Package 'cocorresp'

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2 cocorresp-package

cocorresp-package		Co-cor	respo	onde	nce o	inaly	vsis	ordi	nati	on i	neth	ods	for	com	тин	ıity	eco	l-
Index																		4
	<i>y</i> ,																	
	weights.symcoca .																	
	summary-methods.																	
	simpls																	
	screeplot.symcoca.																	
	scores-methods																	
	scaleChi																	
	resid.symcoca																	
	rescale																	
	points.coca																	
	plot.symcoca																	
	plot.predcoca																	
	permutest.coca																	
	mcChi																	
	loadings																	
	fitted.symcoca																	
	envfit.coca fitCoinertia																	
	crossval																	
	corAxis																	
	coinertial																	
	coinertia																	
																		1/

Description

Fits predictive and symmetric co-correspondence analysis (CoCA) models to relate one data matrix to another data matrix. More specifically, CoCA maximises the weighted covariance between the weighted averaged species scores of one community and the weighted averaged species scores of another community. CoCA attempts to find patterns that are common to both communitities.

Details

The main interface function is coca which accepts a formula or two community data matrices. An appropriate formula is $Y \sim .$, data = X and the associated data object from which . will be looked up. The method argument is used to select from the two forms of Coca: method = "predictive" for predictive Coca (the default), and method = "symmetric" for symmetric Coca.

Author(s)

Original Matlab routines by C.J.F. ter Braak and A.P. Schaffers. R port by Gavin L. Simpson. Function simpls based on simpls.fit (package pls) by Ron Wehrens and Bjorn-Helge Mevik.

Maintainer: Gavin L. Simpson <ucfagls@gmail.com>

beetles 3

beetles

Carabid beetles and vascular plants in Dutch roadside verges

Description

Counts of 126 beetle taxa and abundances of 173 vascular plant taxa (expressed on the 1-9 van der Maarel scale) in 30 road-side verges in the Netherlands.

Usage

```
data(beetles)
data(plants)
data(verges)
```

Details

This is the complete dataset of Raemakers et al (2001). ter Braak and Schaffers (2004) only analyse a subset of 91 beetle taxa.

Source

Raemakers, I.P., Schaffers, A.P., Sykora, V. and Heijerman, T. (2001) The importance of plant communities in road verges as habitat for insects. *Proceedings of the Section Experimental and Applied Entomology if the Netherlands Entomological Society* **12**, 101–106.

ter Braak, C.J.F and Schaffers, A.P. (2004) Co-Correspondence Analysis: a new ordination method to relate two community compositions. *Ecology* **85(3)**, 834–846

Examples

```
data(beetles)
data(plants)
```

biplot-methods

Biplots of co-correspondence analysis models

Description

Produces biplots of the response and predictor from the results of a co-correspondence analysis models.

4 biplot-methods

Usage

```
## S3 method for class 'symcoca'
biplot(x,
   which = "y1",
    choices = 1:2,
   benzecri = TRUE,
    type = NULL,
    xlim = NULL,
   ylim = NULL,
    col.species = "red",
    col.sites = "black",
    pch.species = 3,
    pch.sites = 1,
    cex = 0.7,
   main = ""
    sub = "",
    ylab, xlab,
    ann = par("ann"),
    axes = TRUE,
    ...)
## S3 method for class 'predcoca'
biplot(x,
   which = "response",
    choices = 1:2,
    type = NULL,
    xlim = NULL,
   ylim = NULL,
    col.species = "red",
    col.sites = "black",
    pch.species = 3,
    pch.sites = 1,
    cex = 0.7,
   main = "",
    sub = "",
   ylab, xlab,
    ann = par("ann"),
    axes = TRUE,
    ...)
```

Arguments

```
x an object of class "symcoca", the result of a call to symcoca.

which character; should the response or predictor scores be plotted. Can be specified in several ways: response choices are one from c("y", "Y", "y1", "response"); predictor choices are one from c("x", "X", "y2", "predictor").

choices a vector of length 2 indicating which predictive CoCA axes to plot.
```

biplot-methods 5

benzecri	logical, should a Benzecri plot be drawn? Such plots draw species scores, scaled by the quarter root of the respective eigenvalues, with unscaled site scores. A Benzecri plot is the recommended biplot for symmetric CoCA. See scores.symcoca.
type	one of "points", or "text". Determines how the site and species scores are displayed. If type = "points", scores are plotted as points. If type = "text", then the row names of the scores matrices are plotted.
xlim, ylim	limits for the x and y axes. If non supplied, suitable limits will be determined from the data.
col.species, co	ol.sites, pch.species, pch.sites
	colours and plotting characters used when plotting the species and sites scores.
cex	numeric; scaling factor when drawing points or text labels.
xlab, ylab	labels for the x and y axes. If non supplied suitable labels are formed from the result object.
main, sub	the main and sub titles for the plot.
ann	logical, if TRUE plots are annotated and not if FALSE, currently ignored.
axes	a logical value indicating whether axes and plot border should be drawn on the plot.
	other graphical parameters as in 'par' may also be passed as arguments.

Author(s)

Gavin L. Simpson.

References

Ter Braak, C.J.F and Schaffers, A.P. (2004) Co-Correspondence Analysis: a new ordination method to relate two community compositions. *Ecology* **85(3)**, 834–846

See Also

```
coca, plot.default
```

```
## symmetric CoCA
data(beetles)
data(plants)

## log transform the beetle data
beetles <- log(beetles + 1)

## fit the model
bp.sym <- coca(beetles ~ ., data = plants, method = "symmetric")

## draw a biplot of the beetle results
biplot(bp.sym)

## biplot of both - Fig 1 in ter Braak & Schaffers (2004)</pre>
```

6 bryophyte

```
layout(matrix(1:2, ncol = 2))
biplot(bp.sym, which = "y1", main = "Beetles")
biplot(bp.sym, which = "y2", main = "Plants")
layout(1)

## predictive CoCA
bp.pred <- coca(beetles ~ ., data = plants)

## draw a biplot of the response
biplot(bp.pred)

## recreate Fig 3 in ter Braak & Schaffers (2004)
layout(matrix(1:2, ncol = 2))
biplot(bp.pred, which = "response", main = "Beetles")
biplot(bp.pred, which = "predictor", main = "Plants")
layout(1)</pre>
```

bryophyte

Bryophytes and vascular plants in Carpathian spring meadows

Description

The data consist of observations on 30 bryophyte and 123 vascular plant species in 70 spring meadows (sites, samples). The species data are a subset of only those species occurring in five or more meadows.

