (object, ...)

## Arguments

object an object of class "cma", usually the result of a call to cma.
... arguments passed to or from other methods.

## Value

An object of class "summary.cma" with the components of an object of class cma, plus:
distances a matrix of distances/dissimilarities. Individual columns contain the ordered close modern analogues for individual fossil samples. Rows of this matrix refer to the $k^{\text {th }}$ closest analogue for each fossil sample. See notes below.
samples a matrix of sample names from the reference set that are close modern analogues for a fossil sample. Individual columns contain the ordered close modern analogues for individual fossil samples. Rows of this matrix refer to the $k^{\text {th }}$ closest analogue for each fossil sample. See notes below.

## Note

Currently, only objects of class analog are supported. The number of rows in the returned matrices is equal to the maximum number of close modern analogues identified for an individual fossil sample. If no close modern analogues exist for an individual fossil sample are identified, then the relevant column in "distances" will contain all missing values and in "samples" the string "none". Rows of individual columns will be padded with missing values if the number of close modern analogues for that sample is less than the maximum number of close modern analogues identified for a single sample.

## Author(s)

Gavin L. Simpson

## See Also

cma

## Examples

```
## Not run:
## continue the RLGH example from ?join
example(join)
## analog matching between SWAP and RLGH core
swap.analog <- analog(swapdiat, rlgh, method = "chord")
swap.analog
summary(swap.analog)
## close modern analogues
swap.cma <- cma(swap.analog, cutoff = 0.6)
swap.cma
summary(swap.cma)
## End(Not run)
```


## Description

summary method for class "mat".

## Usage

\#\# S3 method for class 'mat'
summary (object, $k=10$, digits $=\max (2$, getOption("digits") - 4), ...)

## Arguments

object an object of class "cma", usually the result of a call to cma.
$\mathrm{k} \quad$ numeric; maximum modern analogues to use to summarise model fits.
digits numeric; the number of significant digits with which to format results.
... arguments passed to or from other methods.

## Value

A list with the components below. The number of analogues used, $k$ is returned as attribute " k ".
summ a data.frame containing the model fits for training set samples. See notes below.
tbl matrix of summary statistics for an un-weighted model.
tbl.W matrix of summary statistics for a weighted model.
call the matched function call
quantiles the quantiles of the distribution of pairwise dissimilarities for the training set, for "probs $=c(0.01,0.02,0.05,0.1,0.2)$ ".

## Note

The returned component "summ" contains the following:
Obs: the observed responses for the training set samples.
Est: the fitted values of the response for training set samples based on the average of $k$-closest analogues.
Resi: the residuals of the fitted model based on the average of $k$-closest analogues.
W.Est: the fitted values of the response for training set samples based on the weighted average of $k$-closest analogues.
W.Resi: the residuals of the fitted model based on the weighted average of $k$-closest analogues.
minDC: dissimilarity of closest analogue in training set for each training set sample.
minResi: smallest residual for an un-weighted model of size " $k$ ".
k: size of model leading to minimal residual, "minResi".
minW.Resi: smallest residual for a weighted model of size " $k$. W".
$\mathbf{k} . W:$ size of model leading to minimal residual, "minW. Resi".

## Author(s)

Gavin L. Simpson

## See Also

mat, summary.

## Examples

```
## Not run:
## continue the RLGH example from ?join
example(join)
## fit the MAT model using the squared chord distance measure
swap.mat <- mat(swapdiat, swappH, method = "SQchord")
swap.mat
## model summary
summary(swap.mat)
## model summary - evaluating models using k = 1, ..., 20
## analogues instead of the default, 10.
summary(swap.mat, k = 20)
## End(Not run)
```

summary.predict.mat Summarise MAT model predictions

## Description

summary method for objects of class "predict.mat".

## Usage

\#\# S3 method for class 'predict.mat'
summary (object, ...)

## Arguments

object an object of class "predict.mat", usually the result of a call to predict.mat.
... arguments passed to or from other methods.

## Value

An object of class "summary.predict.mat", see predict.mat for more details.

## Author(s)

Gavin L. Simpson

## See Also

predict.mat, mat, bootstrap.mat and summary.

## Examples

```
## Not run:
## continue the RLGH example from ?join
example(join)
## fit the MAT model using the squared chord distance measure
swap.mat <- mat(swapdiat, swappH, method = "SQchord")
## predict for RLGH data
swap.pred <- predict(swap.mat, rlgh, bootstrap = FALSE)
summary(swap.pred)
## End(Not run)
```

swapdiat

SWAP sub-fossil diatom and pH training set

## Description

The Surface Waters Acidifcation Project (SWAP) Palaeolimnology Programme diatom surface sample training set.

