# Package 'pcv' 

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corn Corn data

## Description

NIR spectra and moisture of 80 Corn samples.

## Usage

data(corn)

## Format

A list with two matrices, spectra and moisture.

## Details

This is a part of Corn dataset, which was downloaded from Eigenvector Research, Inc. website (https://eigenvector.com/resources/data-sets/), where it is availble publicly. This dataset contains several NIR spectra of corn samples recorded using different instruments. For our examples we took "mp5" spectra and corrected them using Standard Normal Variate transformation.

## Source

1. Eigenvector Research, Inc. (https://eigenvector.com/resources/data-sets/)

## Description

Returns parameters for cross-validation based on 'cv' value

## Usage

getCrossvalParams(cv, nobj)

Arguments

| cv <br> nobj | settings for cross-validation provided by user <br> number of objects in calibration set |
| :--- | :--- |
| getR | Creates rotation matrix to map a set vectors base 1 to a set of vectors <br> base 2. |

## Description

In both sets vectors should be orthonormal.

## Usage

getR(base1, base2)

## Arguments

$$
\begin{array}{ll}
\text { base1 } & \text { Matrix }(\mathrm{JXA}) \text { with A orthonormal vectors as columns to be rotated }(\mathrm{A}<=\mathrm{J}) \\
\text { base2 } & \text { Matrix }(\mathrm{JXA}) \text { with A orthonormal vectors as columns, base1 should be aligned } \\
\text { with }
\end{array}
$$

## Value

Rotation matrix (JXJ)

```
getxpvorth Generates the orthogonal part for Xpv
```


## Description

Generates the orthogonal part for Xpv

## Usage

getxpvorth(q.k, X.k, PRM)

## Arguments

q.k vector with orthogonal distances for cross-validation set for segment $k$
X.k matrix with local validation set for segment k

PRM projecton matrix for orthogonalization of residuals

## Value

A matrix with orthogonal part for Xpv

```
normalize
```

Normalization rows or columns of a matrix

## Description

Normalization rows or columns of a matrix

## Usage

```
normalize(
    X,
    dim = 1,
    weights = if (dim == 1) 1/sqrt(rowSums(X^2)) else 1/sqrt(colSums(X^2))
)
```


## Arguments

| $X$ | matrix with numeric values |
| :--- | :--- |
| $\operatorname{dim}$ | which dimension to normalize (1 for rows, 2 for columns) |
| weights | vector with normalization weights, by default 2-norm is use |

pcv Compute matrix with pseudo-validation set

## Description

Compute matrix with pseudo-validation set

## Usage

```
    pcv(
        X,
        ncomp = min(round(nrow(X)/nseg) - 1, col(X), 20),
        nseg = 4,
        scale = FALSE
    )
```


## Arguments

X
ncomp number of components for PCA decomposition
nseg number of segments in cross-validation
scale logical, standardize columns of X prior to decompositon or not

## Details

This is the old (original) version of PCV algorithm for PCA models. Use pcvpca instead. Ane check project web-site for details: https://github.com/svkucheryavski/pcv
The method computes pseudo-validation matrix Xpv , based on PCA decomposition of calibration set X and systematic (venetian blinds) cross-validation. It is assumed that data rows are ordered correctly, so systematic cross-validation can be applied

## Value

Pseudo-validation matrix (IxJ)
pcvcrossval Generate sequence of indices for cross-validation

## Description

Generates and returns sequence of object indices for each segment in random segmented crossvalidation

## Usage

pcvcrossval(cv = 1, nobj $=$ NULL, resp $=$ NULL)

## Arguments

cv cross-validation settings, can be a number, a list or a vector with integers.
nobj number of objects in a dataset
resp vector or matrix with response values to use in case of venetian blinds

## Details

Parameter 'cv' defines how to split the rows of the training set. The split is similar to crossvalidation splits, as PCV is based on cross-validation. This parameter can have the following values:

1. A number (e.g. 'cv=4'). In this case this number specifies number of segments for random splits, except 'cv=1' which is a special case for leave-one-out (full cross-validation).
2. A list with 2 values: ‘list("name", nseg)'. In this case '"name"‘ defines the way to make the split, you can select one of the following: ‘"loo"‘ for leave-one-out, ‘"rand"‘ for random splits or ‘"ven"‘ for Venetian blinds (systematic) splits. The second parameter, 'nseg', is a number of segments for splitting the rows into. For example, 'cv = list("ven", 4)', shown in the code examples above, tells PCV to use Venetian blinds splits with 4 segments.
3. A vector with integer numbers, e.g. ' $\mathrm{cv}=\mathrm{c}(1,2,3,1,2,3,1,2,3$ )'. In this case number of values in this vector must be the same as number of rows in the training set. The values specify which segment a particular row will belong to. In case of the example shown here, it is assumed that you have 9 rows in the calibration set, which will be split into 3 segments. The first segment will consist of measurements from rows 1,4 and 7 .