Usage

```
data(bryophyte)
data(vascular)
```

Source

Hajek, M., Hekera, P. and Hajkova, P. (2002) Spring fen vegetations and water chemistry in the Western Carpathian flysch zone. *Folia geobotanica* **37**, 205–224

ter Braak, C.J.F and Schaffers, A.P. (2004) Co-Correspondence Analysis: a new ordination method to relate two community compositions. *Ecology* **85(3)**, 834–846

```
data(bryophyte)
data(vascular)
```

coca 7

coca

Fit Co-Correspondence Analysis Ordination Models

Description

coca is used to fit Co-Correspondence Analysis (CoCA) models. It can fit predictive or symmetric models to two community data matrices containing species abundance data.

Usage

```
coca(y, ...)
## Default S3 method:
coca(y, x, method = c("predictive", "symmetric"),
    reg.method = c("simpls", "eigen"), weights = NULL,
    n.axes = NULL, symmetric = FALSE, quiet = FALSE, ...)
## S3 method for class 'formula'
coca(formula, data, method = c("predictive", "symmetric"),
    reg.method = c("simpls", "eigen"), weights = NULL,
    n.axes = NULL, symmetric = FALSE, quiet = FALSE, ...)
```

Arguments

У	a data frame containing the response community data matrix.
X	a data frame containing the predictor community data matrix.
formula	a symbolic description of the model to be fit. The details of model specification are given below.
data	an optional 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 coca is called.
method	a character string indicating which co-correspondence analysis method to use. One of "predictive"(default), or "symmetric", can be abbreviated.
reg.method	One of "simpls" (default) or "eigen". If method is "predictive" then reg.method controls whether the co-correspondence analysis should be fitted using the SIM-PLS algorithm or via an eigen analysis.
weights	a vector of length nrow(y) of user supplied weights for R_0 . If weights = NULL (default) then the weights are determined from y (default) or x and y (symmetric = TRUE only).
n.axes	the number of CoCA axes to extract. If missing (default) the n. axes is
	$\min(ncol(y), ncol(x), nrow(y), nrow(x)) - 1$

.

8 coca

symmetric if method is "symmetric" then symmetric determines whether weights for R_0

are symmetric and taken as the average of the row sums of x and y (symmetric = TRUE). If symmetric = FALSE (default) then the weights R_0 are taken as the row sums of y unless user defined weights are provided via argument weights.

Ignored if method is "predictive".

quiet logical; suppress messages due to removal of species with no data.

... additional arguments to be passed to lower level methods.

Details

coca is the main user-callable function.

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.

The default is to fit a predictive CoCA model using SIMPLS via a modified version of simpls.fit from package pls. Alternatively, reg.method = "eigen" fits the model using an older, slower eigen analysis version of the SIMPLS algorithm. reg.method = "eigen" is about 100% slower than reg.method = "simpls".

Value

coca returns a list with method and reg. method determining the actual components returned.

nam.dat list with components namY and namX containing the names of the response and

the predictor(s) respectively.

call the matched call.

method the CoCA method used, one of "predictive" or "symmetric".

scores the species and site scores of the fitted model.

loadings the site loadings of the fitted model for the response and the predictor. (Predic-

tive CoCA via SIMPLS only.)

fitted the fitted values for the response. A list with 2 components Yhat (the fitted val-

ues on the original scale) and Yhat1 (the fitted values on the chi-square trans-

formed scale). (Predictive CoCA via SIMPLS only.)

varianceExp list with components Yblock and Xblock containing the variances in the re-

sponse and the predictor respectively, explained by each fitted PLS axis. (Pre-

dictive CoCA via SIMPLS only.)

totalVar list with components Yblock and Xblock containing the total variance in the

response and the predictor respectively. (Predictive CoCA via SIMPLS only.)

lambda the Eigenvalues of the analysis.

n.axes the number of fitted axes

Ychi a list containing the mean-centered chi-square matrices for the response (Ychi1)

and the predictor (Ychi2). (Predictive CoCA only.)

R0 the (possibly user-supplied) row weights used in the analysis.

coca 9

X X-Matrix (symmetric CoCA only).

residuals Residuals of a symmetric model (symmetric CoCA only).

inertia list with components total and residual containing the total and residual in-

ertia for the response and the predictor (symmetric CoCA only).

rowsum a list with the row sums for the response (rsum1) and the preditor (rsum2) (sym-

metric CoCA only).

colsum a list with the column sums for the response (csum1) and the preditor (csum2)

(symmetric CoCA only).

Author(s)

Original Matlab code by C.J.F. ter Braak and A.P. Schaffers. R port by Gavin L. Simpson. Formula method for coca uses a modified version of ordiParseFormula by Jari Oksanen to handle formulea.

References

ter Braak, C.J.F and Schaffers, A.P. (2004) Co-Correspondence Analysis: a new ordination method to relate two community compositions. *Ecology* **85**(3), 834–846

See Also

crossval for cross-validation and permutest.coca for permutation test to determine the number of PLS axes to retain in for predictive CoCA.

summary.predcoca and summary.symcoca for summary methods.

```
## symmetric CoCA
data(beetles)
## log transform the bettle data
beetles <- log(beetles + 1)
data(plants)
## fit the model
bp.sym <- coca(beetles ~ ., data = plants, method = "symmetric")</pre>
bp.sym
summary(bp.sym)
biplot(bp.sym)
                                         # produces a Benzecri biplot
## extract eigenvalues of the analysis
eigenvals(bp.sym)
## correlations between beetle and plant score scores on Co-CA axes
corAxis(bp.sym)
## predictive CoCA using SIMPLS and formula interface
bp.pred <- coca(beetles ~ ., data = plants)</pre>
## should retain only the useful PLS components for a parsimonious model
```

10 coinertia

```
## Leave-one-out crossvalidation - this takes a while
crossval(beetles, plants)
## so 2 axes are sufficient
## permutation test to assess significant PLS components - takes a while
bp.perm <- permutest(bp.pred, permutations = 99)</pre>
bp.perm
## agrees with the Leave-one-out cross-validation
## refit the model with only 2 PLS components
bp.pred <- coca(beetles ~ ., data = plants, n.axes = 2)</pre>
bp.pred
summary(bp.pred)
biplot(bp.pred)
                                     # plots correct scores or loadings
## predictive CoCA using Eigen-analysis
data(bryophyte)
data(vascular)
carp.pred <- coca(y = bryophyte, x = vascular, reg.method = "eigen")</pre>
## determine important PLS components - takes a while
crossval(bryophyte, vascular)
(carp.perm <- permutest(carp.pred, permutations = 99))</pre>
## 2 components again, refit
carp.pred <- coca(y = bryophyte, x = vascular,
                  reg.method = "eigen", n.axes = 2)
carp.pred
## drawn biplot
biplot(carp.pred)
```

coinertia

Co-inertia analysis

Description

Performs a co-inertia of the triplets (Q_1, K_1, R_0) and (Q_2, K_2, R_0) .

