Package: multiblock (via r-universe)

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```
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     interfaces, result- and plotting functions, several real data
     sets and six vignettes covering a range different applications.
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asca

Analysis of Variance Simultaneous Component Analysis - ASCA

Description

This is a quite general and flexible implementation of ASCA.

Usage

```
asca(formula, data, subset, weights, na.action, family, pca.in = FALSE)
```

Arguments

| formula | Model formula accepting a single response (block) and predictor names separated by + signs. |
|-----------|---|
| data | The data set to analyse. |
| subset | Subset of objects |
| weights | Optional object weights. |
| na.action | How to handle NAs (no action implemented). |
| family | Error distributions and link function for Generalized Linear Models. |
| pca.in | Compress response before ASCA (number of components). |

Details

ASCA is a method which decomposes a multivariate response according to one or more design variables. ANOVA is used to split variation into contributions from factors, and PCA is performed on the corresponding least squares estimates, i.e., Y = X1 B1 + X2 B2 + ... + E = T1 P1' + T2 P2' + ... + E. This version of ASCA encompasses variants of LiMM-PCA, generalized ASCA and covariates ASCA. It includes confidence ellipsoids for the balanced fixed effect ASCA.

Value

An asca object containing loadings, scores, explained variances, etc. The object has associated plotting (asca_plots) and result (asca_results) functions.

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References

• Smilde, A., Jansen, J., Hoefsloot, H., Lamers, R., Van Der Greef, J., and Timmerman, M.(2005). ANOVA-Simultaneous Component Analysis (ASCA): A new tool for analyzing designed metabolomics data. Bioinformatics, 21(13), 3043–3048.

- Liland, K.H., Smilde, A., Marini, F., and Næs, T. (2018). Confidence ellipsoids for ASCA models based on multivariate regression theory. Journal of Chemometrics, 32(e2990), 1–13.
- Martin, M. and Govaerts, B. (2020). LiMM-PCA: Combining ASCA+ and linear mixed models to analyse high-dimensional designed data. Journal of Chemometrics, 34(6), e3232.

See Also

Overviews of available methods, multiblock, and methods organised by main structure: basic, unsupervised, asca, supervised and complex. Common functions for computation and extraction of results and plotting are found in asca_results and asca_plots, respectively.

Examples

```
# Load candies data
data(candies)
# Basic ASCA model with two factors
mod <- asca(assessment ~ candy + assessor, data=candies)</pre>
print(mod)
# ASCA model with interaction
mod <- asca(assessment ~ candy * assessor, data=candies)</pre>
print(mod)
# Result plotting for first factor
loadingplot(mod, scatter=TRUE, labels="names")
scoreplot(mod, ellipsoids = "confidence")
# ASCA model with compressed response using 5 principal components
mod.pca <- asca(assessment ~ candy + assessor, data=candies, pca.in=5)</pre>
# Mixed Model ASCA, random assessor
mod.mix <- asca(assessment ~ candy + (1|assessor), data=candies)</pre>
scoreplot(mod.mix)
```

asca_plots

ASCA Result Methods

Description

Various plotting procedures for asca objects.

asca_plots 5

Usage

```
## S3 method for class 'asca'
loadingplot(object, factor = 1, comps = 1:2, ...)
## S3 method for class 'asca'
scoreplot(
  object,
  factor = 1,
  comps = 1:2,
  pch.scores = 19,
  pch.projections = 1,
 gr.col = 1:nlevels(object$effects[[factor]]),
 ellipsoids,
  confidence,
  xlim,
 ylim,
 xlab,
 ylab,
  legendpos,
)
```

Arguments

```
object
                  asca object.
factor
                  integer/character for selecting a model factor.
                  integer vector of selected components.
comps
                  additional arguments to underlying methods.
. . .
                  integer plotting symbol.
pch.scores
pch.projections
                  integer plotting symbol.
                  integer vector of colours for groups.
gr.col
ellipsoids
                  character "confidence" or "data" ellipsoids for balanced fixed effect models.
confidence
                  numeric vector of ellipsoid confidences, default = c(0.4, 0.68, 0.95).
xlim
                  numeric x limits.
                  numeric y limits.
ylim
xlab
                  character x label.
ylab
                  character y label.
                  character position of legend.
legendpos
```

Details

Usage of the functions are shown using generics in the examples in asca. Plot routines are available as scoreplot.asca and loadingplot.asca.

6 asca_results

Value

The plotting routines have no return.

References

- Smilde, A., Jansen, J., Hoefsloot, H., Lamers, R., Van Der Greef, J., and Timmerman, M.(2005). ANOVA-Simultaneous Component Analysis (ASCA): A new tool for analyzing designed metabolomics data. Bioinformatics, 21(13), 3043–3048.
- Liland, K.H., Smilde, A., Marini, F., and Næs, T. (2018). Confidence ellipsoids for ASCA models based on multivariate regression theory. Journal of Chemometrics, 32(e2990), 1–13.
- Martin, M. and Govaerts, B. (2020). LiMM-PCA: Combining ASCA+ and linear mixed models to analyse high-dimensional designed data. Journal of Chemometrics, 34(6), e3232.

See Also

Overviews of available methods, multiblock, and methods organised by main structure: basic, unsupervised, asca, supervised and complex. Common functions for computation and extraction of results are found in asca_results.

asca_results

ASCA Result Methods

Description

Standard result computation and extraction functions for ASCA (asca).

```
## S3 method for class 'asca'
print(x, ...)

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

## S3 method for class 'summary.asca'
print(x, digits = 2, ...)

## S3 method for class 'asca'
loadings(object, factor = 1, ...)

## S3 method for class 'asca'
scores(object, factor = 1, ...)

## S3 method for class 'asca'
projections(object, ...)
```

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Arguments

x asca object.

. . . additional arguments to underlying methods.

object asca object.

digits integer number of digits for printing.

factor integer/character for selecting a model factor.

Details

Usage of the functions are shown using generics in the examples in asca. Explained variances are available (block-wise and global) through blockexpl and print.rosaexpl. Object printing and summary are available through: print.asca and summary.asca. Scores and loadings have their own extensions of scores() and loadings() through scores.asca and loadings.asca. Special to ASCA is that scores are on a factor level basis, while back-projected samples have their own function in projections.asca.

Value

Returns depend on method used, e.g. projections.asca returns projected samples, scores.asca return scores, while print and summary methods return the object invisibly.

References

- Smilde, A., Jansen, J., Hoefsloot, H., Lamers, R., Van Der Greef, J., and Timmerman, M.(2005). ANOVA-Simultaneous Component Analysis (ASCA): A new tool for analyzing designed metabolomics data. Bioinformatics, 21(13), 3043–3048.
- Liland, K.H., Smilde, A., Marini, F., and Næs, T. (2018). Confidence ellipsoids for ASCA models based on multivariate regression theory. Journal of Chemometrics, 32(e2990), 1–13.
- Martin, M. and Govaerts, B. (2020). LiMM-PCA: Combining ASCA+ and linear mixed models to analyse high-dimensional designed data. Journal of Chemometrics, 34(6), e3232.

See Also

Overviews of available methods, multiblock, and methods organised by main structure: basic, unsupervised, asca, supervised and complex. Common functions for plotting are found in asca_plots.

basic

Single- and Two-Block Methods

8 block,data.frame

Description

This documentation covers a range of single- and two-block methods. In particular:

```
• PCA - Principal Component Analysis (pca)
```

- PCR Principal Component Regression (pcr)
- PLSR Partial Least Squares Regression (plsr)
- CCA Canonical Correlation Analysis (cca)
- IFA Interbattery Factor Analysis (ifa)
- GSVD Generalized SVD (gsvd)

See Also

Overviews of available methods, multiblock, and methods organised by main structure: basic, unsupervised, asca, supervised and complex.

Examples

```
data(potato)
X <- potato$Chemical
y <- potato$Sensory[,1,drop=FALSE]

pca.pot <- pca(X, ncomp = 2)
pcr.pot <- pcr(y ~ X, ncomp = 2)
pls.pot <- plsr(y ~ X, ncomp = 2)
cca.pot <- cca(potato[1:2])
ifa.pot <- ifa(potato[1:2])
gsvd.pot <- gsvd(lapply(potato[3:4], t))</pre>
```

block.data.frame

Block-wise indexable data.frame

Description

This is a convenience function for making data. frames that are easily indexed on a block-wise basis.

Usage

```
block.data.frame(X, block_inds = NULL, to.matrix = TRUE)
```

Arguments

| Χ | Either a single data. frame to index or a list of matrices/data.frames |
|------------|--|
| block_inds | Named list of indexes if X is a single data.frame, otherwise NULL. |
| to.matrix | logical indicating if input list elements should be converted to matrices. |

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Value

A data. frame which can be indexed block-wise.

Examples

```
# Random data
M <- matrix(rnorm(200), nrow = 10)
# .. with dimnames
dimnames(M) <- list(LETTERS[1:10], as.character(1:20))
# A named list for indexing
inds <- list(B1 = 1:10, B2 = 11:20)
X <- block.data.frame(M, inds)
str(X)</pre>
```

candies

Sensory assessment of candies.

Description

A dataset containing 9 sensory attributes for 5 candies assessed by 11 trained assessors.

Usage

```
data(candies)
```

Format

A data.frame having 165 rows and 3 variables:

```
assessment Matrix of sensory attributesassessor Factor of assessorscandy Factor of candies
```

References

Luciano G, Næs T. Interpreting sensory data by combining principal component analysis and analysis of variance. Food Qual Prefer. 2009;20(3):167-175.

10 cca

cca

Canonical Correlation Analysis - CCA

Description

This is a wrapper for the stats::cancor function for computing CCA.

Usage

cca(X)

Arguments

Χ

list of input data blocks.

Details

CCA is a method which maximises correlation between linear combinations of the columns of two blocks, i.e. $max(cor(X1 \ x \ a, X2 \ x \ b))$. This is done sequentially with deflation in between, such that a sequence of correlations and weight vectors a and b are associated with a pair of matrices.

Value

multiblock object with associated with printing, scores, loadings. Relevant plotting functions: multiblock_plots and result functions: multiblock_results.

References

Hotelling, H. (1936) Relations between two sets of variates. Biometrika, 28, 321–377.

See Also

Overviews of available methods, multiblock, and methods organised by main structure: basic, unsupervised, asca, supervised and complex. Common functions for computation and extraction of results and plotting are found in multiblock_results and multiblock_plots, respectively.

Examples

```
data(potato)
X <- potato$Chemical
cca.pot <- cca(potato[1:2])</pre>
```

complex 11

Methods With Complex Linkage

Description

This documentation covers a few complex methods. In particular:

- L-PLS Partial Least Squares in L configuration (lpls)
- SO-PLS-PM Sequential and Orthogonalised PLS Path Modeling (sopls_pm)

See Also

Overviews of available methods, multiblock, and methods organised by main structure: basic, unsupervised, asca, supervised and complex.

Examples

```
# L-PLS
sim <- lplsData(I = 30, N = 20, J = 5, K = 6, ncomp = 2)
X1 <- sim$X1; X2 <- sim$X2; X3 <- sim$X3
lp <- lpls(X1,X2,X3) # exo-L-PLS</pre>
```

compnames

Vector of component names

Description

Convenience function for creating a vector of component names based on the dimensions the input object (matrix or object having a score function).

Usage

```
compnames(object, comps, explvar = FALSE, ...)
```

Arguments

| object | An object fitted using the multiblock package. |
|---------|---|
| comps | integer vector of components. |
| explvar | logical indicating if explained variances should be included. |
| | Unused |

Details

This is a copy of compnames from the pls package to work with multiblock objects.

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Value

A character vector of component names.

disco

Distinctive and Common Components with SCA - DISCO

Description

This is a wrapper for the DISCOsca function by Zhengguo Gu for computing DISCO.

Usage

```
disco(X, ncomp = 2, ...)
```

Arguments

X list of input blocks.

ncomp integer number of components to extract.

... additional arguments (not used).

Details

DISCO is a restriction of SCA where Alternating Least Squares is used for estimation of loadings and scores. The SCA solution is rotated towards loadings (in sample linked mode) which are filled with zeros in a pattern resembling distinct, local and common components. When used in sample linked mode and only selecting distinct components, it shares a resemblance to SO-PLS, only in an unsupervised setting. Explained variances are computed as proportion of block variation explained by scores*loadings'.

Value

multiblock object including relevant scores and loadings. Relevant plotting functions: multiblock_plots and result functions: multiblock_results.

References

Schouteden, M., Van Deun, K., Wilderjans, T. F., & Van Mechelen, I. (2014). Performing DISCO-SCA to search for distinctive and common information in linked data. Behavior research methods, 46(2), 576-587.

See Also

Overviews of available methods, multiblock, and methods organised by main structure: basic, unsupervised, asca, supervised and complex.