## Usage

data(swapdiat)

## Format

A data frame of observations on 277 diatom taxa for the 167-lake SWAP diatom-pH training set.

## Details

This data set contains the original 167-lake SWAP diatom-pH training set.
The variable names (colnames) are DIATCODE codes for individual taxa.

## References

Stevenson, A.C., Juggins, S., Birks, H.J.B., Anderson, D.S., Anderson, N.J., Battarbee, R.W., Berge, F., Davis, R.B., Flower, R.J., Haworth, E.Y., Jones, V.J., Kingston, J.C., Kreiser, A.M., Line, J.M., Munro, M.A.R., and Renberg, I. (1995). The Surface Waters Acidification Project Palaeolimnology programme: modern diatom/lake-water chemistry data-set. ENSIS Publishing, London.

## See Also

swappH

## Examples

> data(swapdiat)
swappH
SWAP sub-fossil diatom and $p H$ training set

## Description

The Surface Waters Acidifcation Project (SWAP) Palaeolimnology Programme diatom-pH surface sample training set.

## Usage

data(swappH)

## Format

Numeric vector containing the pH of the 167 lakes from the SWAP diatom- pH training set. The values are the average of 4 quarterly samples.

## References

Stevenson, A.C., Juggins, S., Birks, H.J.B., Anderson, D.S., Anderson, N.J., Battarbee, R.W., Berge, F., Davis, R.B., Flower, R.J., Haworth, E.Y., Jones, V.J., Kingston, J.C., Kreiser, A.M., Line, J.M., Munro, M.A.R., and Renberg, I. (1995). The Surface Waters Acidification Project Palaeolimnology programme: modern diatom/lake-water chemistry data-set. ENSIS Publishing, London.

## See Also

swapdiat

## Examples

```
data(swappH)
str(swappH)
```


## Description

Project passive (e.g. sediment core) samples into an ordination of a set of training samples.

## Usage

```
timetrack(X, passive, env, method = c("cca", "rda"),
            transform = "none", formula, scaling = 3,
            rank = "full", join = "left", correlation = FALSE,
            hill = FALSE, ...)
## S3 method for class 'timetrack'
fitted(object, which = c("passive", "ordination"),
            model = NULL, choices = 1:2, ...)
## S3 method for class 'timetrack'
predict(object, newdata, ...)
## S3 method for class 'timetrack'
scores(x, which = c("ordination", "passive"),
        scaling = x$scaling, choices = 1:2, display = "sites", ...)
## S3 method for class 'timetrack'
plot(x, choices = 1:2, display = c("wa", "lc"),
        order, type = c("p", "n"), ptype = c("l", "p", "o", "b", "n"),
        pch = c(1,2), col = c("black","red"), lty = "solid", lwd = 1,
        xlim = NULL, ylim = NULL, ...)
## S3 method for class 'timetrack'
points(x, choices = 1:2, which = c("passive", "ordination"),
        display = c("wa","lc"), order, ...)
```


## Arguments

$$
\begin{array}{ll}
\text { X } & \text { matrix-like object containing the training set or reference samples. } \\
\text { passive } & \begin{array}{l}
\text { matrix-like object containing the samples to be projected into the ordination of } \\
\text { X. Usually a set of sediment core samples. }
\end{array} \\
\text { env } & \begin{array}{l}
\text { optional data frame of environmental or constraining variables. If provided, } \\
\text { a constrained ordination of } X \text { is performed. If formula is supplied variables } \\
\text { named in formula are looked up with env. }
\end{array} \\
\text { method } & \begin{array}{l}
\text { character, resolving to an ordination function available in vegan. Currently only } \\
\text { "cca", the default, and "rda" are supported. }
\end{array}
\end{array}
$$

| transform | character; the name of the transformation to apply to both $X$ and passive. The |
| :--- | :--- |
| transformations are performed using tran and valid options are given by that |  |
| function's method argument. |  |
| a one-sided model formula; if provided, it defines the right hand side of the |  |
| model formula for the ordination function and is supplied as argument formula |  |
| to the ordination function. E.g.~formula = ~ var1 + var2. If supplied then env |  |
| must also be supplied |  |
| numeric or character; the ordination scaling to apply. Useful options are likely |  |
| to be 1 or 3 where the focus is on the samples. For character, see options in |  |
| scores. cca: character version of the useful scalings are "sites" and "symmetric". |  |
| scaling | See arguments correlation and hill. |
| correlation, hill |  |
| logical; additional arguments passed to predict. cca and predict. rda. See |  |