Usage

coinertia 11

Arguments

y, x	matrices or data frames of the two data sets for which axes of covariance are
	sought.

n.axes numeric; the number of coinertia axes to retain.

weights a vector of weights for the samples. If NULL, weights are chosen from the

row sums of y (symmetric = FALSE) or the average of the row sums of y and

x (symmetric = TRUE).

symmetric logical; should a symmetric set of weights be used. See Details.

... arguments passed to other functions. Currently ignored.

Value

An object of class "coinertia", which is a list with the following components:

scores a list of ordination scores, with components species and sites, each of which

is a list with components Y and X that refer to the scores for the input matrices y

and x respectively.

weights numeric vector of row weights used in the analysis.

lambda numeric vector of Eigenvalues.

n.axes numeric; the number of coinertia axes extracted symmetric logical; was a symmetric analysis performed?

call the matched call.

Author(s)

Gavin L. Simpson, based on original Matlab code by C.J.F. ter Braak and A.P. Schaffers.

References

Ter Braak, C.J.F and Schaffers, A.P. (2004) Co-Correspondence Analysis: a new ordination method to relate two community compositions. *Ecology* **85(3)**, 834–846

Doledec, S and Chessel, D. (1994) Co-inertia analysis: a method for studying species-environment relationships. *Freshwater Biology* **31**, 277–294

See Also

symcoca for the function that calls fitCoinertia and coinertial for co-inertia analysis using identity matrices for K_1 , K_2 , and R_0 .

```
data(beetles, plants)
coin <- coinertia(beetles, plants)
coin</pre>
```

12 coinertiaI

coinertiaI	Coinertia analysis with identity matrices	

Description

Performs a co-inertia of the triplets (Q_1, K_1, R_0) and (Q_2, K_2, R_0) with identity matrices K_1, K_2, R_0 .

Usage

```
coinertiaI(X, Y, fast = TRUE)
```

Arguments

X Species matrix X.Y Species Matrix Y.

fast If "TRUE" only return the row scores of Y.

Details

Argument fast is used to return only the row scores of Y in function permutest.coca, which speeds the permutation test considerably.

Value

If fast = TRUE, a matrix of row scores for matrix Y (see scores below). If fast = FALSE a list with the following components:

A list with components X and Y containing the left and right singular vectors respectivley of the SVD on the triplets.

A list with components X and Y, containing the row scores of the X and Y species matrices. These are the result of a matrix multiplication of X by the left singular vectors and Y by the right singular vectors.

The Eigenvalues of the analysis (the square of the singular values from the SVD.

call the matched function call.

Note

This function is not meant to be called directly by the user. If you wish to use it study the code in permutest.coca to see how it should be called.

Author(s)

Original Matlab code by C.J.F. ter Braak and A.P. Schaffers. R port by Gavin L. Simpson.

References

Doledec, S and Chessel, D. (1994) Co-inertia analysis: a method for studying species-environment relationships. *Freshwater Biology* **31**, 277–294.

corAxis 13

See Also

coinertia

corAxis

Correlation between ordination axes

Description

Calculates the Pearson product-moment correlation coefficient for the site scores of ordination axes.

Usage

```
corAxis(x, ...)
## Default S3 method:
corAxis(x, ...)
## S3 method for class 'symcoca'
corAxis(x, axes = NULL, ...)
```

Arguments

an ordination object. Only methods for objects of class symcoca are currently available.
 axes numeric; the number of axes to calculate the correlation coefficients for. If NULL, coefficients for all axes are returned.
 arguments to be passed on to other methods.

Value

A named vector containing the correlation coefficients for the requested axes.

Note

The arguments for cor are hard coded at their defaults, see cor for details. A more flexible version is planned that will allow arguments to be passed to cor.

Author(s)

Gavin L. Simpson

See Also

cor, for the main analysis function.

14 crossval

Examples

```
## load some data
data(beetles)
data(plants)

## log transform the beetle data
beetles <- log(beetles + 1)

## symmetric Co-CA model
beetles.sym <- coca(beetles ~ ., data = plants, method = "symmetric")

## correlations between axes
corAxis(beetles.sym)</pre>
```

crossval

Cross-validation for predictive Co-Correspondence Analysis models

Description

Performs a leave-one-out cross-validation of a predictive Co-Correspondence Analysis model.

Usage

Arguments

У	the response species matrix.
X	the predictor species matrix.
n.axes	the number of axes to calculate the leave-one-out cross-validation for. Default is to perform the ${\rm CV}$ for all extractable axes.
centre	centre y and x during analysis? Currently ignored as it may not be neccessary.
verbose	if TRUE, the default, print information on the progress of the cross-validation procedure.
object	an object of class crossval as returned by crossval.
axes	the number of axes to summarise results for.
	further arguments to print - currently ignored.

crossval 15

Details

Performs a leave-one-out cross-validation of a predictive Co-Correspondence Analysis model. It can be slow depending on the number of columns in the matrices, and of course the number of sites.

Value

Returns a large list with the following components:

dimx, dimy the dimensions of the input matrices x and y respectively.

press0 the $press_0$ statistic.

n.axes the number of axes tested.

CVfit the cross-validatory fit.

varianceExp list with components Yblock and Xblock containing the variances in the re-

sponse and the predictor respectively, explained by each fitted PLS axis.

totalVar list with components Yblock and Xblock containing the total variance in the

response and the predictor respectively.

nam.dat list with components namY and namX containing the names of the response and

the predictor(s) respectively.

call the R call used.

Note

This function is not a bit out-of-date compared to some of the other functions. It should have a formular interface like coca or work on the results from coca, although that will have to be altered to store a copy of the data?

Author(s)

Gavin L. Simpson, based on Matlab code by C.J.F. ter Braak and A.P. Schaffers.