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Examples

```
data(potato)
potList <- as.list(potato[c(1,2,9)])
pot.disco <- disco(potList)
plot(scores(pot.disco), labels="names")</pre>
```

DISCOsca

DISCO-SCA rotation.

Description

A DISCO-SCA procedure for identifying common and distinctive components. The code is adapted from the orphaned RegularizedSCA package by Zhengguo Gu.

Usage

```
DISCOsca(DATA, R, Jk)
```

Arguments

DATA A matrix, which contains the concatenated data with the same subjects from

multiple blocks. Note that each row represents a subject.

R Number of components $(R \ge 2)$.

Jk A vector containing number of variables in the concatenated data matrix.

Value

Trot_best Estimated component score matrix (i.e., T)

Prot_best Estimated component loading matrix (i.e., P)

comdist A matrix representing common distinctive components. (Rows are data blocks

and columns are components.) 0 in the matrix indicating that the corresponding component of that block is estimated to be zeros, and 1 indicates that (at least one component loading in) the corresponding component of that block is not zero. Thus, if a column in the comdist matrix contains only 1's, then this column is a

common component, otherwise distinctive component.

propExp_component

Proportion of variance per component.

References

Schouteden, M., Van Deun, K., Wilderjans, T. F., & Van Mechelen, I. (2014). Performing DISCO-SCA to search for distinctive and common information in linked data. Behavior research methods, 46(2), 576-587.

14 dummycode

Examples

```
## Not run:
DATA1 <- matrix(rnorm(50), nrow=5)
DATA2 <- matrix(rnorm(100), nrow=5)
DATA <- cbind(DATA1, DATA2)
R <- 5
Jk <- c(10, 20)
DISCOsca(DATA, R, Jk)
## End(Not run)</pre>
```

dummycode

Dummy-coding of a single vector

Description

Flexible dummy-coding allowing for all R's built-in types of contrasts and optional dropping of a factor level to reduce rank defficiency probability.

Usage

```
dummycode(Y, contrast = "contr.sum", drop = TRUE)
```

Arguments

Y vector to dummy code.

contrast Contrast type, default = "contr.sum".

drop logical indicating if one level should be dropped (default = TRUE).

Value

matrix made by dummy-coding the input vector.

Examples

```
vec <- c("a","a","b","b","c","c")
dummycode(vec)</pre>
```

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explvar

Explained predictor variance

Description

Extraction and/or computation of explained variances for various object classes in the multiblock package.

Usage

```
explvar(object)
```

Arguments

object

An object fitted using a method from the multiblock package

Value

A vector of component-wise explained variances for predictors.

Examples

extended.model.frame

Extracting the Extended Model Frame from a Formula or Fit

Description

This function attempts to apply model. frame and extend the result with columns of interactions.

Usage

```
extended.model.frame(formula, data, ..., sep = ".")
```

Arguments

```
formula a model formula or terms object or an R object.
data a data.frame, list or environment (see model.frame).
... further arguments to pass to model.frame.
sep separator in contraction of names for interactions (default = ".").
```

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Value

A data.frame that includes everything a model.frame does plus interaction terms.

Examples

gca

Generalized Canonical Analysis - GCA

Description

This is an interface to both SVD-based (default) and RGCCA-based GCA (wrapping the RGCCA: : rgcca function)

Usage

```
gca(X, ncomp = "max", svd = TRUE, tol = 10^-12, corrs = TRUE, ...)
```

Arguments

| Χ | list of input blocks. |
|-------|--|
| ncomp | integer number of components to extract, either single integer (equal for all blocks), vector (individual per block) or 'max' for maximum possible number of components. |
| svd | logical indicating if Singular Value Decomposition approach should be used (default=TRUE). |
| tol | numeric tolerance for component inclusion (singular values). |
| corrs | logical indicating if correlations should be calculated for RGCCA based approach. |
| | additional arguments for RGCCA approach. |

Details

GCA is a generalisation of Canonical Correlation Analysis to handle three or more blocks. There are several ways to generalise, and two of these are available through gca. The default is an SVD based approach estimating a common subspace and measuring mean squared correlation to this. An alternative approach is available through RGCCA. For the SVD based approach, the ncomp parameter controls the block-wise decomposition while the following the consensus decomposition is limited to the minimum number of components from the individual blocks.

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Value

multiblock object including relevant scores and loadings. Relevant plotting functions: multiblock_plots and result functions: multiblock_results. blockCoef contains canonical coefficients, while blockDecomp contains decompositions of each block.

References

- Carroll, J. D. (1968). Generalization of canonical correlation analysis to three or more sets of variables. Proceedings of the American Psychological Association, pages 227-22.
- Van der Burg, E. and Dijksterhuis, G. (1996). Generalised canonical analysis of individual sensory profiles and instrument data, Elsevier, pp. 221–258.

See Also

Overviews of available methods, multiblock, and methods organised by main structure: basic, unsupervised, asca, supervised and complex. Common functions for computation and extraction of results and plotting are found in multiblock_results and multiblock_plots, respectively.

Examples

```
data(potato)
potList <- as.list(potato[c(1,2,9)])
pot.gca <- gca(potList)
plot(scores(pot.gca), labels="names")</pre>
```

gpa

Generalized Procrustes Analysis - GPA

Description

This is a wrapper for the FactoMineR::GPA function for computing GPA.

Usage

```
gpa(X, graph = FALSE, ...)
```

Arguments

```
X list of input blocks.graph logical indicating if decomposition should be plotted.... additional arguments for RGCCA approach.
```

18 gsvd

Details

GPA is a generalisation of Procrustes analysis, where one matrix is scaled and rotated to be as similar as possible to another one. Through the generalisation, individual scaling and rotation of each input matrix is performed against a common representation which is estimated in an iterative manner.

Value

multiblock object including relevant scores and loadings. Relevant plotting functions: multiblock_plots and result functions: multiblock_results.

References

Gower, J. C. (1975). Generalized procrustes analysis. Psychometrika. 40: 33–51.

See Also

Overviews of available methods, multiblock, and methods organised by main structure: basic, unsupervised, asca, supervised and complex. Common functions for computation and extraction of results and plotting are found in multiblock_results and multiblock_plots, respectively.

Examples

```
data(potato)
potList <- as.list(potato[c(1,2,9)])
pot.gpa <- gpa(potList)
plot(scores(pot.gpa), labels="names")</pre>
```

gsvd

Generalised Singular Value Decomposition - GSVD

Description

This is a wrapper for the geigen::gsvd function for computing GSVD.

Usage

gsvd(X)

Arguments

Χ

list of input data blocks.

Details

GSVD is a generalisation of SVD to two variable-linked matrices where common loadings and block-wise scores are estimated.

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Value

multiblock object with associated with printing, scores, loadings. Relevant plotting functions: multiblock_plots and result functions: multiblock_results.

References

Van Loan, C. (1976) Generalizing the singular value decomposition. SIAM Journal on Numerical Analysis, 13, 76–83.

See Also

Overviews of available methods, multiblock, and methods organised by main structure: basic, unsupervised, asca, supervised and complex. Common functions for computation and extraction of results and plotting are found in multiblock_results and multiblock_plots, respectively.

Examples

```
data(potato)
X <- potato$Chemical
gsvd.pot <- gsvd(lapply(potato[3:4], t))</pre>
```

hogsvd

Higher Order Generalized SVD - HOGSVD

Description

This is a simple implementation for computing HOGSVD

Usage

hogsvd(X)

Arguments

Χ

list of input blocks.

Details

HOGSVD is a generalisation of SVD to two or more blocks. It finds a common set of loadings across blocks and individual sets of scores per block.

Value

multiblock object including relevant scores and loadings. Relevant plotting functions: multiblock_plots and result functions: multiblock_results.

20 hpca

References

Ponnapalli, S. P., Saunders, M. A., Van Loan, C. F., & Alter, O. (2011). A higher-order generalized singular value decomposition for comparison of global mRNA expression from multiple organisms. PloS one, 6(12), e28072.

See Also

Overviews of available methods, multiblock, and methods organised by main structure: basic, unsupervised, asca, supervised and complex. Common functions for computation and extraction of results and plotting are found in multiblock_results and multiblock_plots, respectively.

Examples

```
data(candies)
candyList <- lapply(1:nlevels(candies$candy),function(x)candies$assessment[candies$candy==x,])
can.hogsvd <- hogsvd(candyList)
scoreplot(can.hogsvd, block=1, labels="names")</pre>
```

hpca

Hierarchical Principal component analysis - HPCA

Description

This is a wrapper for the RGCCA::rgcca function for computing HPCA.

Usage

```
hpca(X, ncomp = 2, scale = FALSE, verbose = FALSE, ...)
```

Arguments

X list of input blocks.

ncomp integer number of components to extract.

scale logical indicating if variables should be scaled.

verbose logical indicating if diagnostic information should be printed.

... additional arguments for RGCCA.

Details

HPCA is a hierarchical PCA analysis which combines two or more blocks into a two-level decomposition with block-wise loadings and scores and superlevel common loadings and scores. The method is closely related to the supervised method MB-PLS in structure.

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Value

multiblock object including relevant scores and loadings. Relevant plotting functions: multiblock_plots and result functions: multiblock_results.

References

Westerhuis, J.A., Kourti, T., and MacGregor, J.F. (1998). Analysis of multiblock and hierarchical PCA and PLS models. Journal of Chemometrics, 12, 301–321.

See Also

Overviews of available methods, multiblock, and methods organised by main structure: basic, unsupervised, asca, supervised and complex. Common functions for computation and extraction of results and plotting are found in multiblock_results and multiblock_plots, respectively.

Examples

```
data(potato)
potList <- as.list(potato[c(1,2,9)])
pot.hpca <- hpca(potList)
plot(scores(pot.hpca), labels="names")</pre>
```

ifa

Inter-battery Factor Analysis - IFA

Description

This is a wrapper for the RGCCA::rgcca function for computing IFA.

Usage

```
ifa(X, ncomp = 1, scale = FALSE, verbose = FALSE, ...)
```

Arguments

X list of input data blocks. ncomp integer number of principa

ncomp integer number of principal components to return.

scale logical indicating if variables should be standardised (default=FALSE).

verbose logical indicating if intermediate results should be printed.

... additional arguments to RGCCA::rgcca.

Details

IFA rotates two matrices to align one or more factors against each other, maximising correlations.

jive jive

Value

multiblock object with associated with printing, scores, loadings. Relevant plotting functions: multiblock_plots and result functions: multiblock_results.

References

Tucker, L. R. (1958). An inter-battery method of factor analysis. Psychometrika, 23(2), 111-136.

See Also

Overviews of available methods, multiblock, and methods organised by main structure: basic, unsupervised, asca, supervised and complex. Common functions for computation and extraction of results and plotting are found in multiblock_results and multiblock_plots, respectively.

Examples

```
data(potato)
X <- potato$Chemical
ifa.pot <- ifa(potato[1:2])</pre>
```

jive

Joint and Individual Variation Explained - JIVE

Description

This is a wrapper for the r.jive::jive function for computing JIVE.

Usage

```
jive(X, ...)
```

Arguments

```
X list of input blocks.... additional arguments for r.jive::jive.
```

Details

Jive performs a decomposition of the variation in two or more blocks into low-dimensional representations of individual and joint variation plus residual variation.

Value

multiblock object including relevant scores and loadings. Relevant plotting functions: multiblock_plots and result functions: multiblock_results.

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References

Lock, E., Hoadley, K., Marron, J., and Nobel, A. (2013) Joint and individual variation explained (JIVE) for integrated analysis of multiple data types. Ann Appl Stat, 7 (1), 523–542.

See Also

Overviews of available methods, multiblock, and methods organised by main structure: basic, unsupervised, asca, supervised and complex.

Examples

```
# Too time consuming for testing
data(candies)
candyList <- lapply(1:nlevels(candies$candy),function(x)candies$assessment[candies$candy==x,])
can.jive <- jive(candyList)
summary(can.jive)</pre>
```

lpls

L-PLS regression

Description

Simultaneous decomposition of three blocks connected in an L pattern.

```
lpls(
   X1,
   X2,
   X3,
   ncomp = 2,
   doublecenter = TRUE,
   scale = c(FALSE, FALSE, FALSE),
   type = c("exo"),
   impute = FALSE,
   niter = 25,
   subsetX2 = NULL,
   subsetX3 = NULL,
   ...
)
```

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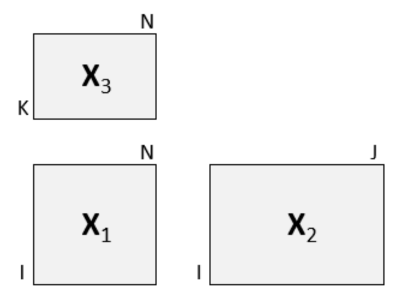
Arguments

| X1 | matrix of size IxN (middle matrix) |
|--------------|---|
| X2 | matrix of size IxJ (left matrix) |
| Х3 | matrix of size KxN (top matrix) |
| ncomp | number of L-PLS components |
| doublecenter | logical indicating if centering should be done both ways for X1 (default=TRUE) |
| scale | logical vector of length three indicating if each of the matrices should be autoscaled. |
| type | character indicating type of L-PLS ("exo"=default, "exo_ort" or "endo") |
| impute | logical indicating if SVD-based imputation of missing data is required. |
| niter | numeric giving number of iterations in component extraction loop. |
| subsetX2 | vector defining optional sub-setting of X2 data. |
| subsetX3 | vector defining optional sub-setting of X3 data. |
| | Additional arguments, not used. |

Details

Two versions of L-PLS are available: exo- and endo-L-PLS which assume an outward or inward relationship between the main block X1 and the two other blocks X2 and X3.