## Value

vector with object indices for each segment
pcvpca Procrustes cross-validation for PCA models

## Description

Procrustes cross-validation for PCA models

## Usage

```
    pcvpca(
        X,
        ncomp = min(nrow(X) - 1, ncol(X), 30),
        cv = list("ven", 4),
        center = TRUE,
        scale = FALSE,
        cv.scope = "global"
    )
```


## Arguments

X
ncomp number of components to use (more than the expected optimal number).
cv which split method to use for cross-validation (see description for details).
center logical, to center or not the data sets
scale logical, to scale or not the data sets
cv.scope scope for center/scale operations inside CV loop: 'global' - using globally computed mean and std or 'local' - recompute new for each local calibration set.

## Details

The method computes Procrustes validation set (PV-set), matrix Xpv, based on PCA decomposition of calibration set ' $X$ ' and cross-validation. See description of the method in [1].
Parameter 'cv' defines how to split the rows of the training set. The split is similar to crossvalidation splits, as PCV is based on cross-validation. This parameter can have the following values:

1. A number (e.g. 'cv $=4$ '). In this case this number specifies number of segments for random splits, except 'cv =1' which is a special case for leave-one-out (full cross-validation).
2. A list with 2 values: 'list("name", nseg)'. In this case '"name"‘ defines the way to make the split, you can select one of the following: ‘"loo"‘ for leave-one-out, ‘"rand"‘ for random splits or ‘"ven"‘ for Venetian blinds (systematic) splits. The second parameter, 'nseg', is a number of segments for
splitting the rows into. For example, 'cv = list("ven", 4)', shown in the code examples above, tells PCV to use Venetian blinds splits with 4 segments.
3. A vector with integer numbers, e.g. ' $\mathrm{cv}=\mathrm{c}(1,2,3,1,2,3,1,2,3$ )'. In this case number of values in this vector must be the same as number of rows in the training set. The values specify which segment a particular row will belong to. In case of the example shown here, it is assumed that you have 9 rows in the calibration set, which will be split into 3 segments. The first segment will consist of measurements from rows 1,4 and 7 .

Parameter 'cv.scope' influences how the Procrustean rule is met. In case of "global" scope, the rule will be met strictly - distances for PV-set and the global model will be identical to the distances from conventional cross-validation. In case of "local" scope, every local model will have its own center and scaling factor and hence the rule will be almost met (the distances will be close but not identical).

## Value

Matrix with PV-set (same size as X)

## References

1. S. Kucheryavskiy, O. Rodionova, A. Pomerantsev. Procrustes cross-validation of multivariate regression models. Analytica Chimica Acta, 1255 (2022) [https://doi.org/10.1016/j.aca.2023.341096]

## Examples

```
# load NIR spectra of Corn samples
data(corn)
X <- corn$spectra
# generate Xpv set based on PCA decomposition with A = 20 and venetian blinds split with 4 segments
Xpv <- pcvpca(X, ncomp = 20, center = TRUE, scale = FALSE, cv = list("ven", 4))
# show the original spectra and the PV-set (as is and mean centered)
oldpar <- par(mfrow = c(2, 2))
matplot(t(X), type = "l", lty = 1, main = "Original data")
matplot(t(Xpv), type = "l", lty = 1, main = "PV-set")
matplot(t(scale(X, scale = FALSE)), type = "l", lty = 1, main = "Original data (mean centered)")
matplot(t(scale(Xpv, scale = FALSE)), type = "l", lty = 1, main = "PV-set (mean centered)")
par(oldpar)
```

pcvper Procrustes cross-validation for PCR models

## Description

Procrustes cross-validation for PCR models

```
Usage
    pcvpcr(
    X,
    Y,
    ncomp = min(nrow(X) - 1, ncol(X), 30),
    cv = list("ven", 4),
    center = TRUE,
    scale = FALSE,
    cv.scope = "global"
    )
```


## Arguments

$X \quad$ matrix with predictors from the training set.
Y vector with response values from the training set.
ncomp number of components to use (more than the expected optimal number).
cv which split method to use for cross-validation (see description of method 'pcvpls()$^{\text {' }}$ for details).
center logical, to center or not the data sets
scale logical, to scale or not the data sets
cv.scope scope for center/scale operations inside CV loop: 'global' — using globally computed mean and std or 'local' - recompute new for each local calibration set.