See Also

The model fitting function coca

```
## load the data sets
data(beetles)
data(plants)

## log transform the bettle data
beetles <- log(beetles + 1)

## predictive CoCA using SIMPLS and formula interface
bp.pred <- coca(beetles ~ ., data = plants)
## should retain only the useful PLS components for a
## parsimonious model</pre>
```

16 envfit.coca

```
## Leave-one-out crossvalidation - this takes a while
## Not run:
crossval(beetles, plants)

## End(Not run)
## so 2 axes are sufficient
```

envfit.coca

Fit an environmental vector or factor to a Co-CA ordination

Description

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

Usage

Arguments

ord	a Co-CA ordination object, the result of a call to coca.
env	a data frame, matrix or vector of environmental/external variable(s) to be fitted to the ordination. The variables may be of a mixed type (factors and continuous variables) in a data frame.
which	character; which of the response or predictor ordinations should be used during fitting of vectors and factors.
choices	numeric; the axes to which vectors and factors are fitted.
scaling	logical; should sacling be applied. See scores.symcoca.
W	weights used in fitting vectors and factors.
na.rm	Remove points with missing values in ordination scores or environmental variables. The operation is casewise; the whole row of data is removed if there is a missing value and na.rm = TRUE.
strata	An integer vector or factor specifying the strata for permutation. If supplied, observations are permuted only within the specified strata.
permutations	Number of permutations for assessing significance of vectors or factors. Set to 0 to skip permutations.
	Arguments passed to vectorfit and factorfit.

fitCoinertia 17

Details

See envfit for details of the method.

Value

Returns an object of class envfit.

Author(s)

Gavin L. Simpson. The code interfaces with and uses code from envfit for the main computations, which was written by Jari Oksanen.

See Also

coca for fitting models. envfit for details of the generic function and the computations performed.

Examples

```
## symmetric CoCA
data(beetles)
data(plants)

## log transform the bettle data
beetles <- log(beetles + 1)

## fit the model
bp.sym <- coca(beetles ~ ., data = plants, method = "symmetric")

## load the environmental data
data(verges)

## fit vectors for the environmental data
sol <- envfit(bp.sym, verges, which = "response")
sol

## plot the response matrix and the fitted vectors
biplot(bp.sym, which = "y1")
plot(sol)</pre>
```

fitCoinertia

Co-inertia analysis

Description

Performs a co-inertia of the triplets (Q_1, K_1, R_0) and (Q_2, K_2, R_0) .

18 fitCoinertia

Usage

```
fitCoinertia(X, Dp, Y, Dq, Dn, n.axes)
## S3 method for class 'fitCoinertia'
summary(object, ...)
```

Arguments

Χ	Q_1 , matrix of expected abundances under row-column independence in the original Y species matrix when treated as a contingency table.
Dp	K_1 , species (column) weights for X.
Υ	Q_2 , matrix of expected abundances under row-column independence in the original X species matrix when treated as a contingency table.
Dq	K_2 , species (column) weights for Y.
Dn	site weights R_0 .
n.axes	number of axes to calculate the co-inertia analysis for.
object	an object of class coinertia.
axes	the number of axes to display when printing.
	arguments passed to other functions. Currently ignored.

Value

A list with the following components:

U1	column weights of X.
U2	column weights of Y.
X1	rowscores of X.
X2	rowscores of Y.

lambda the Eigenvalues (squares of the singular values).

n. axes number of axes requested.
call the matched function call.

Note

This function is not yet meant to be called directly by the user. If you wish to use it directly, see the function definition for symcoca which demonstrates how to prepare the relevant input matrices.

Note that in this function, X corresponds to the input matrix y and Y corresponds to the input matrix x in symcoca. Confusing! This will be changed in a future release but for now the arguments follow those of the original Matlab code - perhaps a little too closely!

fitted.symcoca 19

Author(s)

Original Matlab code by C.J.F. ter Braak and A.P. Schaffers. R port by Gavin L. Simpson.

References

Ter Braak, C.J.F and Schaffers, A.P. (2004) Co-Correspondence Analysis: a new ordination method to relate two community compositions. *Ecology* **85**(3), 834–846

Doledec, S and Chessel, D. (1994) Co-inertia analysis: a method for studying species-environment relationships. *Freshwater Biology* **31**, 277–294

See Also

symcoca for the function that calls fitCoinertia and coinertial for co-inertia analysis using identity matrices for K_1 , K_2 , and R_0

fitted.symcoca

Fitted values of a Symmetric Co-Correpsondence analysis model.

Description

Calculates and extracts the fitted values of a Symmetric Co-Correpsondence analysis model.

Usage

```
## S3 method for class 'symcoca'
fitted(object, which = c("y1","y2"), ...)
```

Arguments

object an object of class "symcoca"

which character; should the response or predictor scores be plotted. Can be specified in

several ways: response choices are one from c("y", "Y", "y1", "response");

predictor choices are one from c("x","X","y2","predictor").

... arguments to be passed to other methods.

Value

A list with the following components:

Y the fitted values for the "response" matrix.

X the fitted values for the "predictor" matrix.

nam.dat a vector containing the names of the "response" and "predictor" matrices re-

spectively. Used for printing the results.

20 loadings

Note

This function needs an update and to allow option to restrict fitted values to specified axes, and the names of the returned objects need making more obvious!

Author(s)

Gavin L. Simpson, based on Matlab code by C.J.F. ter Braak and A.P. Schaffers.

References

Ter Braak, C.J.F and Schaffers, A.P. (2004) Co-Correspondence Analysis: a new ordination method to relate two community compositions. *Ecology* **85(3)**, 834–846

See Also

The model fitting function coca

Examples

```
## symmetric CoCA
data(beetles)
data(plants)

## log transform the bettle data
beetles <- log(beetles + 1)

## fit the model
bp.sym <- coca(beetles ~ ., data = plants, method = "symmetric")

## fitted values
bp.fit <- fitted(bp.sym)
bp.fit

## fitted values for beetles only
beetle.fit <- fitted(bp.sym, which = "y1")</pre>
```

loadings

CoCA species loadings

Description

Extract CoCA species loadings from fitted objects.

loadings 21

Usage

Arguments

x an object resulting from a call to coca.

choices numeric; vector of Co-CA axes to extract loadings for.

which character; should the response or predictor scores be plotted. Can be specified in several ways: response choices are one from c("y", "Y", "y1", "response"); predictor choices are one from c("x", "X", "y2", "predictor").

... additional arguments to be passed to lower level methods.

Details

loadings() is an extractor function to access the loadings of a fitted CoCA model.

This is a generic function, replacing the loadings function, which is preserved as the exported default S3 method. Methods are provided for both predictive and symmetric CoCA.

Value

A list of data frames or a single data frame depending on other arguments.

