

# Package ‘chemometrics’

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**Suggests** gclus

**Description** R companion to the book ``Introduction to Multivariate Statistical Analysis in Chemometrics'' written by K. Varmuza and P. Filzmoser (2009).

**License** GPL (>= 3)

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chemometrics-package *This package is the R companion to the book "Introduction to Multivariate Statistical Analysis in Chemometrics" written by K. Varmuza and P. Filzmoser (2009).*

---

### Description

Included are functions for multivariate statistical methods, tools for diagnostics, multivariate calibration, cross validation and bootstrap, clustering, etc.

### Details

The package can be used to verify the examples in the book. It can also be used to analyze own data.

### Author(s)

P. Filzmoser <P.Filzmoser@tuwien.ac.at>

### References

K. Varmuza and P. Filzmoser: Introduction to Multivariate Statistical Analysis in Chemometrics. CRC Press, Boca Raton, FL, 2009.

---

alr *additive logratio transformation*

---

### Description

A data transformation according to the additive logratio transformation is done.

### Usage

```
alr(X, divisorvar)
```

### Arguments

X                    numeric data frame or matrix  
divisorvar        number of the column of X for the variable to divide with

### Details

The alr transformation is one possibility to transform compositional data to a real space. Afterwards, the transformed data can be analyzed in the usual way.

**Value**

Returns the transformed data matrix with one variable (divisor variable) less.

**Author(s)**

Peter Filzmoser <P.Filzmoser@tuwien.ac.at>

**References**

K. Varmuza and P. Filzmoser: Introduction to Multivariate Statistical Analysis in Chemometrics. CRC Press, Boca Raton, FL, 2009.

**See Also**

[clr,ilr](#)

**Examples**

```
data(glass)
glass_alr <- alr(glass,1)
```

---

ash

*ash data*

---

**Description**

Data from 99 ash samples originating from different biomass, measured on 9 variables; 8 log-transformed variables are added.

**Usage**

```
data(ash)
```

**Format**

A data frame with 99 observations on the following 17 variables.

SOT a numeric vector  
P205 a numeric vector  
SiO2 a numeric vector  
Fe203 a numeric vector  
Al203 a numeric vector  
CaO a numeric vector  
MgO a numeric vector  
Na2O a numeric vector  
K2O a numeric vector

log(P2O5) a numeric vector  
log(SiO2) a numeric vector  
log(Fe2O3) a numeric vector  
log(Al2O3) a numeric vector  
log(CaO) a numeric vector  
log(MgO) a numeric vector  
log(Na2O) a numeric vector  
log(K2O) a numeric vector

### Details

The dependent variable Softening Temperature (SOT) of ash should be modeled by the elemental composition of the ash data. Data from 99 ash samples - originating from different biomass - comprise the experimental SOT (630-1410 centigrades), and the experimentally determined eight mass concentrations the listed elements. Since the distribution of the elements is skewed, the log-transformed variables have been added.

### Source

K. Varmuza and P. Filzmoser: Introduction to Multivariate Statistical Analysis in Chemometrics. CRC Press, Boca Raton, FL, 2009.

### References

K. Varmuza and P. Filzmoser: Introduction to Multivariate Statistical Analysis in Chemometrics. CRC Press, Boca Raton, FL, 2009.

### Examples

```
data(ash)  
str(ash)
```

---

cereal

*Data from cereals*

---

### Description

For 15 cereals an X and Y data set, measured on the same objects, is available. The X data are 145 infrared spectra, and the Y data are 6 chemical/technical properties (Heating value, C, H, N, Starch, Ash). Also the scaled Y data are included (mean 0, variance 1 for each column). The cereals come from 5 groups B=Barley, M=Maize, R=Rye, T=Triticale, W=Wheat.

### Usage

```
data(cereal)
```

**Format**

A data frame with 15 objects and 3 list elements:

X matrix with 15 rows and 145 columns

Y matrix with 15 rows and 6 columns

Ysc matrix with 15 rows and 6 columns

**Details**

The data set can be used for PLS2.

**Source**

K. Varmuza and P. Filzmoser: Introduction to Multivariate Statistical Analysis in Chemometrics. CRC Press, Boca Raton, FL, 2009.

**References**

K. Varmuza and P. Filzmoser: Introduction to Multivariate Statistical Analysis in Chemometrics. CRC Press, Boca Raton, FL, 2009.

**Examples**

```
data(cereal)
names(cereal)
```

---

clr	<i>centered logratio transformation</i>
-----	---

---

**Description**

A data transformation according to the centered logratio transformation is done.

**Usage**

```
clr(X)
```

**Arguments**

X                    numeric data frame or matrix

**Details**

The clr transformation is one possibility to transform compositional data to a real space. Afterwards, the transformed data can be analyzed in the usual way.

**Value**

Returns the transformed data matrix with the same dimension as  $X$ .

**Author(s)**

Peter Filzmoser <P.Filzmoser@tuwien.ac.at>

**References**

K. Varmuza and P. Filzmoser: Introduction to Multivariate Statistical Analysis in Chemometrics. CRC Press, Boca Raton, FL, 2009.

**See Also**

[alr,ilr](#)

**Examples**

```
data(glass)
glass_clr <- clr(glass)
```

---

clvalidity	<i>compute and plot cluster validity</i>
------------	--

---

**Description**

A cluster validity measure based on within- and between-sum-of-squares is computed and plotted for the methods k-means, fuzzy c-means, and model-based clustering.

**Usage**

```
clvalidity(x, clnumb = c(2:10))
```

**Arguments**

x	input data matrix
clnumb	range for the desired number of clusters

**Details**

The validity measure for a number  $k$  of clusters is  $\sum_j W_j$  divided by  $\sum_{j < l} B_{jl}$  with  $W_j$  is the sum of squared distances of the objects in each cluster cluster to its center, and  $B_{jl}$  is the squared distance between the cluster centers of cluster  $j$  and  $l$ .

**Value**

validity	vector with validity measure for the desired numbers of clusters
----------	--

**Author(s)**

Peter Filzmoser <P.Filzmoser@tuwien.ac.at>

**References**

K. Varmuza and P. Filzmoser: Introduction to Multivariate Statistical Analysis in Chemometrics. CRC Press, Boca Raton, FL, 2009.

**See Also**

[princomp](#)

**Examples**

```
data(glass)
require(robustbase)
res <- pcaCV(glass, segments=4, repl=100, cex.lab=1.2, ylim=c(0, 1), las=1)
```

---

delintercept

*Delete intercept from model matrix*

---

**Description**

A utility function to delete any intercept column from a model matrix, and adjust the assign attribute correspondingly.

**Usage**

```
delintercept(mm)
```

**Arguments**

mm                    Model matrix

**Value**

A model matrix without intercept column.

**Author(s)**

B.-H. Mevik and Ron Wehrens

**See Also**

[delete.intercept](#)

---

drawMahal	<i>Draws ellipses according to Mahalanobis distances</i>
-----------	--

---

### Description

For 2-dimensional data a scatterplot is made. Additionally, ellipses corresponding to certain Mahalanobis distances and quantiles of the data are drawn.

### Usage

```
drawMahal(x, center, covariance, quantile = c(0.975, 0.75, 0.5, 0.25), m = 1000,
lwdcrit = 1, ...)
```

### Arguments

x	numeric data frame or matrix with 2 columns
center	vector of length 2 with multivariate center of x
covariance	2 by 2 covariance matrix of x
quantile	vector of quantiles for the Mahalanobis distance
m	number of points where the ellipses should pass through
lwdcrit	line width of the ellipses
...	additional graphics parameters, see <a href="#">par</a>

### Details

For multivariate normally distributed data, a fraction of 1-quantile of data should be outside the ellipses. For center and covariance also robust estimators, e.g. from the MCD estimator, can be supplied.

### Value

A scatterplot with the ellipses is generated.

### Author(s)

Peter Filzmoser <P.Filzmoser@tuwien.ac.at>

### References

K. Varmuza and P. Filzmoser: Introduction to Multivariate Statistical Analysis in Chemometrics. CRC Press, Boca Raton, FL, 2009.

### See Also

[covMcd](#)

**Examples**

```
data(glass)
data(glass.grp)
x=glass[,c(2,7)]
require(robustbase)
x.mcd=covMcd(x)
drawMahal(x,center=x.mcd$center,covariance=x.mcd$cov,quantile=0.975,pch=glass.grp)
```

---

glass

*glass vessels data*

---

**Description**

13 different measurements for 180 archaeological glass vessels from different groups are included.

**Usage**

```
data(glass)
```

**Format**

A data matrix with 180 objects and 13 variables.

**Details**

This is a matrix with 180 objects and 13 columns.

**Source**

Janssen, K.H.A., De Raedt, I., Schalm, O., Veeckman, J.: Microchim. Acta 15 (suppl.) (1998) 253-267. Compositions of 15th - 17th century archaeological glass vessels excavated in Antwerp.

**References**

K. Varmuza and P. Filzmoser: Introduction to Multivariate Statistical Analysis in Chemometrics. CRC Press, Boca Raton, FL, 2009.

**Examples**

```
data(glass)
str(glass)
```

---

glass.grp	<i>glass types of the glass data</i>
-----------	--------------------------------------

---

**Description**

13 different measurements for 180 archaeological glass vessels from different groups are included. These groups are certain types of glasses.

**Usage**

```
data(glass.grp)
```

**Format**

The format is: num [1:180] 1 1 1 1 1 1 1 1 1 1 ...