\(\left.\begin{array}{ll}l \mathrm{lty}, \mathrm{lwd} \& graphical parameters for the plotted time track for ptype!=" \mathrm{p} " . <br>

\mathrm{xlim}, \mathrm{ylim} \& user specified axis limits for the plot.\end{array}\right]\)| arguments passed to other methods. timetrack passes arguments on to tran |
| :--- |
| and the ordination function given in method. fitted passes arguments on to |
| other fitted methods as appropriate. plot passes arguments on to the underly- |
| ing plotting functions. predict passes arguments on to tran for use in applyign |
| the transformation. |

## Details

The timetrack is a way to visualise changes in species composition from sediment core samples within an underlying reference ordination or, usually, training set samples. This technique has been most often applied in situations where the underlying ordination is a constrained ordination and thence the timetrack of sediment core samples within the ordination reflects both the change in species composition and the indicative changes in the constraining variables.
The sediment core samples are projected passively into the underlying ordination. By projected passively, the locations of the core samples are predicted on the basis of the ordination species scores. A common set of species (columns) is required to passively place the sediment samples into the ordination. To achieve this, the left outer join of the species compositions of the training set and passive set is determined; the left outer join results in the passive data matrix having the same set of species (variables; columns) as the training set. Any training set species not in the passive set are added to the passive set with abundance 0 . Any passive species not in the training set are removed from the passive set.

## Value

The plot method results in a plot on the currently active device, whilst the fitted and scores methods return the matrix of fitted locations on the set of ordination axes.
timetrack returns an object of class "timetrack", a list with the following components:
ordination the ordination object, the result of the call to the function of the name method.
fitted.values the matrix of fitted locations for the passive samples on the ordination axes.
method the ordination function used.
formula if supplied, the model formula used to define the ordination model.
scaling the ordination scaling applied.
rank The rank or the number of axes used in the approximation. The default is to use all axes (full rank) of the "model".
model Show constrained ("CCA") or unconstrained ("CA") results.
labels a list of names for the $X$, passive, and env arguments.
call The matched function call.
$X \quad$ The training data.
transform The transformation applied, if any.

## Author(s)

Gavin L. Simpson

## See Also

cca and rda for the underlying ordination functions.

## Examples

```
## load the RLGH and SWAP data sets
data(rlgh, swapdiat)
## Fit the timetrack ordination
mod <- timetrack(swapdiat, rlgh, transform = "hellinger",
        method = "rda")
mod
## Plot the timetrack
plot(mod, ptype = "b", col = c("forestgreen", "orange"), lwd = 2)
## Other options (reorder the time track)
ord <- rev(seq_len(nrow(rlgh)))
plot(mod, choices = 2:3, order = ord, ptype = "b",
    col = c("forestgreen", "orange"), lwd = 2)
## illustrating use of the formula
data(swappH)
mod2 <- timetrack(swapdiat, rlgh, env = data.frame(pH = swappH),
                        transform = "hellinger", method = "rda",
        formula = ~ pH)
mod2
plot(mod2)
## scores and fitted methods
## IGNORE_RDIFF_BEGIN
head(fitted(mod, type = "passive"))
head(scores(mod, type = "passive"))
## IGNORE_RDIFF_END
## predict locations in timetrack for new observations
take <- rlgh[1:50, ]
take <- take[ , colSums(take) > 0]
mod3 <- predict(mod, newdata = take)
class(mod3) ## returns a timetrack object
take <- rlgh[-(1:50), ]
take <- take[ , colSums(take) > 0]
mod4 <- predict(mod, newdata = take)
## build a plot up from base parts
plot(mod, type = "n", ptype = "n")
points(mod, which = "ordination", col = "grey", pch = 19, cex = 0.7)
points(mod3, which = "passive", col = "red")
points(mod4, which = "passive", col = "blue")
## Fit the timetrack ordination - passing scaling args
mod <- timetrack(swapdiat, rlgh, transform = "hellinger",
```

```
    method = "rda", scaling = "sites",
        correlation = TRUE)
    mod
    plot(mod)
```

tortula Morphological data for ten taxa of the genus Tortula

## Description

These data are observations on a series of seven morphological variables for individuals in of the Tortula sect. Rurales De Not. (Pottiaceae, Musci.