The exo_ort algorithm returns orthogonal scores and should be chosen for visual exploration in correlation loading plots. If exo-L-PLS with prediction is the main purpose of the model then the non-orthogonal exo type L-PLS should be chosen for which the predict function has prediction implemented.



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Value

An object of type lpls and multiblock containing all results from the L-PLS analysis. The object type lpls is associated with functions for correlation loading plots, prediction and cross-validation. The type multiblock is associated with the default functions for result presentation (multiblock_results) and plotting (multiblock_plots).

Author(s)

Solve Sæbø (adapted by Kristian Hovde Liland)

References

- Martens, H., Anderssen, E., Flatberg, A., Gidskehaug, L.H., Høy, M., Westad, F., Thybo, A., and Martens, M. (2005). Regression of a data matrix on descriptors of both its rows and of its columns via latent variables: L-PLSR. Computational Statistics & Data Analysis, 48(1), 103 123.
- Sæbø, S., Almøy, T., Flatberg, A., Aastveit, A.H., and Martens, H. (2008). LPLS-regression: a method for prediction and classification under the influence of background information on predictor variables. Chemometrics and Intelligent Laboratory Systems, 91, 121–132.
- Sæbø, S., Martens, M. and Martens H. (2010) Three-block data modeling by endo- and exo-LPLS regression. In Handbook of Partial Least Squares: Concepts, Methods and Applications. Esposito Vinzi, V.; Chin, W.W.; Henseler, J.; Wang, H. (Eds.). Springer.

See Also

Overviews of available methods, multiblock, and methods organised by main structure: basic, unsupervised, asca, supervised and complex. Functions for computation and extraction of results and plotting are found in lpls_results.

Examples

```
# Simulate data set
sim <- lplsData(I = 30, N = 20, J = 5, K = 6, ncomp = 2)
X1 <- sim$X1; X2 <- sim$X2; X3 <- sim$X3
lp <- lpls(X1,X2,X3) # exo-L-PLS</pre>
```

lplsData

L-PLS data simulation for exo-type analysis

Description

Three data blocks are simulated to express covariance in an exo-L-PLS direction (see lpls. Dimensionality and number of underlying components can be controlled.

```
lplsData(I = 30, N = 20, J = 5, K = 6, ncomp = 2)
```

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Arguments

| I | numeric number of rows of X1 and X2 |
|-------|--|
| N | numeric number of columns in $X1$ and $X3$ |
| J | numeric number of columns in X2 |
| K | numeric number of rows in X3 |
| ncomp | numeric number of latent components |

Value

A list of three matrices with dimensions matching in an L-shape.

Author(s)

Solve Sæbø (adapted by Kristian Hovde Liland)

See Also

Overviews of available methods, multiblock, and methods organised by main structure: basic, unsupervised, asca, supervised and complex.

Examples

```
lp <- lplsData(I = 30, N = 20, J = 5, K = 6, ncomp = 2) names(lp)
```

lpls_results

Result functions for L-PLS objects (lpls)

Description

Correlation loading plot, prediction and cross-validation for L-PLS models with class lpls.

```
## S3 method for class 'lpls'
plot(
    x,
    comps = c(1, 2),
    doplot = c(TRUE, TRUE, TRUE),
    level = c(2, 2, 2),
    arrow = c(1, 0, 1),
    xlim = c(-1, 1),
    ylim = c(-1, 1),
    samplecol = 4,
    pathcol = 2,
```

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```
varcol = "grey70",
      varsize = 1,
      sampleindex = 1:dim(x$corloadings$R22)[1],
      pathindex = 1:dim(x$corloadings$R3)[1],
      varindex = 1:dim(x$corloadings$R21)[1],
    )
    ## S3 method for class 'lpls'
    predict(
      object,
      X1new = NULL,
      X2new = NULL,
      X3new = NULL,
      exo.direction = c("X2", "X3"),
    )
    lplsCV(object, segments1 = NULL, segments2 = NULL, trace = TRUE)
Arguments
                     lpls object
    Х
                     integer vector of components.
    comps
    doplot
                     logical indicating if plotting should be performed.
    level
                     integer vector of length 3 for selecting plot symbol. 1=dots. 2=dimnames.
                     integer vector of length 3 indicating arrows (1) or not (0).
    arrow
    xlim
                     numeric x limits.
    ylim
                     numeric y limits.
    samplecol
                     character for sample colours.
    pathcol
                     character for third colour.
                     character for variable colours.
    varcol
                     numeric size of symbols for variables.
    varsize
    sampleindex
                     integer for selecting samples.
    pathindex
                     integer for selecting in third direction.
    varindex
                     integer for selecting variables.
                     Not implemented.
    . . .
                     lpls object.
    object
    X1new
                     matrix of new X1 samples.
                     matrix of new X2 samples.
    X2new
    X3new
                     matrix of new X3 samples.
                     character selecting "X2" or "X3" prediction.
    exo.direction
    segments1
                     list of sample segments.
    segments2
                     list of variable segments.
                     logical indicating if verbose mode should be selected.
    trace
```

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Value

Nothing is return for plotting (plot.lpls), predicted values are returned for predictions (predict.lpls) and cross-validation metrics are returned for for cross-validation (lplsCV).

See Also

Overviews of available methods, multiblock, and methods organised by main structure: basic, unsupervised, asca, supervised and complex.

Examples

```
# Simulate data set
sim < -lplsData(I = 30, N = 20, J = 5, K = 6, ncomp = 2)
X1 <- sim$X1; X2 <- sim$X2; X3 <- sim$X3
# exo-L-PLS:
lp.exo <- lpls(X1,X2,X3, ncomp = 2)
# Predict X1
pred.exo.X2 <- predict(lp.exo, X1new = X1, exo.direction = "X2")</pre>
# Predict X3
pred.exo.X2 <- predict(lp.exo, X1new = X1, exo.direction = "X3")</pre>
# endo-L-PLS:
lp.endo \leftarrow lpls(X1, X2, X3, ncomp = 2, type = "endo")
# Predict X1 from X2 and X3 (in this case fitted values):
pred.endo.X1 <- predict(lp.endo, X2new = X2, X3new = X3)</pre>
# LOO cross-validation horizontally
lp.cv1 <- lplsCV(lp.exo, segments1 = as.list(1:dim(X1)[1]))</pre>
# LOO cross-validation vertically
lp.cv2 <- lplsCV(lp.exo, segments2 = as.list(1:dim(X1)[2]))</pre>
# Three-fold CV, horizontal
lp.cv3 <- lplsCV(lp.exo, segments1 = as.list(1:10, 11:20, 21:30))</pre>
```

maage

Måge plot

Description

Måge plot for SO-PLS (sopls) cross-validation visualisation.

```
maage(
  object,
  expl_var = TRUE,
```

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```
pure.trace = FALSE,
 pch = 20,
 xlab = "# components",
 ylab = ifelse(expl_var, "Explained variance (%)", "RMSECV"),
 xlim = NULL,
 ylim = NULL,
 cex.text = 0.8,
)
maageSeq(
 object,
 compSeq = TRUE,
 expl_var = TRUE,
 pch = 20,
 xlab = "# components",
 ylab = ifelse(expl_var, "Explained variance (%)", "RMSECV"),
 xlim = NULL,
 ylim = NULL,
 cex.text = 0.8,
 col = "gray",
 col.block = c("red", "blue", "darkgreen", "purple", "black", "red", "blue",
    "darkgreen"),
)
```

Arguments

| object | An SO-PLS model (sop1s object) |
|------------|--|
| expl_var | Logical indicating if explained variance (default) or RMSECV should be displayed. |
| pure.trace | Logical indicating if single block solutions should be traced in the plot. |
| pch | Scalar or symbol giving plot symbol. |
| xlab | Label for x-axis. |
| ylab | Label for y-axis. |
| xlim | Plot limits for x-axis (numeric vector). |
| ylim | Plot limits for y-axis (numeric vector). |
| cex.text | Text scaling (scalar) for better readability of plots. |
| | Additional arguments to plot. |
| compSeq | Integer vector giving the sequence of previous components chosen for maageSeq (see example). |
| col | Line colour in plot. |
| col.block | Line colours for blocks (default = c('red','blue','darkgreen','purple','black')) |

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Details

This function can either be used for global optimisation across blocks or sequential optimisation, using maageSeq. The examples below show typical usage.

Value

The maage plot has no return.

See Also

Overviews of available methods, multiblock, and methods organised by main structure: basic, unsupervised, asca, supervised and complex.

Examples

mbpls

Multiblock Partial Least Squares - MB-PLS

Description

A function computing MB-PLS scores, loadings, etc. on the super-level and block-level.

```
mbpls(
  formula,
  data,
  subset,
  na.action,
  X = NULL,
  Y = NULL,
  ncomp = 1,
  scale = FALSE,
  blockScale = c("sqrtnvar", "ssq", "none"),
  ...
)
```

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Arguments

formula Model formula accepting a single response (block) and predictor block names

separated by + signs.

data The data set to analyse.

subset Expression for subsetting the data before modelling.

na.action How to handle NAs (no action implemented).

X list of input blocks. If X is supplied, the formula interface is skipped.

Y matrix of responses.

ncomp integer number of PLS components.

scale logical for autoscaling inputs (default = FALSE).

blockScale Either a character indicating type of block scaling or a numeric vector of

block weights (see Details).

. . . additional arguments to pls::plsr.

Details

MB-PLS is the prototypical component based supervised multiblock method. It was originally formulated as a two-level method with a block-level and a super-level, but it was later discovered that it could be expressed as an ordinary PLS on concatenated weighted X blocks followed by a simple loop for calculating block-level loading weights, loadings and scores. This implementation uses the plsr function on the scaled input blocks (1/sqrt(ncol)) enabling all summaries and plots from the pls package.

Block weighting is performed after scaling all variables and is by default "sqrtnvar": 1/sqrt(ncol(X[[i]])) in each block. Alternatives are "ssq": $1/norm(X[[i]], "F")^2$ and "none": 1/1. Finally, if a numeric vector is supplied, it will be used to scale the blocks after "ssq" scaling, i.e., $Z[[i]] = X[[i]] / norm(X[[i]], "F")^2 * blockScale[i].$

Value

multiblock, mvr object with super-scores, super-loadings, block-scores and block-loading, and the underlying mvr (PLS) object for the super model, with all its result and plot possibilities. Relevant plotting functions: multiblock_plots and result functions: multiblock_results.

References

- Wangen, L.E. and Kowalski, B.R. (1988). A multiblock partial least squares algorithm for investigating complex chemical systems. Journal of Chemometrics, 3, 3–20.
- Westerhuis, J.A., Kourti, T., and MacGregor, J.F. (1998). Analysis of multiblock and hierarchical PCA and PLS models. Journal of Chemometrics, 12, 301–321.

See Also

Overviews of available methods, multiblock, and methods organised by main structure: basic, unsupervised, asca, supervised and complex.

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Examples

```
data(potato)
# Formula interface
mb <- mbpls(Sensory ~ Chemical+Compression, data=potato, ncomp = 5)
# ... or X and Y
mb.XY <- mbpls(X=potato[c('Chemical','Compression')], Y=potato[['Sensory']], ncomp = 5)
identical(mb$scores, mb.XY$scores)
print(mb)
scoreplot(mb, labels="names") # Exploiting mvr object structure from pls package
# Block scaling with emphasis on first block
mbs <- mbpls(Sensory ~ Chemical+Compression, data=potato, ncomp = 5, blockScale = c(10, 1))
scoreplot(mbs, labels="names") # Exploiting mvr object structure from pls package</pre>
```

mbrda

Multiblock Redundancy Analysis - mbRDA

Description

This is a wrapper for the ade4::mbpcaiv function for computing mbRDA.

Usage

```
mbrda(formula, data, subset, na.action, X = NULL, Y = NULL, ncomp = 1, ...)
```

Arguments

formula Model formula accepting a single response (block) and predictor block names

separated by + signs.

data The data set to analyse.

subset Expression for subsetting the data before modelling.

na.action How to handle NAs (no action implemented).

X list of input blocks. Y matrix of responses.

ncomp integer number of PLS components.
... additional arguments to ade4::mbpcaiv.