**Details**

This is a vector with 180 elements referring to the groups.

**Source**

Janssen, K.H.A., De Raedt, I., Schalm, O., Veeckman, J.: *Microchim. Acta* 15 (suppl.) (1998) 253-267. Compositions of 15th - 17th century archaeological glass vessels excavated in Antwerp.

**References**

K. Varmuza and P. Filzmoser: *Introduction to Multivariate Statistical Analysis in Chemometrics*. CRC Press, Boca Raton, FL, 2009.

**Examples**

```
data(glass.grp)
str(glass.grp)
```

---

hyptis	<i>Hyptis data set</i>
--------	------------------------

---

**Description**

30 objects (Wild growing, flowering *Hyptis suaveolens*) and 7 variables (chemotypes), and 2 variables that explain the grouping (4 groups).

**Usage**

```
data(hyptis)
```

**Format**

A data frame with 30 observations on the following 9 variables.

Sabinene a numeric vector

Pinene a numeric vector

Cineole a numeric vector

Terpinene a numeric vector

Fenchone a numeric vector

Terpinolene a numeric vector

Fenchol a numeric vector

Location a factor with levels East-high East-low North South

Group a numeric vector with the group information

**Details**

This data set can be used for cluster analysis.

**References**

P. Grassi, M.J. Nunez, K. Varmuza, and C. Franz: Chemical polymorphism of essential oils of *Hyptis suaveolens* from El Salvador. *Flavour and Fragrance*, 20, 131-135, 2005. K. Varmuza and P. Filzmoser: *Introduction to Multivariate Statistical Analysis in Chemometrics*. CRC Press, Boca Raton, FL, 2009

**Examples**

```
data(hyptis)
str(hyptis)
```

---

ilr

*isometric logratio transformation*

---

**Description**

A data transformation according to the isometric logratio transformation is done.

**Usage**

```
ilr(X)
```

**Arguments**

X numeric data frame or matrix

**Details**

The ilr transformation is one possibility to transform compositional data to a real space. Afterwards, the transformed data can be analyzed in the usual way.

**Value**

Returns the transformed data matrix with one dimension less than X.

**Author(s)**

Peter Filzmoser <P.Filzmoser@tuwien.ac.at>

**References**

K. Varmuza and P. Filzmoser: Introduction to Multivariate Statistical Analysis in Chemometrics. CRC Press, Boca Raton, FL, 2009.

**See Also**

[alr,clr](#)

**Examples**

```
data(glass)
glass_ilr <- ilr(glass)
```

---

knnEval

*kNN evaluation by CV*

---

**Description**

Evaluation for k-Nearest-Neighbors (kNN) classification by cross-validation

**Usage**

```
knnEval(X, grp, train, kfold = 10, knnvec = seq(2, 20, by = 2), plotit = TRUE,
        legend = TRUE, legpos = "bottomright", ...)
```

**Arguments**

X	standardized complete X data matrix (training and test data)
grp	factor with groups for complete data (training and test data)
train	row indices of X indicating training data objects
kfold	number of folds for cross-validation
knnvec	range for k for the evaluation of kNN
plotit	if TRUE a plot will be generated

legend	if TRUE a legend will be added to the plot
legpos	positioning of the legend in the plot
...	additional plot arguments

### Details

The data are split into a calibration and a test data set (provided by "train"). Within the calibration set "kfold"-fold CV is performed by applying the classification method to "kfold"-1 parts and evaluation for the last part. The misclassification error is then computed for the training data, for the CV test data (CV error) and for the test data.

### Value

trainerr	training error rate
testerr	test error rate
cvMean	mean of CV errors
cvSe	standard error of CV errors
cverr	all errors from CV
knnvec	range for k for the evaluation of kNN, taken from input

### Author(s)

Peter Filzmoser <P.Filzmoser@tuwien.ac.at>

### References

K. Varmuza and P. Filzmoser: Introduction to Multivariate Statistical Analysis in Chemometrics. CRC Press, Boca Raton, FL, 2009.

### See Also

[knn](#)

### Examples

```
data(fgl, package="MASS")
grp=fgl$type
X=scale(fgl[,1:9])
k=length(unique(grp))
dat=data.frame(grp,X)
n=nrow(X)
ntrain=round(n*2/3)
require(class)
set.seed(123)
train=sample(1:n,ntrain)
resknn=knnEval(X,grp,train,knnvec=seq(1,30,by=1),legpos="bottomright")
title("kNN classification")
```

---

lassocof	<i>Plot Lasso coefficients</i>
----------	--------------------------------

---

**Description**

Plots the coefficients of Lasso regression

**Usage**

```
lassocof(formula, data, sopt, plot.opt = TRUE, ...)
```

**Arguments**

formula	formula, like $y \sim X$ , i.e., dependent~response variables
data	data frame to be analyzed
sopt	optimal fraction from Lasso regression, see details
plot.opt	if TRUE a plot will be generated
...	additional plot arguments

**Details**

Using the function [lassoCV](#) for cross-validation, the optimal fraction sopt can be determined. Besides a plot for the Lasso coefficients for all values of fraction, the optimal fraction is taken to compute the number of coefficients that are exactly zero.

**Value**

coefficients	regression coefficients for the optimal Lasso parameter
sopt	optimal value for fraction
numb.zero	number of zero coefficients for optimal fraction
numb.nonzero	number of nonzero coefficients for optimal fraction
ind	index of fraction with optimal choice for fraction

**Author(s)**

Peter Filzmoser <P.Filzmoser@tuwien.ac.at>

**References**

K. Varmuza and P. Filzmoser: Introduction to Multivariate Statistical Analysis in Chemometrics. CRC Press, Boca Raton, FL, 2009.

**See Also**

[cv.lars](#), [lassoCV](#)

**Examples**

```
data(PAC)
res=lassocoeff(y~X,data=PAC,sopt=0.3)
```

lassoCV

*CV for Lasso regression***Description**

Performs cross-validation (CV) for Lasso regression and plots the results in order to select the optimal Lasso parameter.

**Usage**

```
lassoCV(formula, data, K = 10, fraction = seq(0, 1, by = 0.05), trace = FALSE,
plot.opt = TRUE, sdfact = 2, legpos = "topright", ...)
```

**Arguments**

formula	formula, like $y \sim X$ , i.e., dependent~response variables
data	data frame to be analyzed
K	the number of segments to use for CV
fraction	fraction for Lasso parameters to be used for evaluation, see details
trace	if 'TRUE', intermediate results are printed
plot.opt	if TRUE a plot will be generated that shows optimal choice for "fraction"
sdfact	factor for the standard error for selection of the optimal parameter, see details
legpos	position of the legend in the plot
...	additional plot arguments

**Details**

The parameter "fraction" is the sum of absolute values of the regression coefficients for a particular Lasso parameter on the sum of absolute values of the regression coefficients for the maximal possible value of the Lasso parameter (unconstrained case), see also [lars](#). The optimal fraction is chosen according to the following criterion: Within the CV scheme, the mean of the SEPs is computed, as well as their standard errors. Then one searches for the minimum of the mean SEPs and adds  $sdfact \cdot standarderror$ . The optimal fraction is the smallest fraction with an MSEP below this bound.

**Value**

cv	MSEP values at each value of fraction
cv.error	standard errors for each value of fraction
SEP	SEP value for each value of fraction
ind	index of fraction with optimal choice for fraction
sopt	optimal value for fraction
fraction	all values considered for fraction

**Author(s)**

Peter Filzmoser <P.Filzmoser@tuwien.ac.at>

**References**

K. Varmuza and P. Filzmoser: Introduction to Multivariate Statistical Analysis in Chemometrics. CRC Press, Boca Raton, FL, 2009.

**See Also**

[cv.lars](#), [lassocoef](#)

**Examples**

```
data(PAC)
# takes some time: # res <- lassoCV(y~X,data=PAC,K=5,fraction=seq(0.1,0.5,by=0.1))
```

---

 lmCV

---

*Repeated Cross Validation for lm*


---

**Description**

Repeated Cross Validation for multiple linear regression: a cross-validation is performed repeatedly, and standard evaluation measures are returned.

**Usage**

```
lmCV(formula, data, repl = 100, segments = 4, segment.type = c("random", "consecutive",
"interleaved"), length.seg, trace = FALSE, ...)
```

**Arguments**

formula	formula, like $y \sim X$ , i.e., dependent~response variables
data	data set including y and X
repl	number of replication for Cross Validation
segments	number of segments used for splitting into training and test data
segment.type	"random", "consecutive", "interleaved" splitting into training and test data
length.seg	number of parts for training and test data, overwrites segments
trace	if TRUE intermediate results are reported
...	additional plotting arguments

**Details**

Repeating the cross-validation with allow for a more careful evaluation.

**Value**

residuals	matrix of size length(y) x repl with residuals
predicted	matrix of size length(y) x repl with predicted values
SEP	Standard Error of Prediction computed for each column of "residuals"
SEPM	mean SEP value
RMSEP	Root MSEP value computed for each column of "residuals"
RMSEPM	mean RMSEP value

**Author(s)**

Peter Filzmoser <P.Filzmoser@tuwien.ac.at>

**References**

K. Varmuza and P. Filzmoser: Introduction to Multivariate Statistical Analysis in Chemometrics. CRC Press, Boca Raton, FL, 2009.