Author(s)

Gavin L. Simpson

See Also

coca for how to fit CoCA models.

```
## symmetric CoCA
data(beetles)
## log transform the bettle data
beetles <- log(beetles + 1)
data(plants)
## fit the model
bp.sym <- coca(beetles ~ ., data = plants, method = "symmetric")</pre>
```

22 mcChi

```
## extract the loadings
lds <- loadings(bp.sym)</pre>
```

mcChi

Standardised chi-square residuals

Description

Scales a matrix, Y, to its standardised chi-square residuals $(o-e)/\sqrt{e}$ (if $R_0=R$, where R contains the row sums of matrix Y) so that further analysis can be unweighted

Usage

```
mcChi(Y, R0, eps = 1e-06)
```

Arguments

Y a matrix for which standardised chi-square residuals are to be calculated.

R0 row weights.

eps tolerance - leave as default.

Details

This function implements equation 8 of ter Braak and Schaffers (2004) by firstly applying equation 7 to form matrix Q using row and column sums of Y as weights, and, secondly, by applying equation 8 to form a matrix of standardised chi-square residuals from Q by pre-multiplication of Q by $\sqrt{R_0}$ and post-multiplication of Q by \sqrt{K} , where K is the column sums of Y.

Value

A list with the following components:

Ychi the matrix of standardised chi-squared residuals of Y

Kn the column sums (K) of Y divided by sum(K)

Note

This function is not intended for casual use by users.

Author(s)

Gavin L. Simpson, based on Matlab code by C.J.F. ter Braak and A.P. Schaffers.

References

Ter Braak, C.J.F and Schaffers, A.P. (2004) Co-Correspondence Analysis: a new ordination method to relate two community compositions. *Ecology* **85(3)**, 834–846

permutest.coca 23

permutest.	coca	Permutation test for predictive co-correspondence analysis models

Description

A permutation test for predictive co-correspondence analysis models to assess the significance of each CoCA ordination axes.

Usage

Arguments

x	an object of class "predcoca".
R0	row weights to use in the analysis. If missing, the default, these are determined from x.
permutations	the number of permutations to perform.
n.axes	The number of axes to test. Defaults to the number of axes stated in x\$n.axes.
verbose	if TRUE, the default, print information on the progress of the permutation test procedure.
object	an object of class "permutest.coca".
	arguments to be passed to other methods.

Details

An alternative approach to cross-validation (see crossval) to select the number of axes to retain in a predictive co-correspondence analysis is to test the statistical significance of each ordination axis using permutation tests.

The test statistic used is the F-ratio based on the fit of the first axis to the response data (ter Braak and Smilauer 2002). The second and subsequent axes are tested by treating previous axes as covariables.

To be precise, this approach does not test the significance of SIMPLS axes, but those of NIPALS-PLS axes (ter Braak and de Jong 1998).

24 permutest.coca

Value

A list with the following components:

pval a vector of *P*-values for each ordination axis.

permstat a vector of values for the test statistic for each axis.

total.inertia the total inertia in the response matrix.

inertia a vector containing the residualised inertia. This is the total inertia in the re-

sponse after removing the inertia explained by all previous axes. For the first

CoCA axis this is, by definition, the total inertia in the response.

fitax a vector containing the amount of inertia in the response matrix explained by

each ordination axis.

pcent.fit a vector containing the fit of each axis to the response as a percentage of the

total inertia (variance).

n. axes the number of axes in the ordination.

call the matched call.

Warning

This function is **slow**. Beware setting argument permutations higher than the default. Determine how long it takes for the default 99 permutations to complete before going crazy and asking for thousands of permutations - you've been warned, have a good book to hand.

Note

Argument R0 is provided for compatibility with the original MATLAB code. The R usage paradigm makes this argument redundant in the current code and it may be invalid to supply different row weights (R_0) as R0. This argument will likely be removed in future versions.

Author(s)

Gavin L. Simpson, based on Matlab code by C.J.F. ter Braak and A.P. Schaffers.

References

ter Braak, C.J.F. and de Jong, S. (1998) The objective function of partial least squares regression. *Journal of Chemometrics* **12**, 41–54.

ter Braak, C.J.F and Schaffers, A.P. (2004) Co-Correspondence Analysis: a new ordination method to relate two community compositions. *Ecology* **85**(3), 834–846.

ter Braak, C.J.F. and Smilauer, P. (2002) Canoco reference manual and CanoDraw for Windows user's guide: software for canonical community ordination. Version 4.5. New York: Microcomputer Power.

See Also

coca, for the model fitting function, crossval, for a leave-one-out cross-validation procedure, which is the preferred way to select axes in a predictive co-correspondence analysis.

plot.predcoca 25

Examples

```
## load some data
data(beetles)
data(plants)

## log transform the bettle data
beetles <- log(beetles + 1)
## predictive CoCA using SIMPLS and formula interface
bp.pred <- coca(beetles ~ ., data = plants)

## should retain only the useful PLS components for a parsimonious model

## Leave-one-out crossvalidation - this takes a while
crossval(beetles, plants)

## so 2 axes are sufficient

## permutation test

## (Testing the first 2 axes & only 50 perms for speed.)
bp.perm <- permutest(bp.pred, permutations = 50, n.axes = 2)
bp.perm</pre>
```

plot.predcoca

Biplots for predictive co-correspondence analysis

Description

Produces biplots of the response and predictor from the results of a predictive co-correspondence analysis.

Usage

```
## S3 method for class 'predcoca'
plot(x,
    which = "response",
    choices = 1:2,
    display = c("species", "sites"),
    type,
    xlim = NULL,
    ylim = NULL,
    main = "",
    sub = "",
    ylab, xlab,
    ann = par("ann"),
    axes = TRUE,
    ...)
```

26 plot.predcoca

Arguments

x	an object of class "predcoca", the result of a call to coca.
which	character; should the response or predictor scores be plotted. Can be specified in several ways: <i>response</i> choices are one from c("y","Y","y1","response"); <i>predictor</i> choices are one from c("x","X","y2","predictor").
choices	a vector of length 2 indicating which predictive CoCA axes to plot.
display	which sets of scores are drawn. See scores.predcoca.
type	one of "points", "text", or "none". Determines how the site and species scores are displayed. If type = "points", scores are plotted as points. If type = "text", then the row names of the scores matrices are plotted. If type = "none", then the scores are not plotted.
xlim, ylim	limits for the x and y axes. If non supplied, suitable limits will be determined from the data.
xlab, ylab	labels for the x and y axes. If non supplied suitable labels are formed from the result object.
main, sub	the main and sub titles for the plot.
ann	logical, if TRUE plots are annotated and not if FALSE, currently ignored.
axes	a logical value indicating whether both axes should be drawn on the plot.
	other graphical parameters as in 'par' may also be passed as arguments.