Details

mbRDA is a multiblock formulation of Redundancy (Data) Analysis. RDA is theoretically between PLS and GCA. Like GCA, RDA does not consider correlations within X, but like PLS it does consider correlations within Y. RDA can also be viewed as a PCR of Y constrained to have scores that are also linear combinations of X. If the adegraphics package is attached, a nice overview can be plotted as plot(mbr\$mbpcaiv) following the example below.

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Value

multiblock, mvr object with scores, block-scores and block-loading. Relevant plotting functions: multiblock_plots and result functions: multiblock_results.

References

Bougeard, S., Qannari, E.M., Lupo, C., and Hanafi, M. (2011). From Multiblock Partial Least Squares to Multiblock Redundancy Analysis. A Continuum Approach. Informatica, 22(1), 11–26.

See Also

Overviews of available methods, multiblock, and methods organised by main structure: basic, unsupervised, asca, supervised and complex.

Examples

mcoa

Multiple Co-Inertia Analysis - MCOA

Description

This is a wrapper for the RGCCA::rgcca function for computing MCOA.

Usage

```
mcoa(X, ncomp = 2, scale = FALSE, verbose = FALSE, ...)
```

Arguments

| Χ | list of input blocks. |
|---------|---|
| ncomp | integer number of components to extract. |
| scale | logical indicating if variables should be scaled. |
| verbose | logical indicating if diagnostic information should be printed. |
| | additional arguments for RGCCA. |

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Details

MCOA resembles GCA and MFA in that it creates a set of reference scores, for which each block's individual scores should correlate maximally too, but also the variance within each block should be taken into account. A single component solution is equivalent to a PCA on concatenated blocks scaled by the so called inverse inertia.

Value

multiblock object including relevant scores and loadings. Relevant plotting functions: multiblock_plots and result functions: multiblock_results.

References

- Le Roux; B. and H. Rouanet (2004). Geometric Data Analysis, From Correspondence Analysis to Structured Data Analysis. Dordrecht. Kluwer: p.180.
- Greenacre, Michael and Blasius, Jörg (editors) (2006). Multiple Correspondence Analysis and Related Methods. London: Chapman & Hall/CRC.

See Also

Overviews of available methods, multiblock, and methods organised by main structure: basic, unsupervised, asca, supervised and complex. Common functions for computation and extraction of results and plotting are found in multiblock_results and multiblock_plots, respectively.

Examples

```
data(potato)
potList <- as.list(potato[c(1,2,9)])
pot.mcoa <- mcoa(potList)
plot(scores(pot.mcoa), labels="names")</pre>
```

mcolors

Colour palette generation from matrix of RGB values

Description

Colour palette generation from matrix of RGB values

```
mcolors(
   n,
   colmatrix = matrix(c(0, 0, 1, 1, 1, 1, 0, 0), 3, 3, byrow = TRUE)
)
```

mfa 35

Arguments

n Integer number of colorus to produce.

colmatrix A numeric matrix of three columns (R,G,B) to generate colour palette from.

Value

A vector of n colours in hexadecimal RGB format.

Examples

```
mcolors(5)
```

mfa

Multiple Factor Analysis - MFA

Description

This is a wrapper for the FactoMineR::MFA function for computing MFA.

Usage

```
mfa(X, type = rep("c", length(X)), graph = FALSE, ...)
```

Arguments

X list of input blocks.

type character vector indicating block types, defaults to rep("c", length(X)) for

continuous values.

graph logical indicating if decomposition should be plotted.

... additional arguments for RGCCA approach.

Details

MFA is a methods typically used to compare several equally sized matrices. It is often used in sensory analyses, where matrices consist of sensory characteristics and products, and each assessor generates one matrix each. In its basic form, MFA scales all matrices by their largest eigenvalue, concatenates them and performs PCA on the result. There are several possibilities for plots and inspections of the model, handling of categorical and continuous inputs etc. connected to MFA.

Value

multiblock object including relevant scores and loadings. Relevant plotting functions: multiblock_plots and result functions: multiblock_results.

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References

Pagès, J. (2005). Collection and analysis of perceived product inter-distances using multiple factor analysis: Application to the study of 10 white wines from the Loire valley. Food Quality and Preference, 16(7), 642–649.

See Also

Overviews of available methods, multiblock, and methods organised by main structure: basic, unsupervised, asca, supervised and complex. Common functions for computation and extraction of results and plotting are found in multiblock_results and multiblock_plots, respectively.

Examples

```
data(potato)
potList <- as.list(potato[c(1,2,9)])
pot.mfa <- mfa(potList)
if(interactive()){
  plot(pot.mfa$MFA)
}</pre>
```

multiblock_plots

Plot Functions for Multiblock Objects

Description

Plotting procedures for multiblock objects.

```
## S3 method for class 'multiblock'
scoreplot(
   object,
   comps = 1:2,
   block = 0,
   labels,
   identify = FALSE,
   type = "p",
   xlab,
   ylab,
   main,
   ...
)

## S3 method for class 'multiblock'
loadingplot(
```

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```
object,
  comps = 1:2,
  block = 0,
  scatter = TRUE,
  labels,
  identify = FALSE,
  type,
  lty,
  lwd = NULL,
  pch,
  cex = NULL,
  col,
  legendpos,
 xlab,
 ylab,
 main,
 pretty.xlabels = TRUE,
  xlim,
  . . .
)
loadingweightplot(object, main = "Loading weights", ...)
## S3 method for class 'multiblock'
biplot(
 Х,
 block = 0,
  comps = 1:2,
 which = c("x", "y", "scores", "loadings"),
 var.axes = FALSE,
  xlabs,
 ylabs,
 main,
)
corrplot(object, ...)
## Default S3 method:
corrplot(object, ...)
## S3 method for class 'mvr'
corrplot(object, ...)
## S3 method for class 'multiblock'
corrplot(
 object,
  comps = 1:2,
```

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```
labels = TRUE,
col = 1:5,
plotx = TRUE,
ploty = TRUE,
blockScores = FALSE,
...
)
```

Arguments

object multiblock object.

comps integer vector giving components, within block, to plot.

block integer/character for block selection.

labels character indicating if "names" or "numbers" should be plot symbols (op-

tional).

identify logical for activating identify to interactively identify points.

type character for selecting type of plot to make. Defaults to "p" (points) for scatter

plots and "l" (lines) for line plots.

xlab character text for x labels.
ylab character text for y labels.
main character text for main header.

... Not implemented.

scatter logical indicating if a scatterplot of loadings should be made (default = TRUE).

1ty Vector of line type specifications (see par for details).

lwd numeric vector of line width specifications.

pch Vector of point specifications (see points for details).

cex numeric vector of plot size expansions (see par for details).
col integer vector of symbol/line colours (see par for details).

legendpos character indicating legend position (if scatter is FALSE), e.g. legendpos

= "topright".

pretty.xlabels logical indicating if xlabels should be more nicely plotted (default = TRUE).

xlim numeric vector of length two, with the x limits of the plot (optional).

x multiblock object.

which character for selecting type of biplot ("x" = default, "y", "scores", "loadings").

var.axes logical indicating if second axes of a biplot should have arrows.

xlabs character vector for labelling first set of biplot points (optional).

ylabs character vector for labelling second set of biplot points (optional).

plotx locical or integer/character. Whether to plot the X correlation loadings,

optionally which block(s). Defaults to TRUE.

ploty logical. Whether to plot the Y correlation loadings. Defaults to TRUE. blockScores logical. Correlation loadings from blockScores (default = FALSE).

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Details

Plot functions for scores, loadings and loading.weights based on the functions found in the pls package.

Value

These plotting routines only generate plots and return no values.

See Also

Overviews of available methods, multiblock, and methods organised by main structure: basic, unsupervised, asca, supervised and complex. Common functions for computation and extraction of results are found in multiblock_results.

Examples

multiblock_results

Result Functions for Multiblock Objects

Description

Standard result computation and extraction functions for multiblock objects.

```
## S3 method for class 'multiblock'
scores(object, block = 0, ...)

## S3 method for class 'multiblock'
loadings(object, block = 0, ...)

## S3 method for class 'multiblock'
print(x, ...)

## S3 method for class 'multiblock'
summary(object, ...)
```

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Arguments

```
object multiblock object.
block integer/character for block selection.
... Not implemented.
x multiblock object.
```

Details

Usage of the functions are shown using generics in the examples below. Object printing and summary are available through: print.multiblock and summary.multiblock. Scores and loadings have their own extensions of scores() and loadings() throught scores.multiblock and loadings.multiblock.

Value

Scores or loadings are returned by scores.multiblock and loadings.multiblock, while print and summary methods invisibly returns the object.

See Also

Overviews of available methods, multiblock, and methods organised by main structure: basic, unsupervised, asca, supervised and complex. Common functions for plotting are found in multiblock_plots, respectively.

Examples

```
data(wine)
sc <- sca(wine[c('Smell at rest', 'View', 'Smell after shaking')], ncomp = 4)
print(sc)
summary(sc)
head(loadings(sc, block = 1))
head(scores(sc))</pre>
```

mvrVal

MSEP, RMSEP and R2 of the MB-PLS model

Description

Functions to estimate the mean squared error of prediction (MSEP), root mean squared error of prediction (RMSEP) and \mathbb{R}^2 (A.K.A. coefficient of multiple determination) for a fitted MB-PLS models. Test-set, cross-validation and calibration-set estimates are implemented.

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Usage

```
## S3 method for class 'mbpls'
R2(
  object,
  estimate,
  newdata,
  ncomp = 1:object$ncomp,
  comps,
  intercept = TRUE,
  se = FALSE,
)
## S3 method for class 'mbpls'
MSEP(
  object,
  estimate,
  newdata,
  ncomp = 1:object$ncomp,
  comps,
  intercept = TRUE,
  se = FALSE,
)
## S3 method for class 'mbpls'
RMSEP(object, ...)
```

Arguments

an mvr object object estimate a character vector. Which estimators to use. Should be a subset of c("all", "train", "CV", "adjCV", "test"). "adjCV" is only available for (R)MSEP. See below for how the estimators are chosen. newdata a data frame with test set data. a vector of positive integers. The components or number of components to use. ncomp, comps See below. intercept logical. Whether estimates for a model with zero components should be returned as well. logical. Whether estimated standard errors of the estimates should be calculated. se Not implemented yet. further arguments sent to underlying functions or (for RMSEP) to MSEP

Details

RMSEP simply calls MSEP and takes the square root of the estimates. It therefore accepts the same arguments as MSEP.

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Several estimators can be used. "train" is the training or calibration data estimate, also called (R)MSEC. For R2, this is the unadjusted R^2 . It is overoptimistic and should not be used for assessing models. "CV" is the cross-validation estimate, and "adjCV" (for RMSEP and MSEP) is the bias-corrected cross-validation estimate. They can only be calculated if the model has been cross-validated. Finally, "test" is the test set estimate, using newdata as test set.

Which estimators to use is decided as follows (see below for pls:mvrValstats). If estimate is not specified, the test set estimate is returned if newdata is specified, otherwise the CV and adjusted CV (for RMSEP and MSEP) estimates if the model has been cross-validated, otherwise the training data estimate. If estimate is "all", all possible estimates are calculated. Otherwise, the specified estimates are calculated.

Several model sizes can also be specified. If comps is missing (or is NULL), length(ncomp) models are used, with ncomp[1] components, ..., ncomp[length(ncomp)] components. Otherwise, a single model with the components comps[1], ..., comps[length(comps)] is used. If intercept is TRUE, a model with zero components is also used (in addition to the above).

The R^2 values returned by "R2" are calculated as 1-SSE/SST, where SST is the (corrected) total sum of squares of the response, and SSE is the sum of squared errors for either the fitted values (i.e., the residual sum of squares), test set predictions or cross-validated predictions (i.e., the PRESS). For estimate = "train", this is equivalent to the squared correlation between the fitted values and the response. For estimate = "train", the estimate is often called the prediction R^2 .

mvrValstats is a utility function that calculates the statistics needed by MSEP and R2. It is not intended to be used interactively. It accepts the same arguments as MSEP and R2. However, the estimate argument must be specified explicitly: no partial matching and no automatic choice is made. The function simply calculates the types of estimates it knows, and leaves the other untouched.

Value

mvrValstats returns a list with components

SSE three-dimensional array of SSE values. The first dimension is the different estimators, the second is the response variables and the third is the models.

SST matrix of SST values. The first dimension is the different estimators and the second is the response variables.

nobj a numeric vector giving the number of objects used for each estimator.

comps the components specified, with 0 prepended if intercept is TRUE.

cumulative TRUE if comps was NULL or not specified.