**See Also**

[mvr](#)

**Examples**

```
data(ash)
set.seed(100)
res=lmCV(SOT~.,data=ash,repl=10)
hist(res$SEP)
```

---

Moutlier

*Plots classical and robust Mahalanobis distances*

---

**Description**

For multivariate outlier detection the Mahalanobis distance can be used. Here a plot of the classical and the robust (based on the MCD) Mahalanobis distance is drawn.

**Usage**

```
Moutlier(X, quantile = 0.975, plot = TRUE, ...)
```

**Arguments**

X	numeric data frame or matrix
quantile	cut-off value (quantile) for the Mahalanobis distance
plot	if TRUE a plot is generated
...	additional graphics parameters, see <a href="#">par</a>

**Details**

For multivariate normally distributed data, a fraction of 1-quantile of data can be declared as potential multivariate outliers. These would be identified with the Mahalanobis distance based on classical mean and covariance. For deviations from multivariate normality center and covariance have to be estimated in a robust way, e.g. by the MCD estimator. The resulting robust Mahalanobis distance is suitable for outlier detection. Two plots are generated, showing classical and robust Mahalanobis distance versus the observation numbers.

**Value**

md	Values of the classical Mahalanobis distance
rd	Values of the robust Mahalanobis distance
cutoff	Value with the outlier cut-off
...	

**Author(s)**

Peter Filzmoser <P.Filzmoser@tuwien.ac.at>

**References**

K. Varmuza and P. Filzmoser: Introduction to Multivariate Statistical Analysis in Chemometrics. CRC Press, Boca Raton, FL, 2009.

**See Also**

[covMcd](#)

**Examples**

```
data(glass)
data(glass.grp)
x=glass[,c(2,7)]
require(robustbase)
res <- Moutlier(glass,quantile=0.975,pch=glass.grp)
```

---

mvr\_dcv

*Repeated double-cross-validation for PLS and PCR*

---

**Description**

Performs a careful evaluation by repeated double-CV for multivariate regression methods, like PLS and PCR.

**Usage**

```
mvr_dcv(formula, ncomp, data, subset, na.action,
  method = c("kernelpls", "widekernelpls", "simpls", "oscorespls", "svdpc"),
  scale = FALSE, repl = 100, sdfact = 2,
  segments0 = 4, segment0.type = c("random", "consecutive", "interleaved"),
  length.seg0, segments = 10, segment.type = c("random", "consecutive", "interleaved"),
  length.seg, trace = FALSE, plot.opt = FALSE, selstrat = "hastie", ...)
```

**Arguments**

formula	formula, like $y \sim X$ , i.e., dependent~response variables
ncomp	number of PLS components
data	data frame to be analyzed
subset	optional vector to define a subset
na.action	a function which indicates what should happen when the data contain missing values
method	the multivariate regression method to be used, see <a href="#">mvr</a>
scale	numeric vector, or logical. If numeric vector, X is scaled by dividing each variable with the corresponding element of 'scale'. If 'scale' is 'TRUE', X is scaled by dividing each variable by its sample standard deviation. If cross-validation is selected, scaling by the standard deviation is done for every segment.
repl	Number of replication for the double-CV
sdfact	factor for the multiplication of the standard deviation for the determination of the optimal number of components
segments0	the number of segments to use for splitting into training and test data, or a list with segments (see <a href="#">mvrCv</a> )
segment0.type	the type of segments to use. Ignored if 'segments0' is a list
length.seg0	Positive integer. The length of the segments to use. If specified, it overrides 'segments' unless 'segments0' is a list
segments	the number of segments to use for selecting the optimal number of components, or a list with segments (see <a href="#">mvrCv</a> )
segment.type	the type of segments to use. Ignored if 'segments' is a list
length.seg	Positive integer. The length of the segments to use. If specified, it overrides 'segments' unless 'segments' is a list
trace	logical; if 'TRUE', the segment number is printed for each segment
plot.opt	if TRUE a plot will be generated that shows the selection of the optimal number of components for each step of the CV
selstrat	method that defines how the optimal number of components is selected, should be one of "diffnext", "hastie", "relchange"; see details
...	additional parameters

## Details

In this cross-validation (CV) scheme, the optimal number of components is determined by an additional CV in the training set, and applied to the test set. The procedure is repeated `repl` times. There are different strategies for determining the optimal number of components (parameter `selstrat`): "diffnext" compares  $MSE + s_{\text{fact}} * sd(MSE)$  among the neighbors, and if the MSE falls outside this bound, this is the optimal number. "hastie" searches for the number of components with the minimum of the mean MSE's. The optimal number of components is the model with the smallest number of components which is still in the range of the  $MSE + s_{\text{fact}} * sd(MSE)$ , where MSE and sd are taken from the minimum. "relchange" is a strategy where the relative change is combined with "hastie": First the minimum of the mean MSE's is searched, and MSE's of larger components are omitted. For this selection, the relative change in MSE compared to the min, and relative to the max, is computed. If this change is very small (e.g. smaller than 0.005), these components are omitted. Then the "hastie" strategy is applied for the remaining MSE's.

## Value

<code>resopt</code>	array [ <code>nrow(Y)</code> x <code>ncol(Y)</code> x <code>repl</code> ] with residuals using optimum number of components
<code>predopt</code>	array [ <code>nrow(Y)</code> x <code>ncol(Y)</code> x <code>repl</code> ] with predicted Y using optimum number of components
<code>optcomp</code>	matrix [ <code>segments0</code> x <code>repl</code> ] optimum number of components for each training set
<code>pred</code>	array [ <code>nrow(Y)</code> x <code>ncol(Y)</code> x <code>ncomp</code> x <code>repl</code> ] with predicted Y for all numbers of components
<code>SEPop</code>	SEP over all residuals using optimal number of components
<code>sIQRopt</code>	spread of inner half of residuals as alternative robust spread measure to the <code>SEPop</code>
<code>SMADopt</code>	MAD of residuals as alternative robust spread measure to the <code>SEPop</code>
<code>MSEPop</code>	MSEP over all residuals using optimal number of components
<code>afinal</code>	final optimal number of components
<code>SEPfinal</code>	vector of length <code>ncomp</code> with final SEP values; use the element <code>afinal</code> for the optimal SEP

## Author(s)

Peter Filzmoser <P.Filzmoser@tuwien.ac.at>

## References

K. Varmuza and P. Filzmoser: Introduction to Multivariate Statistical Analysis in Chemometrics. CRC Press, Boca Raton, FL, 2009.

## See Also

[mvr](#)

## Examples

```
data(NIR)
X <- NIR$xNIR[1:30,] # first 30 observations - for illustration
y <- NIR$yGlcEtOH[1:30,1] # only variable Glucose
NIR.Glc <- data.frame(X=X, y=y)
res <- mvr_dcv(y~,data=NIR.Glc,ncomp=10,method="simpls",repl=10)
```

---

nipals

*PCA calculation with the NIPALS algorithm*

---

## Description

NIPALS is an algorithm for computing PCA scores and loadings.

## Usage

```
nipals(X, a, it = 10, tol = 1e-04)
```

## Arguments

X	numeric data frame or matrix
a	maximum number of principal components to be computed
it	maximum number of iterations
tol	tolerance limit for convergence of the algorithm

## Details

The NIPALS algorithm is well-known in chemometrics. It is an algorithm for computing PCA scores and loadings. The advantage is that the components are computed one after the other, and one could stop at a desired number of components.

## Value

T	matrix with the PCA scores
P	matrix with the PCA loadings

## Author(s)

Peter Filzmoser <P.Filzmoser@tuwien.ac.at>

## References

K. Varmuza and P. Filzmoser: Introduction to Multivariate Statistical Analysis in Chemometrics. CRC Press, Boca Raton, FL, 2009.

## See Also

[princomp](#)

**Examples**

```
data(glass)
res <- nipals(glass,a=2)
```

---

NIR

*NIR data*

---

**Description**

For 166 alcoholic fermentation mashes of different feedstock (rye, wheat and corn) we have 235 variables (X) containing the first derivatives of near infrared spectroscopy (NIR) absorbance values at 1115-2285 nm, and two variables (Y) containing the concentration of glucose and ethanol (in g/L).

**Usage**

```
data(NIR)
```

**Format**

A data frame with 166 objects and 2 list elements:

xNIR data frame with 166 rows and 235 columns

yGlcEtOH data frame with 166 rows and 2 columns

**Details**

The data can be used for linear and non-linear models.

**Source**

B. Liebmann, A. Friedl, and K. Varmuza. Determination of glucose and ethanol in bioethanol production by near infrared spectroscopy and chemometrics. *Anal. Chim. Acta*, 642:171-178, 2009.

**References**

B. Liebmann, A. Friedl, and K. Varmuza. Determination of glucose and ethanol in bioethanol production by near infrared spectroscopy and chemometrics. *Anal. Chim. Acta*, 642:171-178, 2009.

**Examples**

```
data(NIR)
str(NIR)
```

---

nnetEval                      *Neural network evaluation by CV*

---

### Description

Evaluation for Artificial Neural Network (ANN) classification by cross-validation

### Usage

```
nnetEval(X, grp, train, kfold = 10, decay = seq(0, 10, by = 1), size = 30,
maxit = 100, plotit = TRUE, legend = TRUE, legpos = "bottomright", ...)
```

### Arguments

X	standardized complete X data matrix (training and test data)
grp	factor with groups for complete data (training and test data)
train	row indices of X indicating training data objects
kfold	number of folds for cross-validation
decay	weight decay, see <a href="#">nnet</a> , can be a vector with several values - but then "size" can be only one value
size	number of hidden units, see <a href="#">nnet</a> , can be a vector with several values - but then "decay" can be only one value
maxit	maximal number of iterations for ANN, see <a href="#">nnet</a>
plotit	if TRUE a plot will be generated
legend	if TRUE a legend will be added to the plot
legpos	positioning of the legend in the plot
...	additional plot arguments

### Details

The data are split into a calibration and a test data set (provided by "train"). Within the calibration set "kfold"-fold CV is performed by applying the classification method to "kfold"-1 parts and evaluation for the last part. The misclassification error is then computed for the training data, for the CV test data (CV error) and for the test data.