Author(s)

Gavin L. Simpson.

References

Ter Braak, C.J.F and Schaffers, A.P. (2004) Co-Correspondence Analysis: a new ordination method to relate two community compositions. *Ecology* **85(3)**, 834–846

See Also

```
coca, plot.default
```

```
## predictive CoCA
data(beetles)
data(plants)

## log transform the beetle data
beetles <- log(beetles + 1)

## predictive CoCA using SIMPLS and formula interface
bp.pred <- coca(beetles ~ ., data = plants)

## draw the plot for the response scores
plot(bp.pred)</pre>
```

plot.symcoca 27

```
## plot of both
layout(matrix(1:2, ncol = 2))
plot(bp.pred, which = "response", main = "Beetles")
plot(bp.pred, which = "predictor", main = "Plants")
layout(1)
```

plot.symcoca

Plots for symmetric co-correspondence analysis

Description

Produces plots of the response and predictor from the results of a symmetric co-correspondence analysis.

Usage

```
## S3 method for class 'symcoca'
plot(x,
    which = "response",
    choices = 1:2,
    display = c("species", "sites"),
    scaling = FALSE,
    type,
    xlim = NULL,
    ylim = NULL,
    main = "",
    sub = "",
    ylab, xlab,
    ann = par("ann"),
    axes = TRUE,
    ...)
```

Arguments

X	an object of class "symcoca", the result of a call to symcoca.
which	character; should the response or predictor scores be plotted.
choices	a vector of length 2 indicating which predictive CoCA axes to plot.
display	which sets of scores are drawn. See scores.symcoca.
scaling	logical, whether scaling should be applied. See scores.symcoca.
type	one of "points", "text", or "none". Determines how the site and species scores are displayed. If type = "points", scores are plotted as points. If type = "text", then the row names of the scores matrices are plotted. If type = "none", then the scores are not plotted.
xlim, ylim	limits for the x and y axes. If non supplied, suitable limits will be determined from the data.

28 points.coca

xlab, ylab	labels for the x and y axes. If non supplied suitable labels are formed from the result object.
main, sub	the main and sub titles for the plot.
ann	logical, if TRUE plots are annotated and not if FALSE, currently ignored.
axes	a logical value indicating whether both axes should be drawn on the plot.
	other graphical parameters as in 'par' may also be passed as arguments.

Author(s)

Gavin L. Simpson.

References

Ter Braak, C.J.F and Schaffers, A.P. (2004) Co-Correspondence Analysis: a new ordination method to relate two community compositions. *Ecology* **85(3)**, 834–846

See Also

```
coca, plot.default
```

Examples

```
## symmetric CoCA
data(beetles)
data(plants)

## log transform the beetle data
beetles <- log(beetles + 1)

## fit the model
bp.sym <- coca(beetles ~ ., data = plants, method = "symmetric")

## draw a plot of the response scores
plot(bp.sym)

## plot of both
layout(matrix(1:2, ncol = 2))
plot(bp.sym, which = "response", main = "Beetles")
plot(bp.sym, which = "predictor", main = "Plants")
layout(1)</pre>
```

points.coca

Add points to a Co-CA plot

Description

Draws points on the current graphic device based on in supplied coca model object.

points.coca 29

Usage

Arguments

x	an object inheriting from class coca.
display	character; one of the stated choices. Indicates which scores to use to draw points.
which	character; one of the stated choices. Indicates which of the response or predictor data sets is used to select scores from.
choices	The Co-CA axes to draw points for.
scaling	logical; should species scores in a symmetric Co-CA be rescaled?
select	Items to be displayed. This can either be a logical vector which is TRUE for displayed items or a vector of indices of displayed items.
	Arguments passed to other methods

Details

The visual appearance of the plotted points can be controlled by supplying appropriate graphical parameters via the . . . argument. See par for details.

Value

Returns the plotted x and y coordinates as a matrix.

Author(s)

```
Gavin L. Simpson
```

See Also

```
plot methods; plot.predcoca and plot.symcoca.
```

```
## symmetric CoCA
data(beetles)
data(plants)

## log transform the bettle data
beetles <- log(beetles + 1)

## fit the model
bp.sym <- coca(beetles ~ ., data = plants, method = "symmetric")

## draw a plot of the response scores</pre>
```

30 rescale

```
plot(bp.sym, type = "none")
points(bp.sym, display = "sites", col = "blue", pch = 16)
points(bp.sym, display = "species", col = "red", pch = 3, cex = 0.8)
```

rescale

Rescales CoCA species scores

Description

Rescales CoCA species scores to the quarter root of the eigenvalues.

Usage

Arguments

object an R object. Currently only objects of class "symcoca" are supported.
choices numeric; which axes to rescale and return.
display character; which type of scores to rescale.
... other arguments to be passed to rescale methods. Currently not used.

Details

Currently only implemented for objects of class "symcoca".

Value

Returns a list with one or both of the following components:

species rescaled species scores for the response sites rescaled species scores for the preditor

Author(s)

Matlab original by C.J.F. ter Braak and A.P. Schaffers. R port by Gavin L. Simpson.

See Also

```
symcoca
```

resid.symcoca 31

Examples

```
data(bryophyte)
data(vascular)
bryo.sym <- coca(bryophyte ~ ., data = vascular, method = "symmetric")
rescale(bryo.sym, axes = 1:2)</pre>
```

resid.symcoca

Extract Model Residuals

Description

Extracts the residuals of the fitted model of a symmetric CoCA to the response and the predictor.

Usage

```
## S3 method for class 'symcoca'
resid(object, ...)
```

Arguments

object an object of class "symcoca".
... arguments to be passed to other methods.

Value

A list containing the residuals for the response and the predictor with the following components:

Y residuals of the fit to the repsonse.
X residuals of the fit to the predictor.

Author(s)

Gavin L. Simpson

See Also

symcoca

```
data(bryophyte)
data(vascular)
bryo.sym <- coca(bryophyte ~ ., data = vascular, method = "symmetric")
eps <- resid(bryo.sym)</pre>
```

32 scaleChi

sca	1 -/	∼I -	•
SCA	10	n	1

Standardised chi-square residuals

Description

Scales a matrix, Y, to is standardised chi-square residuals $(o-e)/\sqrt{e}$ (given K_n and R_0 metrics derived from an external matrix Y_0) so that further analysis can be unweighted.

Usage

```
scaleChi(Y, Kn, R0, eps = 1e-06)
```

Arguments

Y a matrix for which standardised chi-square residuals are to be calculated.