The other functions return an object of class "mvrVal", with components

val three-dimensional array of estimates. The first dimension is the different estimators, the second is the response variables and the third is the models.

type "MSEP", "RMSEP" or "R2".

comps the components specified, with 0 prepended if intercept is TRUE.

cumulative TRUE if comps was NULL or not specified.

call the function call

pca 43

Author(s)

Kristian Hovde Liland

References

Mevik, B.-H., Cederkvist, H. R. (2004) Mean Squared Error of Prediction (MSEP) Estimates for Principal Component Regression (PCR) and Partial Least Squares Regression (PLSR). *Journal of Chemometrics*, **18**(9), 422–429.

See Also

```
mbpls
```

Examples

```
data(oliveoil, package = "pls")
mod <- pls::plsr(sensory ~ chemical, ncomp = 4, data = oliveoil, validation = "L00")
RMSEP(mod)
## Not run: plot(R2(mod))</pre>
```

рса

Principal Component Analysis - PCA

Description

This is a wrapper for the pls::PCR function for computing PCA.

Usage

```
pca(X, scale = FALSE, ncomp = 1, ...)
```

Arguments

```
X matrix of input data.

scale logical indicating if variables should be standardised (default=FALSE).

ncomp integer number of principal components to return.
```

... additional arguments to pls:pcr.

Details

PCA is a method for decomposing a matrix into subspace components with sample scores and variable loadings. It can be formulated in various ways, but the standard formulation uses singular value decomposition to create scores and loadings. PCA is guaranteed to be the optimal way of extracting orthogonal subspaces from a matrix with regard to the amount of explained variance per component.

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Value

multiblock object with scores, loadings, mean X values and explained variances. Relevant plotting functions: multiblock_plots and result functions: multiblock_results.

References

Pearson, K. (1901) On lines and planes of closest fit to points in space. Philosophical Magazine, 2, 559–572.

See Also

Overviews of available methods, multiblock, and methods organised by main structure: basic, unsupervised, asca, supervised and complex. Common functions for computation and extraction of results and plotting are found in multiblock_results and multiblock_plots, respectively.

Examples

```
data(potato)
X <- potato$Chemical
pca.pot <- pca(X, ncomp = 2)</pre>
```

pcagca

PCA-GCA

Description

PCA-GCA is a methods which aims at estimating subspaces of common, local and distinct variation from two or more blocks.

```
pcagca(
   X,
   commons = 2,
   auto = TRUE,
   auto.par = list(explVarLim = 40, rLim = 0.8),
   manual.par = list(ncomp = 0, ncommon = 0),
   tol = 10^-12
)
```

pcagca 45

Arguments

X list of input blocks

commons numeric giving the highest number of blocks to combine when calculating local

or common scores.

auto logical indicating if automatic choice of complexities should be used.

auto.par named list setting limits for automatic choice of complexities.

manual.par named list for manual choice of blocks. The list consists of ncomp which indi-

cates the number of components to extract from each block and ncommon which is the corresponding for choosing the block combinations (local/common). For the latter, use unique_combos(n_blocks, commons) to see order of local/common blocks. Component numbers will be reduced if simpler models give better pre-

dictions. See example.

tol numeric tolerance for component inclusion (singular values).

Details

The name PCA-GCA comes from the process of first applying PCA to each block, then using GCA to estimate local and common components, and finally orthogonalising the block-wise scores on the local/common ones and re-estimating these to obtain distinct components. The procedure is highly similar to the supervised method PO-PLS, where the PCA steps are exchanged with PLS.

Value

multiblock object including relevant scores and loadings. Relevant plotting functions: multiblock_plots and result functions: multiblock_results. Distinct components are marked as 'D(x), Comp c' for block x and component c while local and common components are marked as "C(x1, x2), Comp c", where x1 and x2 (and more) are block numbers.

References

Smilde, A., Måge, I., Naes, T., Hankemeier, T.,Lips, M., Kiers, H., Acar, E., and Bro, R.(2017). Common and distinct components in data fusion. Journal of Chemometrics, 31(7), e2900.

See Also

Overviews of available methods, multiblock, and methods organised by main structure: basic, unsupervised, asca, supervised and complex. Common functions for computation and extraction of results and plotting are found in multiblock_results and multiblock_plots, respectively.

Examples

```
data(potato)
potList <- as.list(potato[c(1,2,9)])
pot.pcagca <- pcagca(potList)

# Show origin and type of all components
lapply(pot.pcagca$blockScores,colnames)</pre>
```

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```
# Basic multiblock plot
plot(scores(pot.pcagca, block=2), comps=1, labels="names")
```

popls

Parallel and Orthogonalised Partial Least Squares - PO-PLS

Description

This is a basic implementation of PO-PLS with manual and automatic component selections.

Usage

```
popls(
    X,
    Y,
    commons = 2,
    auto = TRUE,
    auto.par = list(explVarLim = 40, rLim = 0.8),
    manual.par = list(ncomp = rep(0, length(X)), ncommon = list())
)
```

Arguments

X list of input blocks

Y matrix of response variable(s)

commons numeric giving the highest number of blocks to combine when calculating local

or common scores.

auto logical indicating if automatic choice of complexities should be used.

auto.par named list setting limits for automatic choice of complexities. See Details.

manual.par named list for manual choice of blocks. The list consists of ncomp which indi-

cates the number of components to extract from each block and ncommon which is the corresponding for choosing the block combinations (local/common). For the latter, use unique_combos(n_blocks, commons) to see order of local/common blocks. Component numbers will be reduced if simpler models give better pre-

dictions. See example.

Details

PO-PLS decomposes a set of input data blocks into common, local and distinct components through a process involving pls and gca. The rLim parameter is a lower bound for the GCA correlation when building common components, while explVarLim is the minimum explained variance for common components and unique components.

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Value

A multiblock object with block-wise, local and common loadings and scores. Relevant plotting functions: multiblock_plots and result functions: multiblock_results.

References

- I Måge, BH Mevik, T Næs. (2008). Regression models with process variables and parallel blocks of raw material measurements. Journal of Chemometrics: A Journal of the Chemometrics Society 22 (8), 443-456
- I Måge, E Menichelli, T Næs (2012). Preference mapping by PO-PLS: Separating common and unique information in several data blocks. Food quality and preference 24 (1), 8-16

See Also

Overviews of available methods, multiblock, and methods organised by main structure: basic, unsupervised, asca, supervised and complex. Common functions for computation and extraction of results and plotting are found in multiblock_results and multiblock_plots, respectively.

Examples

potato

Sensory, rheological, chemical and spectroscopic analysis of potatoes.

Description

A dataset containing 9 blocks of measurements on 26 potatoes. Original dataset can be found at http://models.life.ku.dk/Texture_Potatoes. This version has been pre-processed as follows (corresponding to Liland et al. 2016):

- Variables containing NaN have been removed.
- Chemical and Compression blocks have been scaled by standard deviations.
- NIR blocks have been subjected to SNV (Standard Normal Variate).

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Usage

```
data(potato)
```

Format

A data.frame having 26 rows and 9 variables:

Chemical Matrix of chemical measurements

Compression Matrix of rheological compression data

NIRraw Matrix of near-infrared measurements of raw potatoes

NIRcooked Matrix of near-infrared measurements of cooked potatoes

CPMGraw Matrix of NMR (CPMG) measurements of raw potatoes

CPMGcooked Matrix of NMR (CPMG) measurements of cooked potatoes

FIDraw Matrix of NMR (FID) measurements of raw potatoes

FIDcooked Matrix of NMR (FID) measurements of cooked potatoes

Sensory Matrix of sensory assessments

References

- L.G.Thygesen, A.K.Thybo, S.B.Engelsen, Prediction of Sensory Texture Quality of Boiled Potatoes From Low-field1H NMR of Raw Potatoes. The Role of Chemical Constituents. LWT Food Science and Technology 34(7), 2001, pp 469-477.
- Kristian Hovde Liland, Tormod Næs, Ulf Geir Indahl, ROSA a fast extension of Partial Least Squares Regression for Multiblock Data Analysis, Journal of Chemometrics 30:11 (2016), pp. 651-662.

predict.mbpls

Predict Method for MBPLS

Description

Prediction for the mbpls (MBPLS) model. New responses or scores are predicted using a fitted model and a data.frame or list containing matrices of observations.

```
## S3 method for class 'mbpls'
predict(
  object,
  newdata,
  ncomp = 1:object$ncomp,
  comps,
  type = c("response", "scores"),
  na.action = na.pass,
  ...
)
```

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Arguments

object an mvr object. The fitted model

newdata a data frame. The new data. If missing, the training data is used.

ncomp, comps vector of positive integers. The components to use in the prediction. See below.

type character. Whether to predict scores or response values

na.action function determining what should be done with missing values in newdata. The

default is to predict NA. See na. omit for alternatives.

... further arguments. Currently not used

Details

When type is "response" (default), predicted response values are returned. If comps is missing (or is NULL), predictions for length(ncomp) models with ncomp[1] components, ncomp[2] components, etc., are returned. Otherwise, predictions for a single model with the exact components in comps are returned. (Note that in both cases, the intercept is always included in the predictions. It can be removed by subtracting the Ymeans component of the fitted model.)

When type is "scores", predicted score values are returned for the components given in comps. If comps is missing or NULL, ncomps is used instead.

Value

When type is "response", a three dimensional array of predicted response values is returned. The dimensions correspond to the observations, the response variables and the model sizes, respectively.

When type is "scores", a score matrix is returned.

Note

A warning message like ''newdata' had 10 rows but variable(s) found have 106 rows' means that not all variables were found in the newdata data frame. This (usually) happens if the formula contains terms like yarn\$NIR. Do not use such terms; use the data argument instead. See mvr for details.

Author(s)

Kristian Hovde Liland

See Also

mbpls

Examples

```
data(potato)
mb <- mbpls(Sensory ~ Chemical+Compression, data=potato, ncomp = 5, subset = 1:26 <= 18)
testdata <- subset(potato, 1:26 > 18)

# Predict response
yhat <- predict(mb, newdata = testdata)</pre>
```

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```
# Predict scores and plot
scores <- predict(mb, newdata = testdata, type = "scores")
scoreplot(mb)
points(scores[,1], scores[,2], col="red")
legend("topright", legend = c("training", "test"), col=1:2, pch = 1)</pre>
```

preprocess

Preprocessing of block data

Description

This is an interface to simplify preprocessing of one, a subset or all blocks in a multiblock object, e.g., a data.frame (see block.data.frame) or list. Several standard preprocessing methods are supplied in addition to letting the user supply it's own function.

Usage

```
block.preprocess(
    X,
    block = 1:length(X),
    fun = c("autoscale", "center", "scale", "SNV", "EMSC", "Fro", "FroSq", "SingVal"),
    ...
)
```

Arguments

```
X data.frame or list of data.

block vector of block(s) to preprocess (integers or characters).

fun character or function selecting which preprocessing to apply (see Details).

additional arguments to underlying functions.
```

Details

The fun parameter controls the type of preprocessing to be performed:

- autoscale: centre and scale each feature/variable.
- center: centre each feature/variable.
- scale: scale each feature/variable.
- SNV: Standard Normal Variate correction, i.e., centre and scale each sample across features/variables.
- EMSC: Extended Multiplicative Signal Correction defaulting to basic EMSC (2nd order polynomials). Further parameters are sent to EMSC: :EMSC.
- Fro: Frobenius norm scaling of whole block.
- FroSq: Squared Frobenius norm scaling of whole block (sum of squared values).

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- SingVal: Singular value scaling of whole block (first singular value).
- User defined: If a function is supplied, this will be applied to chosen blocks. Preprocessing can be done for all blocks or a subset. It can also be done in a series of operations to combine preprocessing techniques.

Value

The input multiblock object is preprocessed and returned.

See Also

Overviews of available methods, multiblock, and methods organised by main structure: basic, unsupervised, asca, supervised and complex. Common functions for computation and extraction of results and plotting are found in multiblock_results and multiblock_plots, respectively.

Examples

```
data(potato)
# Autoscale Chemical block
potato <- block.preprocess(potato, block = "Chemical", "autoscale")</pre>
# Apply SNV to NIR blocks
potato <- block.preprocess(potato, block = 3:4, "SNV")</pre>
# Centre Sensory block
potato <- block.preprocess(potato, block = "Sensory", "center")</pre>
# Scale all blocks to unit Frobenius norm
potato <- block.preprocess(potato, fun = "Fro")</pre>
# Effect of SNV
NIR <- (potato$NIRraw + rnorm(26)) * rnorm(26,1,0.2)
NIRc <- block.preprocess(list(NIR), fun = "SNV")[[1]]</pre>
old.par <- par(mfrow = c(2,1), mar = c(4,4,1,1))
matplot(t(NIR), type="l", main = "uncorrected", ylab = "")
matplot(t(NIRc), type="l", main = "corrected", ylab = "")
par(old.par)
```

rosa

Response Oriented Sequential Alternation - ROSA

Description

Formula based interface to the ROSA algorithm following the style of the pls package.