## Details

The method computes pseudo-validation matrix Xpv , based on PCR decomposition of calibration set $\mathrm{X}, \mathrm{y}$ and cross-validation.
Parameter 'cv' defines how to split the rows of the training set. The split is similar to crossvalidation splits, as PCV is based on cross-validation. This parameter can have the following values:

1. A number (e.g. 'cv $=4$ '). In this case this number specifies number of segments for random splits, except 'cv $=1$ ' which is a special case for leave-one-out (full cross-validation).
2. A list with 2 values: 'list("name", nseg)'. In this case '"name"' defines the way to make the split, you can select one of the following: ‘"loo"‘ for leave-one-out, ‘"rand"‘ for random splits or ‘"ven"‘ for Venetian blinds (systematic) splits. The second parameter, 'nseg', is a number of segments for splitting the rows into. For example, 'cv = list("ven", 4)', shown in the code examples above, tells PCV to use Venetian blinds splits with 4 segments.
3. A vector with integer numbers, e.g. ' $\mathrm{cv}=\mathrm{c}(1,2,3,1,2,3,1,2,3)^{\prime}$ '. In this case number of values in this vector must be the same as number of rows in the training set. The values specify which segment a particular row will belong to. In case of the example shown here, it is assumed that you have 9 rows in the calibration set, which will be split into 3 segments. The first segment will consist of measurements from rows 1,4 and 7 .
Parameter 'cv.scope" influences how the Procrustean rule is met. In case of "global" scope, the rule will be met strictly - error of predictions for PV-set and the global model will be identical to the error from conventional cross-validation. In case of "local" scope, every local model will have its own center and hence the rule will be almost met (the errors will be close but not identical).

## Value

Pseudo-validation matrix (same size as X ) with an additional attribute, 'D' which contains the scaling coefficients (ck/c)

## References

1. S. Kucheryavskiy, O. Rodionova, A. Pomerantsev. Procrustes cross-validation of multivariate regression models. Analytica Chimica Acta, 1255 (2022) [https://doi.org/10.1016/j.aca.2023.341096]

## Examples

```
    # load NIR spectra of Corn samples
    data(corn)
    X <- corn$spectra
    Y <- corn$moisture
    # generate Xpv set based on PCA decomposition with A = 20 and venetian blinds split with 4 segments
    Xpv <- pcvpcr(X, Y, ncomp = 20, center = TRUE, scale = FALSE, cv = list("ven", 4))
    # show the original spectra and the PV-set (as is and mean centered)
    oldpar <- par(mfrow = c(2, 2))
    matplot(t(X), type = "l", lty = 1, main = "Original data")
    matplot(t(Xpv), type = "l", lty = 1, main = "PV-set")
    matplot(t(scale(X, scale = FALSE)), type = "l", lty = 1, main = "Original data (mean centered)")
    matplot(t(scale(Xpv, scale = FALSE)), type = "l", lty = 1, main = "PV-set (mean centered)")
    par(oldpar)
```

    pcvpls Procrustes cross-validation for PLS models
    
## Description

Procrustes cross-validation for PLS models

## Usage

pcupls(
X ,
Y,
ncomp $=\min (\operatorname{nrow}(X)-1, n \operatorname{col}(X), 30)$,
center = TRUE,
scale = FALSE,
cv = list("ven", 4),
cv.scope = "global"
)

## Arguments

| $X$ | matrix with predictors from the training set. |
| :--- | :--- |
| Y | vector or matrix with response values from the training set. |
| ncomp | number of components to use (more than the expected optimal number). |
| center | logical, to center or not the data sets |
| scale | logical, to scale or not the data sets |
| cv | which split method to use for cross-validation (see description for details). <br> cv.scope |
| scope for center/scale operations inside CV loop: 'global' - using globally <br> computed mean and std or 'local' - recompute new for each local calibration <br> set. |  |

## Details

The method computes pseudo-validation matrix Xpv, based on PLS decomposition of calibration set X , y and cross-validation.

Parameter 'cv' defines how to split the rows of the training set. The split is similar to crossvalidation splits, as PCV is based on cross-validation. This parameter can have the following values:

1. A number (e.g. ' $\mathrm{cv}=4$ '). In this case this number specifies number of segments for random splits, except ' $\mathrm{cv}=1$ ' which is a special case for leave-one-out (full cross-validation).
2. A list with 2 values: 'list("name", nseg)'. In this case ‘"name"' defines the way to make the split, you can select one of the following: ‘"loo"‘ for leave-one-out, ‘"rand"‘ for random splits or ‘"ven"' for Venetian blinds (systematic) splits. The second parameter, 'nseg', is a number of segments for splitting the rows into. For example, 'cv = list("ven", 4)', shown in the code examples above, tells PCV to use Venetian blinds splits with 4 segments.
3. A vector with integer numbers, e.g. ' $\mathrm{cv}=\mathrm{c}(1,2,3,1,2,3,1,2,3$ )'. In this case number of values in this vector must be the same as number of rows in the training set. The values specify which segment a particular row will belong to. In case of the example shown here, it is assumed that you have 9 rows in the calibration set, which will be split into 3 segments. The first segment will consist of measurements from rows 1,4 and 7 .