Kn the column sums (K) of Y divided by sum(K).

R0 row weights.
eps a tolerance.

Value

Yr the matrix of standardised chi-squared residuals of Y.

Note

This function is not intended for casual use by users.

Author(s)

Gavin L. Simpson, based on Matlab code by C.J.F. ter Braak and A.P. Schaffers.

References

Ter Braak, C.J.F and Schaffers, A.P. (2004) Co-Correspondence Analysis: a new ordination method to relate two community compositions. *Ecology* **85(3)**, 834–846

scores-methods 33

scores-methods Get Species or Si	te Scores from an Ordination
----------------------------------	------------------------------

Description

Function to access either species or site scores for specified axes in co-correspondence analysis ordination methods.

Usage

Arguments

X	an ordination result
display	partial match to access scores for "sites" "species", "loadings" or "xmatrix". The latter two are only available for symcoca.
choices	numeric; the ordination axes to return.
scaling	logical; whether scores should be rescaled by the quarter root of the eigenvalues using rescale.symcoca.
	arguments to be passed to other methods.

Details

Implements a scores method for symmetric co-correspondence analysis ordination results.

Value

A list with one or more components containing matrices of the requested scores:

species	A list with two components, Y and X, containing the species scores for the response matrix Y and the predictor matrix X respectively.
sites	A list with two components, Y and X, containing the site scores for the response matrix Y and the predictor matrix X respectively.
loadings	A list with two components, Y and X containing the loadings for the response and predictor matrix. For symcoca only.
xmatrix	The X matrix. For symcoca only.

34 screeplot.symcoca

Author(s)

Gavin L. Simpson, based on Matlab code by C.J.F. ter Braak and A.P. Schaffers.

References

ter Braak, C.J.F and Schaffers, A.P. (2004) Co-Correspondence Analysis: a new ordination method to relate two community compositions. *Ecology* **85**(3), 834–846

See Also

scores, for further details on the method.

Examples

```
## load some data
data(beetles)
data(plants)

## log transform the bettle data
beetles <- log(beetles + 1)

## fit the model, a symmetric CoCA
bp.sym <- coca(beetles ~ ., data = plants, method = "symmetric")

## extract the scores
scr <- scores(bp.sym)

## predictive CoCA using SIMPLS and formula interface
bp.pred <- coca(beetles ~ ., data = plants)
scr2 <- scores(bp.pred)</pre>
```

screeplot.symcoca

Screeplot for symmetric co-correspondence analysis

Description

Produces a screeplot of the eigenvalues of a symmetric co-correspondence analysis.

Usage

simpls 35

Arguments

x an object of class "symcoca", the result of a call to symcoca.
 type the type of points to draw; see points.
 xlab, ylab labels for the x and y axes. If none supplied suitable labels are formed from the result object.
 other graphical parameters as in 'par' may also be passed as arguments.

Author(s)

Gavin L. Simpson.

See Also

```
coca, screeplot, plot.default
```

Examples

```
## symmetric CoCA
data(beetles)
data(plants)

## log transform the bettle data
beetles <- log(beetles + 1)

## fit the model
bp.sym <- coca(beetles ~ ., data = plants, method = "symmetric")

## draw a screeplot of the eignevalues
screeplot(bp.sym)</pre>
```

 ${\tt simpls}$

Modified version of Sijmen de Jong's SIMPLS

Description

Fits a PLSR model with the SIMPLS algorithm, modified to allow a weighted analysis.

Usage

```
simpls(X, Y, ncomp, stripped = FALSE, ...)
```

36 simpls

Arguments

X a matrix of observations. NAs and Infs are not allowed.

Y a vector or matrix of responses. NAs and Infs are not allowed.

ncomp the number of components to be used in the modelling.

stripped logical. If TRUE the calculations are stripped as much as possible for speed; this

is meant for use with cross-validation or simulations when only the coefficients

are needed. Defaults to FALSE.

... other arguments. Currently ignored.

Details

This function is a modified version of simpls. fit from package pls. Four modification have been made:

1. The input matrices X and Y are not centered,

2. The scores (tt in the code) are not centered,

3. Added code to calculate the total variance in the Y matrix, Ytotvar, and the variance in Y accounted for by each PLS axis, Yvar (See Value below), and

4. Additional components are returned if argument stripped is TRUE.

This function should not be called directly, but through the generic function coca.

SIMPLS is much faster than the NIPALS algorithm, especially when the number of X variables increases, but gives slightly different results in the case of multivariate Y. SIMPLS truly maximises the covariance criterion. According to de Jong, the standard PLS2 algorithms lie closer to ordinary least-squares regression where a precise fit is sought; SIMPLS lies closer to PCR with stable predictions.

Value

A list containing the following components is returned:

coefficients an array of regression coefficients for 1, ..., ncomp components. The dimen-

sions of coefficients are c(nvar, npred, ncomp) with nvar the number of X

variables and npred the number of variables to be predicted in Y.

scores a matrix of scores.

loadings a matrix of loadings.

Yscores a matrix of Y-scores.

Yloadings a matrix of Y-loadings.

projection the projection matrix used to convert X to scores.

Xmeans a vector of means of the X variables.

Ymeans a vector of means of the Y variables.

fitted.values an array of fitted values. The dimensions of fitted.values are c(nobj, npred, ncomp)

with nobj the number samples and npred the number of Y variables.

residuals an array of regression residuals. It has the same dimensions as fitted.values.

summary-methods 37

Xvar a vector with the amount of X-variance explained by each number of compo-

nents.

Yvar a vector with the amount of Y-variance explained by each number of compo-

nents.

Xtotvar Total variance in X. Ytotvar Total variance in Y.

If stripped is TRUE, only the components coefficients, Xmeans and Ymeans, Xvar and Yvar, and Xtotvar and Ytotvar are returned.

Author(s)

Based on simpls.fit by Ron Wehrens and Bjorn-Helge Mevik, with simple modifications by Gavin L. Simpson.

References

de Jong, S. (1993) SIMPLS: an alternative approach to partial least squares regression. *Chemometrics and Intelligent Laboratory Systems*, **18**, 251–263.

See Also

coca

summary-methods

Summarizing Co-CA Model Fits

Description

summary methods for classes "predcoca" and "symcoca". These provide a summary of the main results of a Co-Correspondence Analysis model.

Usage

```
## $3 method for class 'predcoca'
summary(object, axes = NULL, ...)
## $3 method for class 'symcoca'
summary(object, ...)
```

Arguments

object an object of class "predcoca" or "symcoca". Generally the result of a call to

coca.

axes numeric; how many axes to summarise? The default is to display 6 axes or all

available axes, whichever is the smaller.