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Usage

```
rosa(
  formula,
  ncomp,
 Y.add,
  common.comp = 1,
  data,
  subset,
  na.action,
  scale = FALSE,
  weights = NULL,
  validation = c("none", "CV", "L00"),
  internal.validation = FALSE,
  fixed.block = NULL,
  design.block = NULL,
  canonical = TRUE,
)
```

Arguments

formula Model formula accepting a single response (block) and predictor block names

separated by + signs.

ncomp The maximum number of ROSA components.

Y. add Optional response(s) available in the data set.

common.comp Automatically create all combinations of common components up to length

common.comp (default = 1).

data The data set to analyse.

subset Expression for subsetting the data before modelling.

na.action How to handle NAs (no action implemented).

scale Optionally scale predictor variables by their individual standard deviations.

weights Optional object weights.

validation Optional cross-validation strategy "CV" or "LOO".

internal.validation

Optional cross-validation for block selection process, "LOO", "CV3", "CV5", "CV10" (CV-number of segments), or vector of integers (default = FALSE).

(actual These).

fixed.block integer vector with block numbers for each component (0 = not fixed) or list of

length \leq ncomp (element length 0 = not fixed).

design.block integer vector containing block numbers of design blocks

canonical logical indicating if canonical correlation should be use when calculating load-

ing weights (default), enabling B/W maximization, common components, etc. Alternatively (FALSE) a PLS2 strategy, e.g. for spectra response, is used.

.. Additional arguments for cvseg or rosa.fit

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Details

ROSA is an opportunistic method sequentially selecting components from whichever block explains the response most effectively. It can be formulated as a PLS model on concatenated input block with block selection per component. This implementation adds several options that are not described in the literature. Most importantly, it opens for internal validation in the block selection process, making this more robust. In addition it handles design blocks explicitly, enables classification and secondary responses (CPLS), and definition of common components.

Value

An object of classes rosa and mvr having several associated printing (rosa_results) and plotting methods (rosa_plots).

References

Liland, K.H., Næs, T., and Indahl, U.G. (2016). ROSA - a fast extension of partial least squares regression for multiblock data analysis. Journal of Chemometrics, 30, 651–662, doi:10.1002/cem.2824.

See Also

Overviews of available methods, multiblock, and methods organised by main structure: basic, unsupervised, asca, supervised and complex. Common functions for computation and extraction of results and plotting are found in rosa_results and rosa_plots, respectively.

Examples

```
data(potato)
mod <- rosa(Sensory[,1] ~ ., data = potato, ncomp = 10, validation = "CV", segments = 5)
summary(mod)

# For examples of ROSA results and plotting see
# ?rosa_results and ?rosa_plots.</pre>
```

rosa_plots

Plotting functions for ROSA models

Description

Various plotting procedures for rosa objects.

```
## S3 method for class 'rosa'
image(
    x,
    type = c("correlation", "residual", "order"),
    ncomp = x$ncomp,
    col = mcolors(128),
```

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```
legend = TRUE,
mar = c(5, 6, 4, 7),
las = 1,
...
)

## S3 method for class 'rosa'
barplot(
  height,
  type = c("train", "CV"),
  ncomp = height$ncomp,
  col = mcolors(ncomp),
  ...
)
```

Arguments

| X | A rosa object |
|--------|---|
| type | An optional character for selecting the plot type. For image.rosa the options are: "correlation" (default), "residual" or "order". For barplot.rosa the options indicate: explained variance should be based on training data ("train") or cross-validation ("CV"). |
| ncomp | Integer to control the number of components to plot (if fewer than the fitted number of components). |
| col | Colours used for the image and bar plot, defaulting to mcolors(128). |
| legend | Logical indicating if a legend should be included (default = TRUE) for image.rosa. |
| mar | Figure margins, default = $c(5,6,4,7)$ for image.rosa. |
| las | Axis text direction, default = 1 for image.rosa. |
| • • • | Additional parameters passed to loadingplot, image, axis, color.legend, or barplot. |
| height | A rosa object. |

Details

Usage of the functions are shown using generics in the examples below. image.rosa makes an image plot of each candidate score's correlation to the winner or the block-wise response residual. These plots can be used to find alternative block selection for tweaking the ROSA model. barplot.rosa makes barplot of block and component explained variances. loadingweightsplot is an adaptation of pls::loadingplot to plot loading weights.

Value

No return.

References

Liland, K.H., Næs, T., and Indahl, U.G. (2016). ROSA - a fast extension of partial least squares regression for multiblock data analysis. Journal of Chemometrics, 30, 651–662, doi:10.1002/cem.2824.

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See Also

Overviews of available methods, multiblock, and methods organised by main structure: basic, unsupervised, asca, supervised and complex. Common functions for computation and extraction of results in rosa_results.

Examples

```
data(potato)
mod <- rosa(Sensory[,1] ~ ., data = potato, ncomp = 5)
image(mod)
barplot(mod)
loadingweightplot(mod)</pre>
```

rosa_results

Result functions for ROSA models

Description

Standard result computation and extraction functions for ROSA (rosa).

```
## S3 method for class 'rosa'
predict(
  object,
  newdata,
 ncomp = 1:object$ncomp,
  comps,
  type = c("response", "scores"),
  na.action = na.pass,
)
## S3 method for class 'rosa'
coef(object, ncomp = object$ncomp, comps, intercept = FALSE, ...)
## S3 method for class 'rosa'
print(x, ...)
## S3 method for class 'rosa'
summary(
  object,
  what = c("all", "validation", "training"),
  digits = 4,
  print.gap = 2,
```

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```
blockexpl(object, ncomp = object$ncomp, type = c("train", "CV"))
## S3 method for class 'rosaexpl'
print(x, digits = 3, compwise = FALSE, ...)

rosa.classify(object, classes, newdata, ncomp, LQ)
## S3 method for class 'rosa'
scores(object, ...)
## S3 method for class 'rosa'
loadings(object, ...)
```

Arguments

| object | A rosa object. |
|-----------|--|
| newdata | Optional new data with the same types of predictor blocks as the ones used for fitting the object. |
| ncomp | An integer giving the number of components to apply (cumulative). |
| comps | An integer vector giving the exact components to apply (subset). |
| type | For blockexpl: Character indicating which type of explained variance to compute (default = "train", alternative = " CV "). |
| na.action | Function determining what to do with missing values in newdata. |
| | Additional arguments. Currently not implemented. |
| intercept | A logical indicating if coefficients for the intercept should be included (default $=$ FALSE). |
| x | A rosa object. |
| what | A character indicating if summary should include all, validation or training. |
| digits | The number of digits used for printing. |
| print.gap | Gap between columns when printing. |
| compwise | Logical indicating if block-wise (default/FALSE) or component-wise (TRUE) explained variance should be printed. |
| classes | A character vector of class labels. |
| LQ | A character indicating if 'max' (maximum score value), 'lda' or 'qda' should be used when classifying. |
| | |

Details

Usage of the functions are shown using generics in the examples below. Prediction, regression coefficients, object printing and summary are available through: predict.rosa, coef.rosa, print.rosa and summary.rosa. Explained variances are available (block-wise and global) through blockexpl and print.rosaexpl. Scores and loadings have their own extensions of scores() and loadings()

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throught scores.rosa and loadings.rosa. Finally, there is work in progress on classification support through rosa.classify.

When type is "response" (default), predicted response values are returned. If comps is missing (or is NULL), predictions for length(ncomp) models with ncomp[1] components, ncomp[2] components, etc., are returned. Otherwise, predictions for a single model with the exact components in comps are returned. (Note that in both cases, the intercept is always included in the predictions. It can be removed by subtracting the Ymeans component of the fitted model.)

If comps is missing (or is NULL), coef()[, ncomp[i]] are the coefficients for models with ncomp[i] components, for $i=1,\ldots,length(ncomp)$. Also, if intercept = TRUE, the first dimension is nxvar+1, with the intercept coefficients as the first row.

If comps is given, however, coef()[,,comps[i]] are the coefficients for a model with only the component comps[i], i.e., the contribution of the component comps[i] on the regression coefficients.

Value

Returns depend on method used, e.g. predict.rosa returns predicted responses or scores depending on inputs, coef.rosa returns regression coefficients, blockexpl returns an object of class rosaexpl containing block-wise and component-wise explained variance contained in a matrix with attributes.

References

Liland, K.H., Næs, T., and Indahl, U.G. (2016). ROSA - a fast extension of partial least squares regression for multiblock data analysis. Journal of Chemometrics, 30, 651–662, doi:10.1002/cem.2824.

See Also

Overviews of available methods, multiblock, and methods organised by main structure: basic, unsupervised, asca, supervised and complex. Common functions for computation and extraction of results and plotting are found in rosa_results and rosa_plots, respectively.

Examples

```
data(potato)
mod <- rosa(Sensory[,1] ~ ., data = potato, ncomp = 5, subset = 1:20)
testset <- potato[-(1:20),]; testset$Sensory <- testset$Sensory[,1,drop=FALSE]
predict(mod, testset, ncomp=5)
dim(coef(mod, ncomp=5)) # <variables x responses x components>
print(mod)
summary(mod)
blockexpl(mod)
print(blockexpl(mod), compwise=TRUE)
```

58 sca

| sca | Simultaneous C | Component Analysis | - SCA |
|-----|----------------|--------------------|-------|
| | | | |

Description

This is a basic implementation of the SCA-P algorithm (least restricted SCA) with support for both sample- and variable-linked modes.

Usage

```
sca(X, ncomp = 2, scale = FALSE, samplelinked = "auto", ...)
```

Arguments

X list of input blocks.

ncomp integer number of components to extract.

scale logical indicating autoscaling of features (default = FALSE).

samplelinked character/logical indicating if blocks are linked by samples (TRUE) or vari-

ables (FALSE). Using 'auto' (default), this will be determined automatically.

... additional arguments (not used).

Details

SCA, in its original variable-linked version, calculates common loadings and block-wise scores. There are many possible constraints and specialisations. This implementations uses PCA as the backbone, thus resulting in deterministic, ordered components. A parameter controls the linking mode, but if left untouched an attempt is made at automatically determining variable or sample linking.

Value

multiblock object including relevant scores and loadings. Relevant plotting functions: multiblock_plots and result functions: multiblock_results.

References

Levin, J. (1966) Simultaneous factor analysis of several gramian matrices. Psychometrika, 31(3), 413–419.

See Also

Overviews of available methods, multiblock, and methods organised by main structure: basic, unsupervised, asca, supervised and complex. Common functions for computation and extraction of results and plotting are found in multiblock_results and multiblock_plots, respectively.

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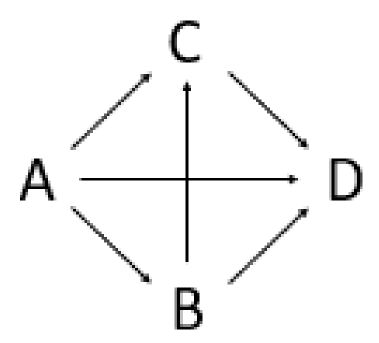
Examples

simulated

Data simulated to have certain characteristics.

Description

A dataset containing simulated data for 4 connected events where A is the starting point and D is the end point. This can be described as a directed acyclic graph (sketched below, moving left->right).



Subpaths include: ABD, AD, ABCD, ACD

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Usage

```
data(simulated)
```

Format

A list of matrices having 200 rows and 10 variables:

- A Simulated matrix A
- **B** Simulated matrix B ...

References

Tormod Næs, Rosaria Romano, Oliver Tomic, Ingrid Måge, Age Smilde, Kristian Hovde Liland, Sequential and orthogonalized PLS (SO-PLS) regression for path analysis: Order of blocks and relations between effects. Journal of Chemometrics, In Press

smbpls

Sparse Multiblock Partial Least Squares - sMB-PLS

Description

sMB-PLS is an adaptation of MB-PLS (mbpls) that enforces sparseness in loading weights when computing PLS components in the global model.

```
smbpls(
  formula,
  data,
  subset,
  na.action,
  X = NULL,
  Y = NULL,
  ncomp = 1,
  scale = FALSE,
  shrink = NULL,
  truncation = NULL,
  trunc.width = 0.95,
  blockScale = c("sqrtnvar", "ssq", "none"),
  ...
)
```

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Arguments

formula Model formula accepting a single response (block) and predictor block names

separated by + signs.

data The data set to analyse.

subset Expression for subsetting the data before modelling.

na.action How to handle NAs (no action implemented).

X list of input blocks. If X is supplied, the formula interface is skipped.

Y matrix of responses.

ncomp integer number of PLS components.

scale logical for autoscaling inputs (default = FALSE).

shrink numeric scalar indicating degree of L1-shrinkage/Soft-Thresholding (optional),

 $0 \le \text{shrink} < 1$.

truncation character indicating type of truncation (optional) "Lenth" uses asymmetric

confidence intervals to determine outlying loading weights. "quantile" uses a

quantile plot approach to determining outliers.

trunc.width numeric indicating confidence of "Lenth type" confidence interval or quantile

in "quantile plot" approach. Default = 0.95.

blockScale Either a character indicating type of block scaling or a numeric vector of

block weights (see Details).