### Value

trainerr	training error rate
testerr	test error rate
cvMean	mean of CV errors
cvSe	standard error of CV errors
cverr	all errors from CV
decay	value(s) for weight decay, taken from input
size	value(s) for number of hidden units, taken from input

**Author(s)**

Peter Filzmoser <P.Filzmoser@tuwien.ac.at>

**References**

K. Varmuza and P. Filzmoser: Introduction to Multivariate Statistical Analysis in Chemometrics. CRC Press, Boca Raton, FL, 2009.

**See Also**

[nnet](#)

**Examples**

```
data(fgl, package="MASS")
grp=fgl$type
X=scale(fgl[,1:9])
k=length(unique(grp))
dat=data.frame(grp,X)
n=nrow(X)
ntrain=round(n*2/3)
require(nnet)
set.seed(123)
train=sample(1:n,ntrain)
resnnet=nnetEval(X,grp,train,decay=c(0,0.01,0.1,0.15,0.2,0.3,0.5,1),
  size=20,maxit=20)
```

---

PAC

*GC retention indices*

---

**Description**

For 209 objects an X-data set (467 variables) and a y-data set (1 variable) is available. The data describe GC-retention indices of polycyclic aromatic compounds (y) which have been modeled by molecular descriptors (X).

**Usage**

```
data(PAC)
```

**Format**

A data frame with 209 objects and 2 list elements:

y numeric vector with length 209

X matrix with 209 rows and 467 columns

**Details**

The data can be used for linear and non-linear models.

**Source**

K. Varmuza and P. Filzmoser: Introduction to Multivariate Statistical Analysis in Chemometrics. CRC Press, Boca Raton, FL, 2009.

**References**

K. Varmuza and P. Filzmoser: Introduction to Multivariate Statistical Analysis in Chemometrics. CRC Press, Boca Raton, FL, 2009.

**Examples**

```
data(PAC)
names(PAC)
```

---

pcaCV	<i>Determine the number of PCA components with repeated cross validation</i>
-------	--

---

**Description**

By splitting data into training and test data repeatedly the number of principal components can be determined by inspecting the distribution of the explained variances.

**Usage**

```
pcaCV(X, amax, center = TRUE, scale = TRUE, repl = 50, segments = 4,
segment.type = c("random", "consecutive", "interleaved"), length.seg, trace = FALSE,
plot.opt = TRUE, ...)
```

**Arguments**

X	numeric data frame or matrix
amax	maximum number of components for evaluation
center	should the data be centered? TRUE or FALSE
scale	should the data be scaled? TRUE or FALSE
repl	number of replications of the CV procedure
segments	number of segments for CV
segment.type	"random", "consecutive", "interleaved" splitting into training and test data
length.seg	number of parts for training and test data, overwrites segments
trace	if TRUE intermediate results are reported
plot.opt	if TRUE the results are shown by boxplots
...	additional graphics parameters, see <a href="#">par</a>

### Details

For cross validation the data are split into a number of segments, PCA is computed (using 1 to amax components) for all but one segment, and the scores of the segment left out are calculated. This is done in turn, by omitting each segment one time. Thus, a complete score matrix results for each desired number of components, and the error matrices of fit can be computed. A measure of fit is the explained variance, which is computed for each number of components. Then the whole procedure is repeated (repl times), which results in repl numbers of explained variance for 1 to amax components, i.e. a matrix. The matrix is presented by boxplots, where each boxplot summarized the explained variance for a certain number of principal components.

### Value

ExplVar	matrix with explained variances, repl rows, and amax columns
MSEP	matrix with MSEP values, repl rows, and amax columns

### Author(s)

Peter Filzmoser <P.Filzmoser@tuwien.ac.at>

### References

K. Varmuza and P. Filzmoser: Introduction to Multivariate Statistical Analysis in Chemometrics. CRC Press, Boca Raton, FL, 2009.

### See Also

[princomp](#)

### Examples

```
data(glass)
x.sc <- scale(glass)
resv <- clvalidity(x.sc,clnumb=c(2:5))
```

---

pcaDiagplot

*Diagnostics plot for PCA*

---

### Description

Score distances and orthogonal distances are computed and plotted.

### Usage

```
pcaDiagplot(X, X.pca, a = 2, quantile = 0.975, scale = TRUE, plot = TRUE, ...)
```

**Arguments**

X	numeric data frame or matrix
X.pca	PCA object resulting e.g. from <a href="#">princomp</a>
a	number of principal components
quantile	quantile for the critical cut-off values
scale	if TRUE then X will be scaled - and X.pca should be from scaled data too
plot	if TRUE a plot is generated
...	additional graphics parameters, see <a href="#">par</a>

**Details**

The score distance measures the outlyingness of the objects within the PCA space using Mahalanobis distances. The orthogonal distance measures the distance of the objects orthogonal to the PCA space. Cut-off values for both distance measures help to distinguish between outliers and regular observations.

**Value**

SDist	Score distances
ODist	Orthogonal distances
critSD	critical cut-off value for the score distances
critOD	critical cut-off value for the orthogonal distances

**Author(s)**

Peter Filzmoser <P.Filzmoser@tuwien.ac.at>

**References**

K. Varmuza and P. Filzmoser: Introduction to Multivariate Statistical Analysis in Chemometrics. CRC Press, Boca Raton, FL, 2009.

**See Also**

[princomp](#)

**Examples**

```
data(glass)
require(robustbase)
glass.mcd <- covMcd(glass)
rpca <- princomp(glass, covmat=glass.mcd)
res <- pcaDiagplot(glass, rpca, a=2)
```

---

pcaVarexpl                      *PCA diagnostics for variables*

---

**Description**

Diagnostics of PCA to see the explained variance for each variable.

**Usage**

```
pcaVarexpl(X, a, center = TRUE, scale = TRUE, plot = TRUE, ...)
```

**Arguments**

X	numeric data frame or matrix
a	number of principal components
center	centring of X (FALSE or TRUE)
scale	scaling of X (FALSE or TRUE)
plot	if TRUE make plot with explained variance
...	additional graphics parameters, see <a href="#">par</a>

**Details**

For a desired number of principal components the percentage of explained variance is computed for each variable and plotted.

**Value**

ExplVar                      explained variance for each variable

**Author(s)**

Peter Filzmoser <P.Filzmoser@tuwien.ac.at>

**References**

K. Varmuza and P. Filzmoser: Introduction to Multivariate Statistical Analysis in Chemometrics. CRC Press, Boca Raton, FL, 2009.

**See Also**

[princomp](#)

**Examples**

```
data(glass)
res <- pcaVarexpl(glass,a=2)
```

---

Phenyl

*Phenyl data set*

---

### **Description**

The data consist of mass spectra from 600 chemical compounds, where 300 contain a phenyl substructure (group 1) and 300 compounds do not contain this substructure (group 2). The mass spectra have been transformed to 658 variables, containing the mass spectral features. The 2 groups are coded as -1 (group 1) and +1 (group 2), and is provided as first last variable.

### **Usage**

```
data(Phenyl)
```

### **Format**

A data frame with 600 observations on the following 659 variables.

grp a numeric vector

spec.V1 a numeric vector

spec.V2 a numeric vector

spec.V3 a numeric vector

spec.V4 a numeric vector

spec.V5 a numeric vector

spec.V6 a numeric vector

spec.V7 a numeric vector

spec.V8 a numeric vector

spec.V9 a numeric vector

spec.V10 a numeric vector

spec.V11 a numeric vector

spec.V12 a numeric vector

spec.V13 a numeric vector

spec.V14 a numeric vector

spec.V15 a numeric vector

spec.V16 a numeric vector

spec.V17 a numeric vector

spec.V18 a numeric vector

spec.V19 a numeric vector

spec.V20 a numeric vector

spec.V21 a numeric vector

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### Details

The data set can be used for classification in high dimensions.

### Source

K. Varmuza and P. Filzmoser: Introduction to Multivariate Statistical Analysis in Chemometrics. CRC Press, Boca Raton, FL, 2009.

### References

K. Varmuza and P. Filzmoser: Introduction to Multivariate Statistical Analysis in Chemometrics. CRC Press, Boca Raton, FL, 2009.

### Examples

```
data(Phenyl)
str(Phenyl)
```

---

plotcompmvr

*Component plot for repeated DCV*

---

### Description

Generate plot showing optimal number of components for Repeated Double Cross-Validation

### Usage

```
plotcompmvr(mvr_dcvobj, ...)
```

### Arguments

mvr\_dcvobj      object from repeated double-CV, see [mvr\\_dcv](#)  
...              additional plot arguments

### Details

After running repeated double-CV, this plot helps to decide on the final number of components.

**Value**

optcomp            optimal number of components  
 compdistrib        frequencies for the optimal number of components

**Author(s)**

Peter Filzmoser <P.Filzmoser@tuwien.ac.at>

**References**

K. Varmuza and P. Filzmoser: Introduction to Multivariate Statistical Analysis in Chemometrics. CRC Press, Boca Raton, FL, 2009.