## Usage

data(tortula)

## Format

tortula is a data frame of seven morphological measurements on 14 individuals from the genus Tortula.

Taxon factor; the species of Tortula
Hydroid logical; presence of hydroid cells
LeafOutline ordered; shape of the leaf outline
Denticulation ordered; degree of denticulation
ApexShape ordered; shape of the leaf apex
Length numeric, leaf length
Diameter numeric; leaf diameter
Papillae numeric; number of papillae per cell
The last three variables are the average of ten replicate samples from the same herbarium capsule.

## Source

The data were presented in Podani (1999).

## References

Podani, J. (1999) Extending Gower's coefficient of similarity to ordinal characters. Taxon 48, 331340.

## Examples

data(tortula)
head(tortula)
str(tortula)

## Description

Provides common data transformations and standardizations useful for palaeoecological data. The function acts as a wrapper to function decostand in package vegan for several of the available options.

The formula method allows a convenient method for selecting or excluding subsets of variables before applying the chosen transformation.

## Usage

\#\# Default S3 method:
$\operatorname{tran}(x$, method, $a=1, b=0, p=2$, base $=\exp (1)$, na. $\mathrm{rm}=$ FALSE, na.value $=0, \ldots$ )
\#\# S3 method for class 'formula'
tran(formula, data $=$ NULL, subset $=$ NULL, na.action = na.pass, ...)

## Arguments

| x |  |
| :--- | :--- |
| method | A matrix-like object. <br> transformation or standardization method to apply. See Details for available <br> options. |
| a | Constant to multiply $x$ by. method $=$ "log" only. Can be a vector, in which case <br> the vector of values to multiply each column of $x$ by. |
| b | Constant to add to $x$ before taking logs. method $=$ "log" only. Can be a vector, <br> in which case the vector of values to add to each column of $x$. |
| p | The power to use in the power transformation. <br> the base with respect to which logarithms are computed. See log for further <br> details. The default is to compute natural logarithms. |
| na.rm | Should missing values be removed before some computations? |
| na.value | The value with which to replace missing values (NA). |
| $\ldots$ | Arguments passed to decostand, or other tran methods. |
| formula model formula describing the variables to be transformed. The formula should |  |

See model.frame for details on these arguments. data will generally be the object or environment within which the variables in the forumla are searched for.

## Details

The function offers following transformation and standardization methods for community data:

- sqrt: take the square roots of the observed values.
- cubert: take the cube root of the observed values.
- rootroot: take the fourth root of the observed values. This is also known as the root root transformation (Field et al 1982).
- log: take the logarithms of the observed values. The tansformation applied can be modified by constants a and b and the base of the logarithms. The transformation applied is $x^{*}=$ $\log _{\text {base }}(a x+b)$
- $\log 1 \mathrm{p}$ : computes $\log (1+x)$ accurately also for $|x| \ll 1$ via $\log 1 \mathrm{p}$. Note the arguments a and b have no effect in this method.
- expm1: computes $\exp (x)-1)$ accurately for $|x| \ll 1$ via expm1.
- reciprocal: returns the multiplicative inverse or reciprocal, $1 / x$, of the observed values.
- freq: divide by column (variable, species) maximum and multiply by the number of non-zero items, so that the average of non-zero entries is 1 (Oksanen 1983).
- center: centre all variables to zero mean.
- range: standardize values into range $0 \ldots$. If all values are constant, they will be transformed to 0 .
- percent: convert observed count values to percentages.
- proportion: convert observed count values to proportions.
- standardize: scale $x$ to zero mean and unit variance.
- pa: scale $x$ to presence/absence scale $(0 / 1)$.
- missing: replace missing values with na.value.
- chi. square: divide by row sums and square root of column sums, and adjust for square root of matrix total (Legendre \& Gallagher 2001). When used with the Euclidean distance, the distances should be similar to the the Chi-square distance used in correspondence analysis. However, the results from cmdscale would still differ, since CA is a weighted ordination method.
- hellinger: square root of observed values that have first been divided by row (site) sums (Legendre \& Gallagher 2001).
- wisconsin: applies the Wisconsin double standardization, where columns (species, variables) are first standardized by maxima and then sites (rows) by site totals.
- pcent2prop: convert percentages to proportions.
- prop2pcent: convert proportions to percentages.
- logRatio: applies a log ransformation (see log above) to the data, then centres the data by rows (by subtraction of the mean for row $i$ from the observations in row $i$ ). Using this transformation subsequent to PCA results in Aitchison's Log Ratio Analysis (LRA), a means of dealing with closed compositional data such as common in palaeoecology (Aitchison, 1983).
- power: applies a power tranformation.
- rowCentre, rowCenter: Centres $x$ by rows through the subtraction of the corresponding row mean from the observations in the row.
- colCentre colCenter: Centres x by columns through the subtraction of the corresponding column mean from the observations in the row.
- none none: no transformation is applied.