Parameter 'cv.scope' influences how the Procrustean rule is met. In case of "global" scope, the rule will be met strictly - error of predictions for PV-set and the global model will be identical to the error from conventional cross-validation. In case of "local" scope, every local model will have its own center and hence the rule will be almost met (the errors will be close but not identical).

## Value

Pseudo-validation matrix (same size as X ) with an additional attribute, 'D' which contains the scaling coefficients (ck/c)

## References

1. S. Kucheryavskiy, O. Rodionova, A. Pomerantsev. Procrustes cross-validation of multivariate regression models. Analytica Chimica Acta, 1255 (2022) [https://doi.org/10.1016/j.aca.2023.341096]

## Examples

```
# load NIR spectra of Corn samples
data(corn)
X <- corn$spectra
Y <- corn$moisture
# generate Xpv set based on PCA decomposition with A = 20 and venetian blinds split with 4 segments
Xpv <- pcvpls(X, Y, ncomp = 20, center = TRUE, scale = FALSE, cv = list("ven", 4))
# show the original spectra and the PV-set (as is and mean centered)
oldpar <- par(mfrow = c(2, 2))
matplot(t(X), type = "l", lty = 1, main = "Original data")
matplot(t(Xpv), type = "l", lty = 1, main = "PV-set")
matplot(t(scale(X, scale = FALSE)), type = "l", lty = 1, main = "Original data (mean centered)")
matplot(t(scale(Xpv, scale = FALSE)), type = "l", lty = 1, main = "PV-set (mean centered)")
par(oldpar)
# show the heatmap with the scaling coefficients
plotD(Xpv)
```

pcvreg Procrustes cross-validation for multivariate regression models

## Description

This is a generic method, use 'pcvpls()' or ' $\operatorname{pcvpcr}()^{\prime}$ instead.

## Usage

```
pcvreg(
    X,
    Y,
    ncomp = min(nrow(X) - 1, ncol(X), 30),
    cv = list("ven", 4),
    center = TRUE,
    scale = FALSE,
    funlist = list(),
    cv.scope = "global"
)
```


## Arguments

$X \quad$ matrix with predictors from the training set.
Y vector with response values from the training set.
ncomp number of components to use (more than the expected optimal number).

| cv | which split method to use for cross-validation (see description of method 'pcv- <br> pls()' for details). |
| :--- | :--- |
| center | logical, to center or not the data sets |
| scale | logical, to scale or not the data sets |
| funlist | list with functions for particular implementation |
| cv.scope | scope for center/scale operations inside CV loop: 'global' - using globally <br> computed mean and std or 'local' - recompute new for each local calibration <br> set. |

plotD | Plots heatmap for scaling coefficients obtained when generating PV- |
| :--- |
| set for PCR or PLS |

## Description

Plots heatmap for scaling coefficients obtained when generating PV-set for PCR or PLS

## Usage

```
plotD(
    Xpv,
    colmap = colorRampPalette(c("blue", "white", "red"))(256),
    lim = c(-2, 4),
    xlab = "Components",
    ylab = "Segments",
)
```


## Arguments

| Xpv | PV-set generated by 'pcvpcr()' or 'pcvpls()'. |
| :--- | :--- |
| colmap | colormap - any with 256 colors. |
| lim | limits for color map (smallest/largest expected value), centered around 1. |
| xlab | label for x-axis |
| ylab | label for y-axis |
| $\ldots$ | any other parameters for method 'image' |

## Value

No return value, just creates a plot.

```
rotationMatrixToX1 Creates a rotation matrix to map a vectorx to [1 0 0 ... 0]
```


## Description

Creates a rotation matrix to map a vector x to $\left[\begin{array}{llll}1 & 0 & 0 & \ldots\end{array}\right]$

## Usage

rotationMatrixToX1(x)

## Arguments

$x \quad$ Vector (sequence with J coordinates)

## Value

Rotation matrix (JxJ)
simpls SIMPLS algorithm

## Description

SIMPLS algorithm for calibration of PLS model

## Usage

simpls(X, Y, ncomp)

## Arguments

$\mathrm{X} \quad$ a matrix with x values (predictors)
$Y \quad$ a matrix with y values (responses)
ncomp number of components to calculate

## Value

a list with computed weights, x - and y -loadings for PLS regression model.

## References

[1]. S. de Jong. SIMPLS: An Alternative approach to partial least squares regression. Chemometrics and Intelligent Laboratory Systems, 18, 1993 (251-263).

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