**See Also**

[mvr](#)

**Examples**

```
data(NIR)
X <- NIR$xNIR[1:30,]        # first 30 observations - for illustration
y <- NIR$yGlcEtOH[1:30,1] # only variable Glucose
NIR.Glc <- data.frame(X=X, y=y)
res <- mvr_dcv(y~., data=NIR.Glc, ncomp=10, method="simpls", repl=10)
plot2 <- plotcompmvr(res)
```

---

plotcompprm

*Component plot for repeated DCV of PRM*

---

**Description**

Generate plot showing optimal number of components for Repeated Double Cross-Validation of Partial Robust M-regression

**Usage**

```
plotcompprm(prmdcvobj, ...)
```

**Arguments**

prmdcvobj            object from repeated double-CV of PRM, see [prmdcv](#)  
 ...                    additional plot arguments

**Details**

After running repeated double-CV for PRM, this plot helps to decide on the final number of components.

**Value**

optcomp            optimal number of components  
 compdistrib       frequencies for the optimal number of components

**Author(s)**

Peter Filzmoser <P.Filzmoser@tuwien.ac.at>

**References**

K. Varmuza and P. Filzmoser: Introduction to Multivariate Statistical Analysis in Chemometrics. CRC Press, Boca Raton, FL, 2009.

**See Also**

[prm](#)

**Examples**

```
data(NIR)
X <- NIR$xNIR[1:30,]        # first 30 observations - for illustration
y <- NIR$yGlcEtOH[1:30,1] # only variable Glucose
NIR.Glc <- data.frame(X=X, y=y)
res <- prm_dcv(X,y,a=4,repl=2)
plot2 <- plotcompprm(res)
```

---

plotpredmvr

*Plot predictions from repeated DCV*

---

**Description**

Generate plot showing predicted values for Repeated Double Cross Validation

**Usage**

```
plotpredmvr(mvr_dcvobj, optcomp, y, X, method = "simpls", ...)
```

**Arguments**

mvr\_dcvobj        object from repeated double-CV, see [mvr\\_dcv](#)  
 optcomp           optimal number of components  
 y                 data from response variable  
 X                 data with explanatory variables  
 method           the multivariate regression method to be used, see [mvr](#)  
 ...               additional plot arguments

**Details**

After running repeated double-CV, this plot visualizes the predicted values.

**Value**

A plot is generated.

**Author(s)**

Peter Filzmoser <P.Filzmoser@tuwien.ac.at>

**References**

K. Varmuza and P. Filzmoser: Introduction to Multivariate Statistical Analysis in Chemometrics. CRC Press, Boca Raton, FL, 2009.

**See Also**

[mvr](#)

**Examples**

```
data(NIR)
X <- NIR$xNIR[1:30,]      # first 30 observations - for illustration
y <- NIR$yGlcEtOH[1:30,1] # only variable Glucose
NIR.Glc <- data.frame(X=X, y=y)
res <- mvr_dcv(y~,data=NIR.Glc,ncomp=10,method="simpls",repl=10)
plot3 <- plotpredmvr(res,opt=7,y,X,method="simpls")
```

---

plotpredprm

*Plot predictions from repeated DCV of PRM*

---

**Description**

Generate plot showing predicted values for Repeated Double Cross Validation of Partial Robust M-regression

**Usage**

```
plotpredprm(prmdcvobj, optcomp, y, X, ...)
```

**Arguments**

prmdcvobj	object from repeated double-CV of PRM, see <a href="#">prmdcv</a>
optcomp	optimal number of components
y	data from response variable
X	data with explanatory variables
...	additional plot arguments

**Details**

After running repeated double-CV for PRM, this plot visualizes the predicted values. The result is compared with predicted values obtained via usual CV of PRM.

**Value**

A plot is generated.

**Author(s)**

Peter Filzmoser <P.Filzmoser@tuwien.ac.at>

**References**

K. Varmuza and P. Filzmoser: Introduction to Multivariate Statistical Analysis in Chemometrics. CRC Press, Boca Raton, FL, 2009.

**See Also**

[prm](#)

**Examples**

```
data(NIR)
X <- NIR$xNIR[1:30,]      # first 30 observations - for illustration
y <- NIR$yGlcEtOH[1:30,1] # only variable Glucose
NIR.Glc <- data.frame(X=X, y=y)
res <- prm_dcv(X,y,a=4,repl=2)
plot3 <- plotpredprm(res,opt=res$afinal,y,X)
```

---

plotprm

*Plot results from robust PLS*

---

**Description**

The predicted values and the residuals are shown for robust PLS using the optimal number of components.

**Usage**

```
plotprm(prmobj, y, ...)
```

**Arguments**

prmobj	resulting object from CV of robust PLS, see <a href="#">prm_cv</a>
y	vector with values of response variable
...	additional plot arguments

**Details**

Robust PLS based on partial robust M-regression is available at [prm](#). Here the function `prm_cv` has to be used first, applying cross-validation with robust PLS. Then the result is taken by this routine and two plots are generated for the optimal number of PLS components: The measured versus the predicted y, and the predicted y versus the residuals.

**Value**

A plot is generated.

**Author(s)**

Peter Filzmoser <P.Filzmoser@tuwien.ac.at>

**References**

K. Varmuza and P. Filzmoser: Introduction to Multivariate Statistical Analysis in Chemometrics. CRC Press, Boca Raton, FL, 2009.

**See Also**

[prm](#)

**Examples**

```
data(cereal)
set.seed(123)
res <- prm_cv(cereal$X, cereal$Y[, 1], a=5, segments=4, plot.opt=FALSE)
plotprm(res, cereal$Y[, 1])
```

---

plotresmvr

*Plot residuals from repeated DCV*

---

**Description**

Generate plot showing residuals for Repeated Double Cross Validation

**Usage**

```
plotresmvr(mvrdcvobj, optcomp, y, X, method = "simpls", ...)
```

**Arguments**

<code>mvrdcvobj</code>	object from repeated double-CV, see <a href="#">mvr_dcv</a>
<code>optcomp</code>	optimal number of components
<code>y</code>	data from response variable
<code>X</code>	data with explanatory variables
<code>method</code>	the multivariate regression method to be used, see <a href="#">mvr</a>
<code>...</code>	additional plot arguments

**Details**

After running repeated double-CV, this plot visualizes the residuals.

**Value**

A plot is generated.

**Author(s)**

Peter Filzmoser <P.Filzmoser@tuwien.ac.at>

**References**

K. Varmuza and P. Filzmoser: Introduction to Multivariate Statistical Analysis in Chemometrics. CRC Press, Boca Raton, FL, 2009.

**See Also**

[mvr](#)

**Examples**

```
data(NIR)
X <- NIR$xNIR[1:30,] # first 30 observations - for illustration
y <- NIR$yGlcEtOH[1:30,1] # only variable Glucose
NIR.Glc <- data.frame(X=X, y=y)
res <- mvr_dcv(y~, data=NIR.Glc, ncomp=10, method="simpls", repl=10)
plot4 <- plotresmvr(res, opt=7, y, X, method="simpls")
```

---

plotresprm

*Plot residuals from repeated DCV of PRM*

---

**Description**

Generate plot showing residuals for Repeated Double Cross Validation for Partial Robust M-regression

**Usage**

```
plotresprm(prm_dcvobj, optcomp, y, X, ...)
```

**Arguments**

prm_dcvobj	object from repeated double-CV of PRM, see <a href="#">prm_dcv</a>
optcomp	optimal number of components
y	data from response variable
X	data with explanatory variables
...	additional plot arguments

**Details**

After running repeated double-CV for PRM, this plot visualizes the residuals. The result is compared with predicted values obtained via usual CV of PRM.

**Value**

A plot is generated.

**Author(s)**

Peter Filzmoser <P.Filzmoser@tuwien.ac.at>

**References**

K. Varmuza and P. Filzmoser: Introduction to Multivariate Statistical Analysis in Chemometrics. CRC Press, Boca Raton, FL, 2009.

**See Also**

[prm](#)

**Examples**

```
data(NIR)
X <- NIR$XNIR[1:30,]      # first 30 observations - for illustration
y <- NIR$yGlcEtOH[1:30,1] # only variable Glucose
NIR.Glc <- data.frame(X=X, y=y)
res <- prm_dcv(X,y,a=4,repl=2)
plot4 <- plotresprm(res,opt=res$afinal,y,X)
```

---

plotRidge

*Plot results of Ridge regression*

---

**Description**

Two plots from Ridge regression are generated: The MSE resulting from Generalized Cross Validation (GCV) versus the Ridge parameter lambda, and the regression coefficients versus lambda. The optimal choice for lambda is indicated.

**Usage**

```
plotRidge(formula, data, lambda = seq(0.5, 50, by = 0.05), ...)
```

**Arguments**

formula	formula, like $y \sim X$ , i.e., dependent~response variables
data	data frame to be analyzed
lambda	possible values for the Ridge parameter to evaluate
...	additional plot arguments

**Details**

For all values provided in lambda the results for Ridge regression are computed. The function [lm.ridge](#) is used for cross-validation and Ridge regression.

**Value**

predicted	predicted values for the optimal lambda
lambdaopt	optimal Ridge parameter lambda from GCV

**Author(s)**

Peter Filzmoser <P.Filzmoser@tuwien.ac.at>

**References**

K. Varmuza and P. Filzmoser: Introduction to Multivariate Statistical Analysis in Chemometrics. CRC Press, Boca Raton, FL, 2009.