## Value

Returns the suitably transformed or standardized $x$. If $x$ is a data frame, the returned value is likewise a data frame. The returned object also has an attribute "tran" giving the name of applied transformation or standardization "method".

## Author(s)

Gavin L. Simpson. Much of the functionality of tran is provided by decostand, written by Jari Oksanen.

## References

Aitchison, J. (1983) Principal components analysis of compositional data. Biometrika 70(1); 57-65.
Field, J.G., Clarke, K.R., \& Warwick, R.M. (1982) A practical strategy for analysing multispecies distributions patterns. Marine Ecology Progress Series 8; 37-52.

Legendre, P. \& Gallagher, E.D. (2001) Ecologically meaningful transformations for ordination of species data. Oecologia 129; 271-280.

Oksanen, J. (1983) Ordination of boreal heath-like vegetation with principal component analysis, correspondence analysis and multidimensional scaling. Vegetatio 52; 181-189.

## See Also

decostand

## Examples

```
data(swapdiat)
## convert percentages to proportions
sptrans <- tran(swapdiat, "pcent2prop")
## apply Hellinger transformation
spHell <- tran(swapdiat, "hellinger")
## Dummy data to illustrate formula method
d <- data.frame(A = runif(10), B = runif(10), C = runif(10))
## simulate some missings
d[sample(10,3), 1] <- NA
## apply tran using formula
tran(~ . - B, data = d, na.action = na.pass,
    method = "missing", na.value = 0)
```


## varExpl Variance explained by ordination axes

## Description

Extracts information about the variance explained by ordination axes and expresses it in a variety of ways.

## Usage

```
varExpl(object, ...)
## S3 method for class 'cca'
varExpl(object, axes = 1L, cumulative = FALSE,
    pcent = FALSE, ...)
## S3 method for class 'prcurve'
varExpl(object, pcent = FALSE, ...)
```


## Arguments

object an R object of an appropriate type. Currently only for objects that inherit from classes "cca" or "prcurve".
axes numeric vector indicating which axes to compute variance explained for.
cumulative logical; should the variance be explained as a cumulative sum over the axes?
pcent logical; should the variance explained be expressed as a percentage of the total variance.
... additional arguments passed to other methods. Currently not used.

## Value

A numeric vector variance explained by each axis.

## Author(s)

Gavin L. Simpson

## See Also

See cca and prcurve for functions that produce objects that varExpl() can work with.

## Examples

```
data(abernethy)
## Remove the Depth and Age variables
abernethy2 <- abernethy[, -(37:38)]
## Fit PCA
aber.pca <- rda(abernethy2)
## Distance along the first PCA axis
varExpl(aber.pca)
```

    wa Weighted averaging transfer functions
    
## Description

Implements the weighted averaging transfer function methodology. Tolerance down-weighting and inverse and classicial deshrinking are supported.

## Usage

```
wa(x, ...)
    ## Default S3 method:
    wa(x, env,
        deshrink = c("inverse", "classical", "expanded", "none", "monotonic"),
        tol.dw = FALSE, useN2 = TRUE,
        na.tol = c("min","mean","max"),
        small.tol = c("min","mean","fraction","absolute"),
        min.tol = NULL, f = 0.1, ...)
    ## S3 method for class 'formula'
    wa(formula, data, subset, na.action,
    deshrink = c("inverse", "classical", "expanded", "none", "monotonic"),
    tol.dw = FALSE, useN2 = TRUE, na.tol = c("min","mean","max"),
    small.tol = c("min","mean","fraction","absolute"), min.tol = NULL,
    f = 0.1,\ldots, model = FALSE)
    ## S3 method for class 'wa'
    fitted(object, ...)
    ## S3 method for class 'wa'
    residuals(object, ...)
    ## S3 method for class 'wa'
```

```
coef(object, ...)
waFit(x, y, tol.dw, useN2, deshrink, na.tol, small.tol,
    min.tol, f)
```