... additional arguments to pls::plsr.

Details

Two versions of sparseness are supplied: Soft-Threshold PLS, also known as Sparse PLS, and Truncation PLS. The former uses L1 shrinkage of loading weights, while the latter comes in two flavours, both estimating inliers and outliers. The "Lenth" method uses asymmetric confidence intervals around the median of a loading weigh vector to estimate inliers. The "quantile" method uses a quantile plot approach to estimate outliers as deviations from the estimated quantile line. As with ordinary MB-PLS scaled input blocks (1/sqrt(ncol)) are used.

Block weighting is performed after scaling all variables and is by default "sqrtnvar": 1/sqrt(ncol(X[[i]])) in each block. Alternatives are "ssq": $1/norm(X[[i]], "F")^2$ and "none": 1/1. Finally, if a numeric vector is supplied, it will be used to scale the blocks after "ssq" scaling, i.e., $Z[[i]] = X[[i]] / norm(X[[i]], "F")^2 * blockScale[i]$.

Value

multiblock, mvr object with super-scores, super-loadings, block-scores and block-loading, and the underlying mvr (PLS) object for the super model, with all its result and plot possibilities. Relevant plotting functions: multiblock_plots and result functions: multiblock_results.

References

 Sæbø, S.; Almøy, T.; Aarøe, J. & Aastveit, A. ST-PLS: a multi-directional nearest shrunken centroid type classifier via PLS Journal of Chemometrics: A Journal of the Chemometrics Society, Wiley Online Library, 2008, 22, 54-62. 62 sopls

• Lê Cao, K.; Rossouw, D.; Robert-Granié, C. & Besse, P. A sparse PLS for variable selection when integrating omics data Statistical applications in genetics and molecular biology, 2008, 7.

- Liland, K.; Høy, M.; Martens, H. & Sæbø, S. Distribution based truncation for variable selection in subspace methods for multivariate regression Chemometrics and Intelligent Laboratory Systems, 2013, 122, 103-111.
- Karaman, I.; Nørskov, N.; Yde, C.; Hedemann, M.; Knudsen, K. & Kohler, A. Sparse multiblock PLSR for biomarker discovery when integrating data from LC–MS and NMR metabolomics Metabolomics, 2015, 11, 367-379.

See Also

Overviews of available methods, multiblock, and methods organised by main structure: basic, unsupervised, asca, supervised and complex.

Examples

```
data(potato)
# Truncation MB-PLS
# Loading weights inside 60% confidence intervals around the median are set to 0.
tmb <- smbpls(Sensory ~ Chemical+Compression, data=potato, ncomp = 5,</pre>
              truncation = "Lenth", trunc.width = 0.6)
# Alternative XY-interface
tmb.XY <- smbpls(X=potato[c('Chemical','Compression')], Y=potato[['Sensory']], ncomp = 5,</pre>
              truncation = "Lenth", trunc.width = 0.6)
identical(tmb, tmb.XY)
scoreplot(tmb, labels="names") # Exploiting mvr object structure from pls package
loadingweightplot(tmb, labels="names")
# Soft-Threshold / Sparse MB-PLS
# Loading weights are subtracted by 60% of maximum value.
smb <- smbpls(X=potato[c('Chemical','Compression')], Y=potato[['Sensory']],</pre>
              ncomp = 5, shrink = 0.6)
print(smb)
scoreplot(smb, labels="names") # Exploiting mvr object structure from pls package
loadingweightplot(smb, labels="names")
# Emphasis may be different for blocks
smb <- smbpls(X=potato[c('Chemical','Compression')], Y=potato[['Sensory']],</pre>
              ncomp = 5, shrink = 0.6, blockScale = c(1, 10)
```

sopls

Sequential and Orthogonalized PLS (SO-PLS)

Description

Function for computing standard SO-PLS based on the interface of the pls package.

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Usage

```
sopls(
  formula,
  ncomp,
  max_comps = min(sum(ncomp), 20),
  data,
  subset,
  na.action,
  scale = FALSE,
  validation = c("none", "CV", "L00"),
  sequential = FALSE,
  segments = 10,
  sel.comp = "opt",
  progress = TRUE,
  ...
)
```

Arguments

| formula | Model formula accepting a single response (block) and predictor block names separated by + signs. |
|------------|--|
| ncomp | Numeric vector of components per block or scalar of overall maximum components. |
| max_comps | $Maximum\ total\ number\ of\ components\ from\ all\ blocks\ combined\ (<=sum(ncomp)).$ |
| data | The data set to analyse. |
| subset | Expression for subsetting the data before modelling. |
| na.action | How to handle NAs (no action implemented). |
| scale | Logical indicating if variables should be scaled. |
| validation | Optional cross-validation strategy "CV" or "LOO". |
| sequential | Logical indicating if optimal components are chosen sequentially or globally (default=FALSE). |
| segments | Optional number of segments or list of segments for cross-validation. (See [pls::cvsegments()]). |
| sel.comp | Character indicating if sequential optimal number of components should be chosen as minimum RMSECV ('opt', default) or by Chi-square test ('chi'). |
| progress | Logical indicating if a progress bar should be displayed while cross-validating. |
| | Additional arguments to underlying methods. |

Details

SO-PLS is a method which handles two or more input blocks by sequentially performing PLS on blocks against a response and orthogonalising the remaining blocks on the extracted components. Component number optimisation can either be done globally (best combination across blocks) or sequentially (determine for one block, move to next, etc.).

Value

An sopls, mvr object with scores, loadings, etc. associated with printing (sopls_results) and plotting methods (sopls_plots).

References

Jørgensen K, Mevik BH, Næs T. Combining designed experiments with several blocks of spectroscopic data. Chemometr Intell Lab Syst. 2007;88(2): 154–166.

See Also

SO-PLS result functions, sopls_results, SO-PLS plotting functions, sopls_plots, SO-PLS Måge plot, maage, and SO-PLS path-modelling, SO_TDI. Overviews of available methods, multiblock, and methods organised by main structure: basic, unsupervised, asca, supervised and complex.

Examples

sopls_plots

Scores, loadings and plots for sopls objects

Description

Extraction of scores and loadings and adaptation of scoreplot, loadingplot and biplot from package pls for sopls objects.

```
## S3 method for class 'sopls'
loadings(object, ncomp = "all", block = 1, y = FALSE, ...)
## S3 method for class 'sopls'
scores(object, ncomp = "all", block = 1, y = FALSE, ...)
## S3 method for class 'sopls'
scoreplot(
  object,
```

```
comps = 1:2,
 ncomp = NULL,
 block = 1,
  labels,
  identify = FALSE,
  type = "p",
 xlab,
 ylab,
)
## S3 method for class 'sopls'
loadingplot(
 object,
  comps = 1:2,
  ncomp = NULL,
 block = 1,
  scatter = TRUE,
  labels,
  identify = FALSE,
  type,
  lty,
  1wd = NULL,
  pch,
  cex = NULL,
  col,
 legendpos,
 xlab,
 ylab,
 pretty.xlabels = TRUE,
 xlim,
)
## S3 method for class 'sopls'
corrplot(
 object,
 comps = 1:2,
  ncomp = NULL,
 block = 1,
 labels = TRUE,
  col = 1:5,
 plotx = TRUE,
 ploty = TRUE,
)
## S3 method for class 'sopls'
```

```
biplot(
    x,
    comps = 1:2,
    ncomp = "all",
    block = 1,
    which = c("x", "y", "scores", "loadings"),
    var.axes = FALSE,
    xlabs,
    ylabs,
    main,
    ...
)
```

Arguments

object sopls object

ncomp integer vector giving components from all blocks before block (see next ar-

gument).

block integer indicating which block to extract components from.

y logical extract Y loadings/scores instead of X loadings/scores (default = FALSE).

.. further arguments sent to the underlying plot function(s)

comps integer vector giving components, within block, to plot (see Details regarding

combination of blocks).

labels character indicating if "names" or "numbers" should be plot symbols (op-

tional).

identify logical for activating identify to interactively identify points.

type character for selecting type of plot to make. Defaults to "p" (points) for scatter

plots and "l" (lines) for line plots.

xlab character text for x labels. ylab character text for y labels.

scatter logical indicating if a scatterplot of loadings should be made (default = TRUE).

1ty Vector of line type specifications (see par for details).

lwd numeric vector of line width specifications.

pch Vector of point specifications (see points for details).

cex numeric vector of plot size expansions (see par for details).

col integer vector of symbol/line colours (see par for details).

legendpos character indicating legend position (if scatter is FALSE), e.g. legendpos

= "topright".

pretty.xlabels logical indicating if xlabels should be more nicely plotted (default = TRUE).

xlim numeric vector of length two, with the x limits of the plot (optional).

plotx locical or integer/character. Whether to plot the *X* correlation loadings,

optionally which block(s). Defaults to TRUE.

| ploty | logical. Whether to plot the Y correlation loadings. Defaults to TRUE. |
|----------|--|
| x | sopls object |
| which | character for selecting type of biplot (" x " = default, " y ", "scores", "loadings"). |
| var.axes | logical indicating if second axes of a biplot should have arrows. |
| xlabs | character vector for labelling first set of biplot points (optional). |
| ylabs | character vector for labelling second set of biplot points (optional). |
| main | character for setting the main title of a plot. |

Details

If comps is supplied as a list for scoreplot, it is assumed that its elements refer to each of the blocks up to block number block. For instance comps = list(1, 0, 1:2) will select 1 component from the first block, no components from the second block and the first two components from the last block. This must be matched by ncomp, specifying how many components were selected before block number block.

See Also

Overviews of available methods, multiblock, and methods organised by main structure: basic, unsupervised, asca, supervised and complex. Common functions for computation and extraction of results are found in sopls_results.

#' @return The score and loading functions return scores and loadings, while plot functions have no return (except use of 'identify').

Examples

```
data(potato)
so <- sopls(Sensory ~ Chemical + Compression + NIRraw, data=potato, ncomp=c(5,5,5))
# Loadings
loadings(so, ncomp=c(3), block=2)[, 1:3]
# Scores
scores(so, block=1)[, 1:4]
# Default plot from first block
scoreplot(so)
# Second block with names
scoreplot(so, ncomp=c(3), block=2, labels="names")
# Scatterplot matrix
scoreplot(so, ncomp=c(3,2), block=3, comps=1:3)
# Combination of blocks (see Details)
scoreplot(so, ncomp=c(3,2), block=3, comps=list(1,0,1))
# Default plot from first block
loadingplot(so, scatter=TRUE)
```

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```
# Second block with names
loadingplot(so, ncomp=c(3), block=2, labels="names", scatter=TRUE)

# Scatterplot matrix
loadingplot(so, ncomp=c(3,2), block=3, comps=1:3, scatter=TRUE)

# Correlation loadings
corrplot(so, block=2, ncomp=1)

# Default plot from first block
biplot(so)
```

sopls_results

Result functions for SO-PLS models

Description

Standard result functions for SO-PLS (sopls).

```
## S3 method for class 'sopls'
predict(
 object,
  newdata,
  ncomp = object$ncomp,
  type = c("response", "scores"),
 na.action = na.pass,
)
## S3 method for class 'sopls'
coef(object, ncomp = object$ncomp, intercept = FALSE, ...)
## S3 method for class 'sopls'
print(x, ...)
## S3 method for class 'sopls'
summary(
 object,
 what = c("all", "validation", "training"),
 digits = 4,
 print.gap = 2,
)
classify(object, ...)
```

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```
## S3 method for class 'sopls'
classify(object, classes, newdata, ncomp, LQ = "LDA", ...)
## S3 method for class 'sopls'
R2(object, estimate, newdata, ncomp = "all", individual = FALSE, ...)
## S3 method for class 'sopls'
RMSEP(object, estimate, newdata, ncomp = "all", individual = FALSE, ...)
pcp(object, ...)
## S3 method for class 'sopls'
pcp(object, ncomp, ...)
## Default S3 method:
pcp(object, X, ...)
cvanova(pred, ...)
## Default S3 method:
cvanova(pred, true, absRes = TRUE, ...)
## S3 method for class 'sopls'
cvanova(pred, comps, absRes = TRUE, ...)
## S3 method for class 'cvanova'
print(x, ...)
## S3 method for class 'cvanova'
summary(object, ...)
## S3 method for class 'cvanova'
plot(x, ...)
## S3 method for class 'sopls'
residuals(object, ...)
```

Arguments

object A sopls object.

newdata Optional new data with the same types of predictor blocks as the ones used for

fitting the object.

ncomp An integer vector giving the exact components to apply.

type A character for predict indicating if responses or scores should be predicted

(default = "response", or "scores"), for summary indicating which type of ex-

plained variance to compute (default = "train", alternative = "CV").

na.action Function determining what to do with missing values in newdata.