**See Also**

[lm.ridge](#), [plotRidge](#)

**Examples**

```
data(PAC)
res=plotRidge(y~X,data=PAC,lambda=seq(1,20,by=0.5))
```

---

plotSEPMvr

*Plot SEP from repeated DCV*

---

**Description**

Generate plot showing SEP values for Repeated Double Cross Validation

**Usage**

```
plotSEPMvr(mvrdcvobj, optcomp, y, X, method = "simpls", complete = TRUE, ...)
```

**Arguments**

mvrdcvobj	object from repeated double-CV, see <a href="#">mvr_dcv</a>
optcomp	optimal number of components
y	data from response variable
X	data with explanatory variables
method	the multivariate regression method to be used, see <a href="#">mvr</a>

complete	if TRUE the SEPcv values are drawn and computed for the same range of components as included in the mvr_dcvobj object; if FALSE only optcomp components are computed and their results are displayed
...	additional plot arguments

### Details

After running repeated double-CV, this plot visualizes the distribution of the SEP values.

### Value

SEPdcv	all SEP values from repeated double-CV
SEPcv	SEP values from classical CV

### Author(s)

Peter Filzmoser <P.Filzmoser@tuwien.ac.at>

### References

K. Varmuza and P. Filzmoser: Introduction to Multivariate Statistical Analysis in Chemometrics. CRC Press, Boca Raton, FL, 2009.

### See Also

[mvr](#)

### Examples

```
data(NIR)
X <- NIR$xNIR[1:30,] # first 30 observations - for illustration
y <- NIR$yGlcEtOH[1:30,1] # only variable Glucose
NIR.Glc <- data.frame(X=X, y=y)
res <- mvr_dcv(y~, data=NIR.Glc, ncomp=10, method="simpls", repl=10)
plot1 <- plotSEPmvr(res, opt=7, y, X, method="simpls")
```

---

plotSEPprm

*Plot trimmed SEP from repeated DCV of PRM*

---

### Description

Generate plot showing trimmed SEP values for Repeated Double Cross Validation for Partial Robust M-Regression (PRM)

### Usage

```
plotSEPprm(prm_dcvobj, optcomp, y, X, complete = TRUE, ...)
```

**Arguments**

prmdcvobj	object from repeated double-CV of PRM, see <a href="#">prmdcv</a>
optcomp	optimal number of components
y	data from response variable
X	data with explanatory variables
complete	if TRUE the trimmed SEPcv values are drawn and computed from <a href="#">prmdcv</a> for the same range of components as included in the prmdcvobj object; if FALSE only optcomp components are computed and their results are displayed
...	additional arguments of <a href="#">prmdcv</a>

**Details**

After running repeated double-CV for PRM, this plot visualizes the distribution of the SEP values. While the gray lines represent the resulting trimmed SEP values from repeated double CV, the black line is the result for standard CV with PRM, and it is usually too optimistic.

**Value**

SEPdcv	all trimmed SEP values from repeated double-CV
SEPcv	trimmed SEP values from usual CV

**Author(s)**

Peter Filzmoser <P.Filzmoser@tuwien.ac.at>

**References**

K. Varmuza and P. Filzmoser: Introduction to Multivariate Statistical Analysis in Chemometrics. CRC Press, Boca Raton, FL, 2009.

**See Also**

[prmdcv](#)

**Examples**

```
data(NIR)
X <- NIR$xNIR[1:30,]      # first 30 observations - for illustration
y <- NIR$yGlcEtOH[1:30,1] # only variable Glucose
NIR.Glc <- data.frame(X=X, y=y)
res <- prmdcv(X,y,a=4,repl=2)
plot1 <- plotSEPprm(res,opt=res$afinal,y,X)
```

---

plotsom

*Plot SOM results*

---

## Description

Plot results of Self Organizing Maps (SOM).

## Usage

```
plotsom(obj, grp, type = c("num", "bar"), margins = c(3,2,2,2), ...)
```

## Arguments

obj	result object from <a href="#">som</a>
grp	numeric vector or factor with group information
type	type of presentation for output, see details
margins	plot margins for output, see <a href="#">par</a>
...	additional graphics parameters, see <a href="#">par</a>

## Details

The results of Self Organizing Maps (SOM) are plotted either in a table with numbers (type="num") or with barplots (type="bar"). There is a limitation to at most 9 groups. A summary table is returned.

## Value

sumtab	Summary table
--------	---------------

## Author(s)

Peter Filzmoser <P.Filzmoser@tuwien.ac.at>

## References

K. Varmuza and P. Filzmoser: Introduction to Multivariate Statistical Analysis in Chemometrics. CRC Press, Boca Raton, FL, 2009.

## See Also

[som](#)

**Examples**

```

data(glass)
require(som)
Xs <- scale(glass)
Xn <- Xs/sqrt(apply(Xs^2,1,sum))
X_SOM <- som(Xn,xdim=4,ydim=4) # 4x4 fields
data(glass.grp)
res <- plotsom(X_SOM,glass.grp,type="bar")

```

---

pls1\_nipals

*PLS1 by NIPALS*


---

**Description**

NIPALS algorithm for PLS1 regression (y is univariate)

**Usage**

```
pls1_nipals(X, y, a, it = 50, tol = 1e-08, scale = FALSE)
```

**Arguments**

X	original X data matrix
y	original y-data
a	number of PLS components
it	number of iterations
tol	tolerance for convergence
scale	if TRUE the X and y data will be scaled in addition to centering, if FALSE only mean centering is performed

**Details**

The NIPALS algorithm is the originally proposed algorithm for PLS. Here, the y-data are only allowed to be univariate. This simplifies the algorithm.

**Value**

P	matrix with loadings for X
T	matrix with scores for X
W	weights for X
C	weights for Y
b	final regression coefficients

**Author(s)**

Peter Filzmoser <P.Filzmoser@tuwien.ac.at>

**References**

K. Varmuza and P. Filzmoser: Introduction to Multivariate Statistical Analysis in Chemometrics. CRC Press, Boca Raton, FL, 2009.

**See Also**

[mvr](#), [pls2\\_nipals](#)

**Examples**

```
data(PAC)
res <- pls1_nipals(PAC$X,PAC$y,a=5)
```

---

pls2\_nipals

*PLS2 by NIPALS*

---

**Description**

NIPALS algorithm for PLS2 regression (y is multivariate)

**Usage**

```
pls2_nipals(X, Y, a, it = 50, tol = 1e-08, scale = FALSE)
```

**Arguments**

X	original X data matrix
Y	original Y-data matrix
a	number of PLS components
it	number of iterations
tol	tolerance for convergence
scale	if TRUE the X and y data will be scaled in addition to centering, if FALSE only mean centering is performed

**Details**

The NIPALS algorithm is the originally proposed algorithm for PLS. Here, the Y-data matrix is multivariate.

**Value**

P	matrix with loadings for X
T	matrix with scores for X
Q	matrix with loadings for Y
U	matrix with scores for Y
D	D-matrix within the algorithm
W	weights for X
C	weights for Y
B	final regression coefficients

**Author(s)**

Peter Filzmoser <P.Filzmoser@tuwien.ac.at>

**References**

K. Varmuza and P. Filzmoser: Introduction to Multivariate Statistical Analysis in Chemometrics. CRC Press, Boca Raton, FL, 2009.

**See Also**

[mvr](#), [pls1\\_nipals](#)

**Examples**

```
data(cereal)
res <- pls2_nipals(cereal$X, cereal$Y, a=5)
```

---

pls\_eigen

*Eigenvector algorithm for PLS*

---

**Description**

Computes the PLS solution by eigenvector decompositions.

**Usage**

```
pls_eigen(X, Y, a)
```

**Arguments**

X	X input data, centered (and scaled)
Y	Y input data, centered (and scaled)
a	number of PLS components

**Details**

The X loadings (P) and scores (T) are found by the eigendecomposition of  $X'YY'X$ . The Y loadings (Q) and scores (U) come from the eigendecomposition of  $Y'XX'Y$ . The resulting P and Q are orthogonal. The first score vectors are the same as for standard PLS, subsequent score vectors different.

**Value**

P	matrix with loadings for X
T	matrix with scores for X
Q	matrix with loadings for Y
U	matrix with scores for Y

**Author(s)**

Peter Filzmoser <P.Filzmoser@tuwien.ac.at>

**References**

K. Varmuza and P. Filzmoser: Introduction to Multivariate Statistical Analysis in Chemometrics. CRC Press, Boca Raton, FL, 2009.

**See Also**

[mvr](#)

**Examples**

```
data(cereal)
res <- pls_eigen(cereal$X,cereal$Y,a=5)
```

---

prm

*Robust PLS*

---

**Description**

Robust PLS by partial robust M-regression.

**Usage**

```
prm(X, y, a, fairct = 4, opt = "l1m",usesvd=FALSE)
```

**Arguments**

X	predictor matrix
y	response variable
a	number of PLS components
fairct	tuning constant, by default fairct=4
opt	if "l1m" the mean centering is done by the l1-median, otherwise if "median" the coordinate-wise median is taken
usesvd	if TRUE, SVD will be used if X has more columns than rows

**Details**

M-regression is used to robustify PLS, with initial weights based on the FAIR weight function.