## Arguments

x
env, y
deshrink
tol.dw
useN2 logical; should Hill's N2 values be used to produce un-biased tolerances?
na. tol character; method to use to replace missing (NA) tolerances in WA computations. Missing values are replaced with the minimum, average or maximum tolerance observed that is not missing.
small.tol character; method to replace small tolerances. See Details.
min.tol numeric; threshold below which tolerances are treated as being 'small'. Default is not to replace small tolerances.
$\mathrm{f} \quad$ numeric, $0<f<1$; fraction of environmental gradient env to replace small tolerances with if small.tol = "fraction" is specified.
formula a model formula
data an optional data frame, list or environment (or object coercible by as. data. frame to a data frame) containing the variables specified on the RHS of the model formula. If not found in data, the variables are taken from environment (formula), typically the environment from which wa is called.
subset an optional vector specifying a subset of observations to be used in the fitting process.
na.action a function which indicates what should happen when the data contain NAs. The default is set by the na.action setting of options, and is na.fail if that is unset. The 'factory-fresh' default is na.omit. Another possible value is NULL, no action. Value na. exclude can be useful.
model logical. If TRUE the model frame is returned.
object an Object of class "wa", the result of a call to wa.
... arguments to other methods.

## Details

A typical model has the form response ~ terms where response is the (numeric) response vector (the variable to be predicted) and terms is a series of terms which specifies a linear predictor for response. A terms specification of the form first + second indicates all the terms in first together with all the terms in second with duplicates removed. A specification of . is shorthand for all terms in data not already included in the model.

Species that have very small tolerances can dominate reconstructed values if tolerance down-weighting is used. In wa, small tolerances are defined as a tolerance that is $<$ min.tol. The default is to not replace small tolerances, and the user needs to specify suitable values of min.tol. Function tolerance may be of use in computing tolerances before fitting the WA model.
Small tolerances can be adjusted in several ways:
min small tolerances are replaced by the smallest observed tolerance that is greater than, or equal to, min.tol. With this method, the replaced values will be no smaller than any other observed tolerance. This is the default in analogue.
mean small tolerances are replaced by the average observed tolerance from the set that are greater than, or equal to, min.tol.
fraction small tolerances are replaced by the fraction, $f$, of the observed environmental gradient in the training set, env.
absolute small tolerances are replaced by min.tol.
Function waFit is the workhorse implementing the actual WA computations. It performs no checks on the input data and returns a simple list containing the optima, tolernances, model tolerances, fitted values, coefficients and the numbers of samples and species. See Value below for details of each component.

## Value

An object of class "wa", a list with the following components:

| wa.optima | The WA optima for each species in the model. |
| :--- | :--- |
| tolerances | The actual tolerances calculated (these are weighted standard deviations). <br> model.tol |
| The tolerances used in the WA model computations. These will be similar to <br> tol, but will no contain any NAs and any small tolerances will have been replaced <br> with the appropriate value. |  |
| fitted.values | The fitted values of the response for each of the training set samples. |
| residuals | Model residuals. |
| coefficients | Deshrinking coefficients. Note that in the case of deshrink = "monotonic" <br> this is a list with components sm (the representation of the smooth term as re- <br> turned by smoothCon) and p (solutions to the least squares fit with monotonic <br> constraints, the result of a call to pcls). |
| rmse | The RMSE of the model. |
| r.squared | The coefficient of determination of the observed and fitted values of the re- <br> sponse. |
| avg.bias, max.bias |  |
| n. samp, n.spp | The average and maximum bias statistics. |
| The number of samples and species in the training set. |  |
| dol.dw | The deshrinking regression method used. <br> logical; was tolerance down-weighting applied? |
| call | The matched function call. |


| orig.x | The training set species data. |
| :--- | :--- |
| orig.env | The response data for the training set. |
| options.tol | A list, containing the values of the arguments useN2, na.tol, small.tol, min.tol, <br> and $f$. |
| terms, model | Model terms and model.frame components. Only returned by the formula <br> method of wa. |

## Author(s)

Gavin L. Simpson and Jari Oksanen

## See Also

mat for an alternative transfer function method.