. . . arguments to be passed to other methods.

38 summary-methods

Value

A list with the some of the following components:

cocaScores The site and/or species scores for the axes requested.

call The call used to fit the model.

lambda The eigenvalues for the axes requested. Not for predcoca.simpls.

namY, namX the names of the response and predictor either supplied by the user or derived

from the original call.

loadings a list with two components loadings1 and loadings2, which refer to the re-

sponse and the predictor matrices respectively. (Only for predictive CoCA mod-

els.)

varianceExp a list with components Yblock and Xblock containing the amount of variance

explained on each CoCA axis in the response and the predictor respectively.

(Only for predictive CoCA models.)

totalVar a list with components Yblock and Xblock containing the total variance in the

response and the predictor data sets respectively

inertia a list with components total and residual containing the total and residual in-

ertia (variance) in the response and the predictor matrices of a symmetric CoCA

model. (Only for symmetric CoCA models.)

scaling the scaling used/requested. (Only for symmetric CoCA models.)

Author(s)

Gavin L. Simpson

See Also

The model fitting function coca

```
## symmetric CoCA
data(beetles)
data(plants)

## log transform the bettle data
beetles <- log(beetles + 1)

## fit the model
bp.sym <- coca(beetles ~ ., data = plants, method = "symmetric")
summary(bp.sym)

## Predictive CoCA
bp.pred <- coca(beetles ~ ., data = plants)
summary(bp.pred, axes = 1:2)</pre>
```

weights.symcoca 39

weights.symcoca

Exctract CoCA model weights

Description

Extractor function to identify and select appropriate analysis weights from Co-correspondence Analysis models.

Usage

```
## S3 method for class 'symcoca'
weights(object, ...)
## S3 method for class 'predcoca'
weights(object, ...)
```

Arguments

```
object an object of class "symcoca" or "predcoca".
... arguments passed to other methods. Not used.
```

Value

A numeric vector of common site weights is currently returned. These correspond to R_0 in ter Braak and Schaffers (2004).

Author(s)

Gavin L. Simpson

References

ter Braak, C.J.F and Schaffers, A.P. (2004) Co-Correspondence Analysis: a new ordination method to relate two community compositions. *Ecology* **85**(3), 834–846.

See Also

envfit.coca uses these weights to estimate weighted correlations in ordination space. See coca for details on creating CoCA models.

```
## symmetric CoCA
data(beetles)
data(plants)
## log transform the bettle data
```

40 weights.symcoca

```
beetles <- log(beetles + 1)
## fit the model
bp.sym <- coca(beetles ~ ., data = plants, method = "symmetric")
## weights == R[0]
weights(bp.sym)</pre>
```

Index

*Topic datasets	*Topic package
beetles, 3	cocorresp-package, 2
bryophyte, 6	*Topic regression
*Topic hplot	coca, 7
biplot-methods, 3	simpls, 35
plot.predcoca, 25	*Topic univar
plot.symcoca, 27	corAxis, 13
screeplot.symcoca,34	*Topic utilities
*Topic htest	loadings, 20
envfit.coca, 16	weights.symcoca, 39
*Topic methods	
biplot-methods, 3	beetles, 3
loadings, 20	biplot-methods, 3
plot.symcoca, 27	biplot.predcoca(biplot-methods), 3
points.coca, 28	biplot.symcoca(biplot-methods), 3
scores-methods, 33	bryophyte, 6
screeplot.symcoca, 34	
weights.symcoca, 39	coca, 2, 5, 7, 15–17, 20, 21, 24, 26, 28, 29,
*Topic models	35–39
coca, 7	cocorresp (cocorresp-package), 2
coinertia, 10	cocorresp-package, 2
coinertiaI, 12	coinertia, 10, <i>13</i>
fitCoinertia, 17	coinertiaI, 11, 12, 19
*Topic multivariate	cor, <i>13</i>
coca, 7	corAxis, 13
coinertia, 10	crossval, 9, 14, 23, 24
coinertiaI, 12	Ci+ 17
crossval, 14	envfit, 17
envfit.coca, 16	envfit.coca, 16, <i>39</i>
fitCoinertia, 17	factorfit, 16
fitted.symcoca, 19	fitCoinertia, 17
mcChi, 22	fitted.symcoca, 19
permutest.coca, 23	Titted. Syllicoca, 19
rescale, 30	loadings, 20, 21
resid.symcoca, 31	lodaings, 20, 27 lodaings.default (loadings), 20
scaleChi, 32	100011165.0010011165/, 20
scores-methods, 33	mcChi, 22
simpls, 35	,
summary-methods, 37	ordiParseFormula,9
3	,-

INDEX

par, 29
permutest.coca, 9, 12, 23
plants (beetles), 3
plot, 29
plot.default, 5, 26, 28, 35
plot.predcoca, 25, 29
plot.symcoca, 27, 29
points, 35
points.coca, 28
predcoca.simpls, 38
print.crossval (crossval), 14
print.fitCoinertia (fitCoinertia), 17
print.fitted.symcoca(fitted.symcoca),
19
print.permutest.coca (permutest.coca),
23
print.predcoca(coca), 7
print.summary.crossval (crossval), 14
<pre>print.summary.fitCoinertia</pre>
(fitCoinertia), 17
<pre>print.summary.permutest.coca</pre>
(permutest.coca), 23
print.summary.predcoca
(summary-methods), 37
print.summary.symcoca
(summary-methods), 37
print.symcoca (coca), 7
p. 1
rescale, 30
rescale.symcoca, 33
resid.symcoca, 31
1 0014.03 1110004, 31
scaleChi, 32
scores, 33, 34
scores-methods, 33
scores.predcoca, 26
scores.predcoca (scores-methods), 33
scores.symcoca, 5, 16, 27
scores.symcoca (scores-methods), 33
screeplot, 35
screeplot.symcoca, 34
simpls, 35
simpls.fit, 8, 36, 37
summary-methods, 37
summary.crossval (crossval), 14
summary.fitCoinertia(fitCoinertia), 17
summary.permutest.coca
(permutest.coca), 23

```
summary.predcoca (summary-methods), 37 summary.symcoca, 9 summary.symcoca (summary-methods), 37 symcoca, 4, 11, 13, 18, 19, 27, 30, 31, 33, 35 vascular (bryophyte), 6 vectorfit, 16 verges (beetles), 3 weights.predcoca (weights.symcoca), 39 weights.symcoca, 39
```