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. . . Additional arguments. Currently not implemented.

intercept A logical indicating if coefficients for the intercept should be included (default

= FALSE).

x A sopls object.

what A character indicating if summary should include all, validation or training.

digits The number of digits used for printing.

print.gap Gap between columns when printing.

classes A character vector of class labels.

LQ A character indicating if 'max' (maximum score value), 'lda' or 'qda' should

be used when classifying.

estimate A character indicating if 'train', 'CV' or 'test' results should be displayed.

individual A logical indicating if results for individual responses should be displayed.

X A list of data blocks.

pred An object holding the CV-predicted values (sopls, matrix or list of vectors)

true A numeric of true response values for CVANOVA.

absRes A logical indicating if absolute (TRUE) or squared (FALSE) residuals should

be computed.

comps An integer vector giving the exact components to apply.

Details

The parameter ncomp controls which components to apply/extract, resulting in the sequence of components leading up to the specific choice, i.e. ncomp = c(2,2,1) results in the sequence 1,0,0; 2,0,0; 2,1,0; 2,2,0; 2,2,1. Usage of the functions are shown using generics in the examples below. Prediction, regression coefficients, object printing and summary are available through: predict.sopls, coef.sopls, print.sopls and summary.sopls. Explained variances and RMSEP are available through R2.sopls and RMSEP.sopls. Principal components of predictions are available through pcp.sopls. Finally, there is work in progress on classification support through classify.sopls.

Value

Returns depend on method used, e.g. predict.sopls returns predicted responses or scores depending on inputs, coef.sopls return regression coefficients, while print and summary methods return the object invisibly.

References

Jørgensen K, Mevik BH, Næs T. Combining designed experiments with several blocks of spectroscopic data. Chemometr Intell Lab Syst. 2007;88(2): 154–166.

See Also

Overviews of available methods, multiblock, and methods organised by main structure: basic, unsupervised, asca, supervised and complex. Common functions for plotting are found in sopls_plots.

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Examples

```
mod \leftarrow sopls(Sensory[,1] \sim ., data = potato[c(1:3,9)], ncomp = 5, subset = 1:20)
testset <- potato[-(1:20),]; testset$Sensory <- testset$Sensory[,1,drop=FALSE]</pre>
predict(mod, testset, ncomp=c(2,1,2))
dim(coef(mod, ncomp=c(3,0,1))) # < variables x responses x components>
R2(mod, ncomp = c(4,1,2))
print(mod)
summary(mod)
# PCP from sopls object
modMulti \leftarrow sopls(Sensory \sim ., data = potato[c(1:3,9)], ncomp = 5, validation = "CV", segment = 5)
(PCP \leftarrow pcp(modMulti, c(2,1,2)))
scoreplot(PCP)
# PCP from matrices
preds <- modMulti$validation$Ypred[,,"2,1,2"]</pre>
PCP_default <- pcp(preds, potato[1:3])</pre>
# CVANOVA
modCV \leftarrow sopls(Sensory[,1] \sim ., data = potato[c(1:3,9)], ncomp = 5, validation = "CV", segment = 5)
summary(cva <- cvanova(modCV, "2,1,2"))</pre>
plot(cva)
```

SO_TDI

Total, direct, indirect and additional effects in SO-PLS-PM.

Description

SO-PLS-PM is the use of SO-PLS for path-modelling. This particular function is used to compute effects (explained variances) in sub-paths of the directed acyclic graph.

```
sopls_pm(
   X,
   Y,
   ncomp,
   max_comps = min(sum(ncomp), 20),
   sel.comp = "opt",
   computeAdditional = FALSE,
   sequential = FALSE,
   B = NULL,
   k = 10,
   type = "consecutive",
   simultaneous = TRUE
)
```

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```
## S3 method for class 'S0_TDI'
    print(x, showComp = TRUE, heading = "SO-PLS path effects", digits = 2, ...)
    sopls_pm_multiple(
      Χ,
      ncomp,
      max_comps = min(sum(ncomp), 20),
      sel.comp = "opt",
      computeAdditional = FALSE,
      sequential = FALSE,
      B = NULL
      k = 10,
      type = "consecutive"
    )
    ## S3 method for class 'SO_TDI_multiple'
    print(x, heading = "SO-PLS path effects", digits = 2, ...)
Arguments
                     A list of input blocks (of type matrix).
    Χ
    Υ
                     A matrix of response(s).
                     An integer vector giving the number of components per block or a single inte-
    ncomp
                     ger for common number of components.
    max_comps
                     Maximum total number of components.
                     A character or integer vector indicating the type ("opt" - minimum error /
    sel.comp
                     "chi" - chi-squared reduced) or exact number of components in selections.
    computeAdditional
                     A logical indicating if additional components should be computed.
    sequential
                     A logical indicating if sequential component optimization should be applied.
                     An integer giving the number of bootstrap replicates for variation estimation.
                     An integer indicating number of cross-validation segments (default = 10).
                     A character for selecting type of cross-validation segments (default = "con-
    type
                     secutive").
                     logical indicating if simultaneous orthogonalisation on intermediate blocks
    simultaneous
                     should be performed (default = TRUE).
                     An object of type SO_TDI.
    showComp
                     A logical indicating if components should be shown in print (default = TRUE).
                     A character giving the heading of the print.
    heading
    digits
                     An integer for selecting number of digits in print.
                     Not implemented
```

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Details

sopls_pm computes 'total', 'direct', 'indirect' and 'additional' effects for the 'first' versus the 'last' input block by cross-validated explained variances. 'total' is the explained variance when doing regression of 'first' -> 'last'. 'indirect' is the the same, but controlled for the intermediate blocks. 'direct' is the left-over part of the 'total' explained variance when subtracting the 'indirect'. Finally, 'additional' is the added explained variance of 'last' for each block following 'first'.

sopls_pm_multiple is a wrapper for sopls_pm that repeats the calculation for all pairs of blocks from 'first' to 'last'. Where sopls_pm has a separate response, Y, signifying the 'last' block, sopls_pm_multiple has multiple 'last' blocks, depending on sub-path, thus collects the response(s) from the list of blocks X.

When sel.comp = "opt", the number of components for all models are optimized using cross-validation within the ncomp and max_comps supplied. If sel.comp is "chi", an optimization is also performed, but parsimonious solutions are sought through a chi-square chriterion. When setting sel.comp to a numeric vector, exact selection of number of components is performed.

When setting B to a number, e.g. 200, the procedures above are repeated B times using bootstrapping to estimate standard deviations of the cross-validated explained variances.

Value

An object of type SO_TDI containing total, direct and indirect effects, plus possibly additional effects and standard deviations (estimated by bootstrapping).

References

- Menichelli, E., Almøy, T., Tomic, O., Olsen, N. V., & Næs, T. (2014). SO-PLS as an exploratory tool for path modelling. Food quality and preference, 36, 122-134.
- Næs, T., Romano, R., Tomic, O., Måge, I., Smilde, A., & Liland, K. H. (2020). Sequential and orthogonalized PLS (SO-PLS) regression for path analysis: Order of blocks and relations between effects. Journal of Chemometrics, e3243.

See Also

Overviews of available methods, multiblock, and methods organised by main structure: basic, unsupervised, asca, supervised and complex.

Examples

```
# Single path for the potato data:
data(potato)
pot.pm <- sopls_pm(potato[1:3], potato[['Sensory']], c(5,5,5), computeAdditional=TRUE)
pot.pm

# Corresponding SO-PLS model:
# so <- sopls(Sensory ~ ., data=potato[c(1,2,3,9)], ncomp=c(5,5,5), validation="CV", segments=10)
# maageSeq(pot.so, compSeq = c(3,2,4))

# All path in the forward direction for the wine data:
data(wine)
pot.pm.multiple <- sopls_pm_multiple(wine, ncomp = c(4,2,9,8))</pre>
```

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```
pot.pm.multiple
```

statis

Structuration des Tableaux à Trois Indices de la Statistique - STATIS

Description

This is a wrapper for the ade4::statis function for computing STATIS.

Usage

```
statis(X, ncomp = 3, scannf = FALSE, tol = 1e-07, ...)
```

Arguments

X list of input blocks.

ncomp integer number of components to extract.

scannf logical indicating if eigenvalue bar plot shoulde be displayed.

tol numeric eigenvalue threshold tolerance.

... additional arguments (not used).

Details

STATIS is a method, related to MFA, for analysing two or more blocks. It also decomposes the data into a low-dimensional subspace but uses a different scaling of the individual blocks.

Value

multiblock object including relevant scores and loadings. Relevant plotting functions: multiblock_plots and result functions: multiblock_results.

References

Lavit, C.; Escoufier, Y.; Sabatier, R.; Traissac, P. (1994). The ACT (STATIS method). Computational Statistics & Data Analysis. 18: 97

See Also

Overviews of available methods, multiblock, and methods organised by main structure: basic, unsupervised, asca, supervised and complex. Common functions for computation and extraction of results and plotting are found in multiblock_results and multiblock_plots, respectively.

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Examples

```
data(candies)
candyList <- lapply(1:nlevels(candies$candy),function(x)candies$assessment[candies$candy==x,])
can.statis <- statis(candyList)
plot(scores(can.statis), labels="names")</pre>
```

supervised

Supervised Multiblock Methods

Description

Collection of supervised multiblock methods:

- MB-PLS Multiblock Partial Least Squares (mbpls)
- sMB-PLS Sparse Multiblock Partial Least Squares (smbpls)
- SO-PLS Sequential and Orthogonalized PLS (sopls)
- PO-PLS Parallel and Orthogonalized PLS (popls)
- ROSA Response Oriented Sequential Alternation (rosa)
- mbRDA Multiblock Redundancy Analysis (mbrda)

See Also

Overviews of available methods, multiblock, and methods organised by main structure: basic, unsupervised, asca, supervised and complex. Common functions for computation and extraction of results and plotting are found in multiblock_results and multiblock_plots, respectively.

Examples

```
data(potato)
mb <- mbpls(Sensory ~ Chemical + Compression, data=potato, ncomp = 5)
print(mb)

# Convert data.frame with AsIs objects to list of matrices
potatoList <- lapply(potato, unclass)
mbr <- mbrda(Sensory ~ Chemical + Compression, data=potatoList, ncomp = 10)
print(mbr)
scoreplot(mbr, labels="names")</pre>
```

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unique_combos

Unique combinations of blocks

Description

Compute a list of all possible block combinations where the number of blocks in each combination is limited by parameters min_level and max_level.

Usage

```
unique_combos(n_block, max_level, min_level = 2)
```

Arguments

n_block integer number of input blocks.

max_level integer maximum number of blocks per combination.
min_level integer minimum number of blocks per combination.

Details

This function is used for minimal redundancy implementations of rosa and sopls and for lookups into computed components.

Value

A list of unique block combinations.

Examples

```
unique_combos(3, 2)
```

unsupervised

Unsupervised Multiblock Methods

Description

Collection of unsupervised multiblock methods:

- SCA Simultaneous Component Analysis (sca)
- GCA Generalized Canonical Analysis (gca)
- GPA Generalized Procrustes Analysis (gpa)
- MFA Multiple Factor Analysis (mfa)
- PCA-GCA (pcagca)

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- DISCO Distinctive and Common Components with SCA (disco)
- HPCA Hierarchical Principal component analysis (hpca)
- MCOA Multiple Co-Inertia Analysis (mcoa)
- JIVE Joint and Individual Variation Explained (jive)
- STATIS Structuration des Tableaux à Trois Indices de la Statistique (statis)
- HOGSVD Higher Order Generalized SVD (hogsvd)

Details

Original documentation of STATIS: statis. JIVE, STATIS and HOGSVD assume variable linked matrices/data.frames, while SCA handles both links.

See Also

Overviews of available methods, multiblock, and methods organised by main structure: basic, unsupervised, asca, supervised and complex. Common functions for computation and extraction of results and plotting are found in multiblock_results and multiblock_plots, respectively.

Examples

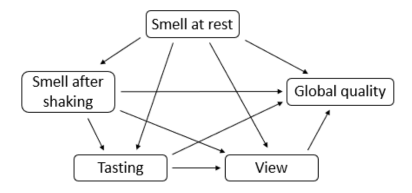
wine

Wines of Val de Loire

Description

This dataset contains sensory assessment of 21 wines. The assessments are grouped according to the tasting process and thus have a natural ordering with a all blocks pointing forward to all remaining blocks in the process.

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Usage

data(wine)

Format

A data.frame having 21 rows and 5 variables:

Smell at rest Matrix of sensory assessments

View Matrix of sensory assessments

Smell after shaking Matrix of sensory assessments

Tasting Matrix of sensory assessments

Global quality Matrix of sensory assessments

References

Escofier B, Pages L. Analyses Factorielles Simples and Multiples. Paris: Dunod; 1988.

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