## Examples

```
data(ImbrieKipp)
data(SumSST)
## fit the WA model
mod <- wa(SumSST ~., data = ImbrieKipp)
mod
## extract the fitted values
fitted(mod)
## residuals for the training set
residuals(mod)
## deshrinking coefficients
coef(mod)
## diagnostics plots
par(mfrow = c(1,2))
plot(mod)
par(mfrow = c(1,1))
## caterpillar plot of optima and tolerances
caterpillarPlot(mod) ## observed tolerances
caterpillarPlot(mod, type = "model") ## with tolerances used in WA model
## plot diagnostics for the WA model
par(mfrow = c(1,2))
plot(mod)
par(mfrow = c(1,1))
## tolerance DW
mod2 <- wa(SumSST ~ ., data = ImbrieKipp, tol.dw = TRUE,
    min.tol = 2, small.tol = "min")
```

```
mod2
## compare actual tolerances to working values
with(mod2, rbind(tolerances, model.tol))
## tolerance DW
mod3 <- wa(SumSST ~ ., data = ImbrieKipp, tol.dw = TRUE,
    min.tol = 2, small.tol = "mean")
mod3
## fit a WA model with monotonic deshrinking
mod4 <- wa(SumSST ~., data = ImbrieKipp, deshrink = "monotonic")
mod4
## extract the fitted values
fitted(mod4)
## residuals for the training set
residuals(mod4)
```

weightedCor Weighted correlation test of WA reconstruction

## Description

Weighted correlation between WA optima from training set and axis 1 scores of constrained ordination fitted to fossil data with WA model predictions for fossil samples as constraints.

## Usage

```
## Default S3 method:
weightedCor(x, env, fossil, method = c("rda", "cca"),
        test = TRUE, type = c("simulate", "permute"), sim = 999,
        verbose = TRUE, ...)
    ## S3 method for class 'weightedCor'
    plot(x,
    type = c("bubble", "null"),
    weighted = TRUE,
    size = 0.25,
    xlab = paste(x$env, "WA Optima"),
    ylab = "Axis 1 Score",
    xlim,
    main = "",
    sub = NULL,
    border = "gray75",
    col = "gray75",
```

```
obscol = "red",
fg = "black", ...)
```


## Arguments

x
training set covariates, a matrix-like object usually of species/proxy data. For the plot method, an object of class "weightedCor", the result of a call to weightedCor.
env training set response, a vector usually of environmental data.
fossil matrix of fossil/core species/proxy data for which a reconstruction is sought.
method constrained ordination method. One of "rda" and "cca". Currently only "rda" is supported.
test logical; should the observed correlation be tested?
type the type of test to apply. One of "simulate" or "permute". The latter is currently not implemented. For the plot method, the type of plot to produce.
sim numeric; number of simulations or permutations to permform as part of the test
verbose logical; should the progress of the test be shown via a progress bar?
$\ldots \quad$ arguments passed to other methods. In the case of the plot method, additional graphical parameters can be supplied.
weighted logical; should the null distribution plotted be of the weighted or normal correlation.
size numeric; the size of the largest bubble in inches. See symbols and argument inches for details.
xlim,xlab,ylab, main, sub graphical parameters with their usual meaning.
border, col The border and fill colours for the histogram bars.
fg The colour of the bubbles drawn on the bubble plot.
obscol The colour of the indicator for the observed correlation.

## Value

The plot method produces a plot on the current device. weightedCor() returns a list with the following components:
wtdCorrel, Correl
numeric; the observed weighted and standard correlation.
data data frame; containing the training set WA Optima, axis 1 species scores, and mean abundance for each species.
ord the fitted constrained ordination.
model the fitted WA model.
method the ordination method used.
ndist the null distribution produced. NULL if argument test was FALSE.
sim numeric; the number of simulations or permutations used to test the observed correlations.
weightedCor
type the type of test performed.
env the deparsed version of env argument. Used for plotting.
call the matched function call.

## Author(s)

Gavin L. Simpson

## References

Telford R.J. and Birks, H.J.B. (2011) A novel method for assessing the statistical significance of quantitative reconstructions inferred from biotic assemblages. Quanternary Science Reviews 30:1272-1278.

See Also
wa for details on fitting weighted average models.

## Examples

```
data(ImbrieKipp, SumSST, V12.122)
Cor <- weightedCor(ImbrieKipp, env = SumSST,
    fossil = V12.122, type = "simulate", sim = 49)
Cor
plot(Cor)
plot(Cor, type = "null")
```


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