**Value**

coef	vector with regression coefficients
intercept	coefficient for intercept
wy	vector of length(y) with residual weights
wt	vector of length(y) with weights for leverage
w	overall weights
scores	matrix with PLS X-scores
loadings	matrix with PLS X-loadings
fitted.values	vector with fitted y-values
mx	column means of X
my	mean of y

**Author(s)**

Peter Filzmoser <P.Filzmoser@tuwien.ac.at>

**References**

S. Serneels, C. Croux, P. Filzmoser, and P.J. Van Espen. Partial robust M-regression. *Chemometrics and Intelligent Laboratory Systems*, Vol. 79(1-2), pp. 55-64, 2005.

**See Also**

[mvr](#)

**Examples**

```
data(PAC)
res <- prn(PAC$X, PAC$y, a=5)
```

pru\_cv

*Cross-validation for robust PLS***Description**

Cross-validation (CV) is carried out with robust PLS based on partial robust M-regression. A plot with the choice for the optimal number of components is generated. This only works for univariate y-data.

**Usage**

```
pru_cv(X, y, a, fairct = 4, opt = "median", subset = NULL, segments = 10,
       segment.type = "random", trim = 0.2, sdfact = 2, plot.opt = TRUE)
```

**Arguments**

X	predictor matrix
y	response variable
a	number of PLS components
fairct	tuning constant, by default fairct=4
opt	if "l1m" the mean centering is done by the l1-median, otherwise by the coordinate-wise median
subset	optional vector defining a subset of objects
segments	the number of segments to use or a list with segments (see <a href="#">mvrCv</a> )
segment.type	the type of segments to use. Ignored if 'segments' is a list
trim	trimming percentage for the computation of the SEP
sdfact	factor for the multiplication of the standard deviation for the determination of the optimal number of components, see <a href="#">mvr_dcv</a>
plot.opt	if TRUE a plot will be generated that shows the selection of the optimal number of components for each step of the CV, see <a href="#">mvr_dcv</a>

**Details**

A function for robust PLS based on partial robust M-regression is available at [pru](#). The optimal number of robust PLS components is chosen according to the following criterion: Within the CV scheme, the mean of the trimmed SEPs  $SEP_{trimave}$  is computed for each number of components, as well as their standard errors  $SEP_{trimse}$ . Then one searches for the minimum of the  $SEP_{trimave}$  values and adds  $sdfact * SEP_{trimse}$ . The optimal number of components is the most parsimonious model that is below this bound.

**Value**

predicted	matrix with length(y) rows and a columns with predicted values
SEPall	vector of length a with SEP values for each number of components
SEPtrim	vector of length a with trimmed SEP values for each number of components
SEPj	matrix with segments rows and a columns with SEP values within the CV for each number of components
SEPtrimj	matrix with segments rows and a columns with trimmed SEP values within the CV for each number of components
optcomp	final optimal number of PLS components
SEPOpt	trimmed SEP value for final optimal number of PLS components

**Author(s)**

Peter Filzmoser <P.Filzmoser@tuwien.ac.at>

**References**

K. Varmuza and P. Filzmoser: Introduction to Multivariate Statistical Analysis in Chemometrics. CRC Press, Boca Raton, FL, 2009.

**See Also**

[prm](#)

**Examples**

```
data(cereal)
set.seed(123)
res <- prm_cv(cereal$X, cereal$Y[,1], a=5, segments=4, plot.opt=TRUE)
```

---

prm\_dcv

*Repeated double-cross-validation for robust PLS*

---

**Description**

Performs a careful evaluation by repeated double-CV for robust PLS, called PRM (partial robust M-estimation).

**Usage**

```
prm_dcv(X, Y, a=10, repl=10, segments0=4, segments=7, segment0.type="random",
  segment.type="random", sdfact=2, fairct=4, trim=0.2, opt="median", plot.opt=FALSE, ...)
```

**Arguments**

X	predictor matrix
Y	response variable
a	number of PLS components
repl	Number of replication for the double-CV
segments0	the number of segments to use for splitting into training and test data, or a list with segments (see <a href="#">mvrCv</a> )
segments	the number of segments to use for selecting the optimal number of components, or a list with segments (see <a href="#">mvrCv</a> )
segment0.type	the type of segments to use. Ignored if 'segments0' is a list
segment.type	the type of segments to use. Ignored if 'segments' is a list
sdfact	factor for the multiplication of the standard deviation for the determination of the optimal number of components, see <a href="#">mvr_dcv</a>
fairct	tuning constant, by default fairct=4
trim	trimming percentage for the computation of the SEP
opt	if "l1m" the mean centering is done by the l1-median, otherwise if "median", by the coordinate-wise median
plot.opt	if TRUE a plot will be generated that shows the selection of the optimal number of components for each step of the CV
...	additional parameters

**Details**

In this cross-validation (CV) scheme, the optimal number of components is determined by an additional CV in the training set, and applied to the test set. The procedure is repeated `repl` times. The optimal number of components is the model with the smallest number of components which is still in the range of the  $MSE + sdfact * sd(MSE)$ , where `MSE` and `sd` are taken from the minimum.

**Value**

b	estimated regression coefficients
intercept	estimated regression intercept
resopt	array [nrow(Y) x ncol(Y) x repl] with residuals using optimum number of components
predopt	array [nrow(Y) x ncol(Y) x repl] with predicted Y using optimum number of components
optcomp	matrix [segments0 x repl] optimum number of components for each training set
residcomp	array [nrow(Y) x ncomp x repl] with residuals using optimum number of components
pred	array [nrow(Y) x ncol(Y) x ncomp x repl] with predicted Y for all numbers of components
SEPal1	matrix [ncomp x repl] with SEP values

SEPtrim	matrix [ncomp x repl] with trimmed SEP values
SEPcomp	vector of length ncomp with trimmed SEP values; use the element afinal for the optimal trimmed SEP
afinal	final optimal number of components
SEPOpt	trimmed SEP over all residuals using optimal number of components

**Author(s)**

Peter Filzmoser <P.Filzmoser@tuwien.ac.at>

**References**

K. Varmuza and P. Filzmoser: Introduction to Multivariate Statistical Analysis in Chemometrics. CRC Press, Boca Raton, FL, 2009.

**See Also**

[mvr](#)

**Examples**

```
data(NIR)
X <- NIR$xNIR[1:30,]      # first 30 observations - for illustration
y <- NIR$yGlcEtOH[1:30,1] # only variable Glucose
NIR.Glc <- data.frame(X=X, y=y)
res <- prm_dcv(X,y,a=3,repl=2)
```

---

ridgeCV

*Repeated CV for Ridge regression*

---

**Description**

Performs repeated cross-validation (CV) to evaluate the result of Ridge regression where the optimal Ridge parameter lambda was chosen on a fast evaluation scheme.

**Usage**

```
ridgeCV(formula, data, lambdaopt, repl = 5, segments = 10,
        segment.type = c("random", "consecutive", "interleaved"), length.seg,
        trace = FALSE, plot.opt = TRUE, ...)
```

**Arguments**

formula	formula, like $y \sim X$ , i.e., dependent~response variables
data	data frame to be analyzed
lambdaopt	optimal Ridge parameter lambda
repl	number of replications for the CV
segments	the number of segments to use for CV, or a list with segments (see <a href="#">mvrCv</a> )
segment.type	the type of segments to use. Ignored if 'segments' is a list
length.seg	Positive integer. The length of the segments to use. If specified, it overrides 'segments' unless 'segments' is a list
trace	logical; if 'TRUE', the segment number is printed for each segment
plot.opt	if TRUE a plot will be generated that shows the predicted versus the observed y-values
...	additional plot arguments

**Details**

Generalized Cross Validation (GCV) is used by the function [lm.ridge](#) to get a quick answer for the optimal Ridge parameter. This function should make a careful evaluation once the optimal parameter lambda has been selected. Measures for the prediction quality are computed and optionally plots are shown.

**Value**

residuals	matrix of size $\text{length}(y) \times \text{repl}$ with residuals
predicted	matrix of size $\text{length}(y) \times \text{repl}$ with predicted values
SEP	Standard Error of Prediction computed for each column of "residuals"
SEPM	mean SEP value
sMAD	MAD of Prediction computed for each column of "residuals"
sMADm	mean of MAD values
RMSEP	Root MSEP value computed for each column of "residuals"
RMSEPM	mean RMSEP value

**Author(s)**

Peter Filzmoser <[P.Filzmoser@tuwien.ac.at](mailto:P.Filzmoser@tuwien.ac.at)>

**References**

K. Varmuza and P. Filzmoser: Introduction to Multivariate Statistical Analysis in Chemometrics. CRC Press, Boca Raton, FL, 2009.

**See Also**

[lm.ridge](#), [plotRidge](#)

**Examples**

```
data(PAC)
res=ridgeCV(y~X,data=PAC,lambdaopt=4.3,repl=5,segments=5)
```

RPvectors

*Generating random projection directions***Description**

A matrix with pandom projection (RP) directions (columns) is generated according to a chosen distributions; optionally the random vectors are orthogonalized.

**Usage**

```
RPvectors(a, m, ortho = "none", distr = "uniform", par_unif = c(-1, 1),
par_norm = c(0, 1), par_eq = c(-1, 0, 1), par_uneq = c(-sqrt(3), 0, sqrt(3)),
par_uneqprob = c(1/6, 2/3, 1/6))
```

**Arguments**

a	number of generated vectors ( $\geq 1$ )
m	dimension of generated vectors ( $\geq 2$ )
ortho	orthogonalization of vectors: "none" ... no orthogonalization (default); "onfly" ... orthogonalization on the fly after each generated vector; "end" ... orthogonalization at the end, after the whole random matrix was generated
distr	distribution of generated random vector components: "uniform" ... uniformly distributed in range par_unif (see below); default U[-1, +1]; "normal" ... normally distributed with parameters par_norm (see below); typical N(0, 1); "randeq" ... random selection of values par_eq (see below) with equal probabilities; typically -1, 0, +1; "randuneq" ... random selection of values par_uneq (see below) with probabilties par_uneqprob (see below); typical $-(3)^{0.5}$ with probability 1/6; 0 with probability 2/3; $+(3)^{0.5}$ with probability 1/6
par_unif	parameters for range for distr=="uniform"; default to c(-1,1)
par_norm	parameters for mean and sdev for distr=="normal"; default to c(0,1)
par_eq	values for distr=="randeq" which are replicated; default to c(-1,0,1)
par_uneq	values for distr=="randuneq" which are replicated with probabilties par_uneqprob; default to c(-sqrt(3),0,sqrt(3))
par_uneqprob	probabilities for distr=="randuneq" to replicate values par_uneq; default to c(1/6,2/3,1/6)

**Details**

The generated random projections can be used for dimension reduction of multivariate data. Suppose we have a data matrix  $X$  with  $n$  rows and  $m$  columns. Then the call `B <- RPvectors(a,m)` will produce a matrix  $B$  with the random directions in its columns. The matrix product  $X$  times  $t(B)$  results in a matrix of lower dimension  $a$ . There are several options to generate the projection directions, like orthogonal directions, and different distributions with different parameters to generate the random numbers. Random Projection (RP) can have comparable performance for dimension reduction like PCA, but gives a big advantage in terms of computation time.

**Value**

The value returned is the matrix  $B$  with  $a$  columns of length  $m$ , representing the random vectors

**Author(s)**

Peter Filzmoser <P.Filzmoser@tuwien.ac.at>

**References**

K. Varmuza, P. Filzmoser, and B. Liebmann. Random projection experiments with chemometric data. *Journal of Chemometrics*. To appear.

**Examples**

```
B <- RPvectors(a=5,m=10)
res <- t(B)
```

---

sd\_trim

*Trimmed standard deviation*

---

**Description**

The trimmed standard deviation as a robust estimator of scale is computed.

**Usage**

```
sd_trim(x, trim=0.2, const=TRUE)
```

**Arguments**

<code>x</code>	numeric vector, data frame or matrix
<code>trim</code>	trimming proportion; should be between 0 and 0.5
<code>const</code>	if TRUE, the appropriate consistency correction is done

**Details**

The trimmed standard deviation is defined as the average trimmed sum of squared deviations around the trimmed mean. A consistency factor for normal distribution is included. However, this factor is only available now for trim equal to 0.1 or 0.2. For different trimming percentages the appropriate constant needs to be used. If the input is a data matrix, the trimmed standard deviation of the columns is computed.

**Value**

Returns the trimmed standard deviations of the vector  $x$ , or in case of a matrix, of the columns of  $x$ .

**Author(s)**

Peter Filzmoser <P.Filzmoser@tuwien.ac.at>

**References**

K. Varmuza and P. Filzmoser: Introduction to Multivariate Statistical Analysis in Chemometrics. CRC Press, Boca Raton, FL, 2009.

**See Also**

[sd,mean](#)

**Examples**

```
x <- c(rnorm(100),100) # outlier 100 is included
sd(x) # classical standard deviation
sd_trim(x) # trimmed standard deviation
```

---

stepwise

*Stepwise regression*

---

**Description**

Stepwise regression, starting from the empty model, with scope to the full model

**Usage**

```
stepwise(formula, data, k, startM, maxTime = 1800, direction = "both",
writeFile = FALSE, maxsteps = 500, ...)
```

**Arguments**

formula	formula, like $y \sim X$ , i.e., dependent~response variables
data	data frame to be analyzed
k	sensible values are $\log(\text{nrow}(x))$ for BIC or 2 for AIC; if not provided -> BIC
startM	optional, the starting model; provide a binary vector
maxTime	maximal time to be used for algorithm
direction	either "forward" or "backward" or "both"
writeFile	if TRUE results are shown on the screen
maxsteps	maximum number of steps
...	additional plot arguments

**Details**

This function is similar to the function [step](#) for stepwise regression. It is especially designed for cases where the number of regressor variables is much higher than the number of objects. The formula for the full model (scope) is automatically generated.

**Value**

usedTime	time that has been used for algorithm
bic	BIC values for different models
models	matrix with no. of models rows and no. of variables columns, and 0/1 entries defining the models

**Author(s)**

Leonhard Seyfang and (marginally) Peter Filzmoser <P.Filzmoser@tuwien.ac.at>

**References**

K. Varmuza and P. Filzmoser: Introduction to Multivariate Statistical Analysis in Chemometrics. CRC Press, Boca Raton, FL, 2009.

**See Also**

[step](#)

**Examples**

```
data(NIR)
X <- NIR$XNIR[1:30,]      # first 30 observations - for illustration
y <- NIR$yGlcEtOH[1:30,1] # only variable Glucose
NIR.Glc <- data.frame(X=X, y=y)
res <- stepwise(y~., data=NIR.Glc, maxsteps=2)
```

svmEval

*Support Vector Machine evaluation by CV***Description**

Evaluation for Support Vector Machines (SVM) by cross-validation

**Usage**

```
svmEval(X, grp, train, kfold = 10, gamvec = seq(0, 10, by = 1), kernel = "radial",
degree = 3, plotit = TRUE, legend = TRUE, legpos = "bottomright", ...)
```

**Arguments**

X	standardized complete X data matrix (training and test data)
grp	factor with groups for complete data (training and test data)
train	row indices of X indicating training data objects
kfold	number of folds for cross-validation
gamvec	range for gamma-values, see <a href="#">svm</a>
kernel	kernel to be used for SVM, should be one of "radial", "linear", "polynomial", "sigmoid", default to "radial", see <a href="#">svm</a>
degree	degree of polynome if kernel is "polynomial", default to 3, see <a href="#">svm</a>
plotit	if TRUE a plot will be generated
legend	if TRUE a legend will be added to the plot
legpos	positioning of the legend in the plot
...	additional plot arguments

**Details**

The data are split into a calibration and a test data set (provided by "train"). Within the calibration set "kfold"-fold CV is performed by applying the classification method to "kfold"-1 parts and evaluation for the last part. The misclassification error is then computed for the training data, for the CV test data (CV error) and for the test data.

**Value**

trainerr	training error rate
testerr	test error rate
cvMean	mean of CV errors
cvSe	standard error of CV errors
cverr	all errors from CV
gamvec	range for gamma-values, taken from input

**Author(s)**

Peter Filzmoser <P.Filzmoser@tuwien.ac.at>

**References**

K. Varmuza and P. Filzmoser: Introduction to Multivariate Statistical Analysis in Chemometrics. CRC Press, Boca Raton, FL, 2009.

**See Also**

[svm](#)

**Examples**

```
data(fgl,package="MASS")
grp=fgl$type
X=scale(fgl[,1:9])
k=length(unique(grp))
dat=data.frame(grp,X)
n=nrow(X)
ntrain=round(n*2/3)
require(e1071)
set.seed(143)
train=sample(1:n,ntrain)
ressvm=svmEval(X,grp,train,gamvec=c(0,0.05,0.1,0.2,0.3,0.5,1,2,5),
  legpos="topright")
title("Support vector machines")
```

---

treeEval

*Classification tree evaluation by CV*

---

**Description**

Evaluation for classification trees by cross-validation

**Usage**

```
treeEval(X, grp, train, kfold = 10, cp = seq(0.01, 0.1, by = 0.01), plotit = TRUE,
  legend = TRUE, legpos = "bottomright", ...)
```

**Arguments**

X	standardized complete X data matrix (training and test data)
grp	factor with groups for complete data (training and test data)
train	row indices of X indicating training data objects
kfold	number of folds for cross-validation
cp	range for tree complexity parameter, see <a href="#">rpart</a>

plotit	if TRUE a plot will be generated
legend	if TRUE a legend will be added to the plot
legpos	positioning of the legend in the plot
...	additional plot arguments

### Details

The data are split into a calibration and a test data set (provided by "train"). Within the calibration set "kfold"-fold CV is performed by applying the classification method to "kfold"-1 parts and evaluation for the last part. The misclassification error is then computed for the training data, for the CV test data (CV error) and for the test data.

### Value

trainerr	training error rate
testerr	test error rate
cvMean	mean of CV errors
cvSe	standard error of CV errors
cverr	all errors from CV
cp	range for tree complexity parameter, taken from input

### Author(s)

Peter Filzmoser <P.Filzmoser@tuwien.ac.at>

### References

K. Varmuza and P. Filzmoser: Introduction to Multivariate Statistical Analysis in Chemometrics. CRC Press, Boca Raton, FL, 2009.

### See Also

[rpart](#)

### Examples

```
data(fgl, package="MASS")
grp=fgl$type
X=scale(fgl[,1:9])
k=length(unique(grp))
dat=data.frame(grp,X)
n=nrow(X)
ntrain=round(n*2/3)
require(rpart)
set.seed(123)
train=sample(1:n,ntrain)
par(mar=c(4,4,3,1))
restree=treeEval(X,grp,train,cp=c(0.01,0.02:0.05,0.1,0.15,0.2:0.5,1))
title("Classification trees")
```

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