Package ‘OmicsPLS’

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Title Data Integration with Two-Way Orthogonal Partial Least Squares
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Description Performs the O2PLS data integration method for two datasets yielding joint and data-specific parts for each dataset.
The algorithm automatically switches to a memory-efficient approach to fit O2PLS to high dimensional data.
It provides a rigorous and a faster alternative cross-validation method to select the number of components,
as well as functions to report proportions of explained variation and to construct plots of the results.
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R topics documented:

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adjR2

Gridwise adjusted R2 for O2PLS

Description

For (a grid of) values for \(a, nx\) and \(ny\), loocv calculates the R2 of the joint part. Parallel computing is supported on Windows with package parallel.

Usage

\[
\text{adjR2}(X, Y, a = 1:2, a2 = 1, b2 = 1, \text{func} = \text{o2m}, \text{parall} = \text{F}, \\
\text{cl} = \text{NULL}, \text{stripped} = \text{TRUE}, p_{\text{thresh}} = 3000, q_{\text{thresh}} = p_{\text{thresh}}, \\
tol = 1e-10, \text{max}_\text{iterations} = 100)
\]

Arguments

- **X**: Numeric matrix. Vectors will be coerced to matrix with as.matrix (if this is possible)
- **Y**: Numeric matrix. Vectors will be coerced to matrix with as.matrix (if this is possible)
- **a**: Vector of integers. Contains the numbers of joint components.
- **a2**: Vector of integers. Contains the numbers of orthogonal components in \(X\).
- **b2**: Vector of integers. Contains the numbers of orthogonal components in \(Y\).
- **func**: Function to fit the O2PLS model with. Only o2m and o2m_stripped are supported.
- **parall**: Integer. Should a parallel cluster be set up using package parallel (Windows)? Best is to leave it to FALSE.
**crossval_o2m**

Object of class 'cluster'. If parall is TRUE and cl is not NULL, calculations are parallelized over workers in cl.

stripped Logical. Use the stripped version of o2m (usually when cross-validating)?

p_thresh Integer. If X has more than p_thresh columns, a power method optimization is used, see o2m2

q_thresh Integer. If Y has more than q_thresh columns, a power method optimization is used, see o2m2

tol double. Threshold for power method iteration

max_iterations Integer, Maximum number of iterations for power method

**Details**

The use of this function is to calculate the R2 of the joint part, while varying the number of orthogonal components. Adding more joint components will increase the R2!

A parallelized version is built in -tested on windows-, use package parallel and set parall=TRUE to activate this. There should not be already a cluster object with the name cl. In case of some error, don’t forget to invoke stopCluster(cl) to end the cluster. See Task Manager (Windows) to verify that the workers are spanned/ended.

See loocv for more intuition.

**Value**

Matrix with two rows:

adjR2X Contains the joint R2 in X

adjR2Y Contains the joint R2 in Y

---

crossval_o2m Cross-validate procedure for O2PLS

**Description**

Cross-validate procedure for O2PLS

**Usage**

crossval_o2m(X, Y, a, ax, ay, nr_folds, nr_cores = 1, stripped = TRUE, p_thresh = 3000, q_thresh = p_thresh, tol = 1e-10, max_iterations = 100)
Arguments

- **X**
  Numeric matrix. Vectors will be coerced to matrix with `as.matrix` (if this is possible)

- **Y**
  Numeric matrix. Vectors will be coerced to matrix with `as.matrix` (if this is possible)

- **a**
  Vector of positive integers. Denotes the numbers of joint components to consider.

- **ax**
  Vector of non-negative integers. Denotes the numbers of X-specific components to consider.

- **ay**
  Vector of non-negative integers. Denotes the numbers of Y-specific components to consider.

- **nr_folds**
  Positive integer. Number of folds to consider. Note: `kcv=N` gives leave-one-out CV. Note that CV with less than two folds does not make sense.

- **nr_cores**
  Positive integer. Number of cores to use for CV. You might want to use `detectCores()`. Defaults to 1.

- **stripped**
  Logical. Use the stripped version of o2m (usually when cross-validating)?

- **p_thresh**
  Integer. If X has more than `p_thresh` columns, a power method optimization is used, see `o2m2`

- **q_thresh**
  Integer. If Y has more than `q_thresh` columns, a power method optimization is used, see `o2m2`

- **tol**
  Double. Threshold for power method iteration

- **max_iterations**
  Integer, Maximum number of iterations for power method

Details

This is the standard CV approach. It minimizes the sum of the prediction errors of X and Y over a three-dimensional grid of integers. Parallelization is possible on all platforms. On Windows it uses `makePSOCKcluster`, then exports all necessary objects to the workers, and then calls `parLapply`. On OSX and Linux the more friendly `mclapply` is used, which uses forking. A print method is defined, printing the minimizers and minimum in a readable way. Also the elapsed time is tracked and reported.

Value

List of class "cvo2m" with the original and sorted Prediction errors and the number of folds used.

Examples

```r
local({
  X = scale(jitter(tcrossprod(rnorm(100),runif(10))))
  Y = scale(jitter(tcrossprod(rnorm(100),runif(10))))
  crossval_o2m(X, Y, a = 1:4, ax = 1:2, ay = 1:2,
               nr_folds = 5, nr_cores = 1)
})
```
crossval_o2m_adjR2  

*Adjusted Cross-validate procedure for O2PLS*

**Description**

Combines CV with R2 optimization

**Usage**

```r
crossval_o2m_adjR2(X, Y, a, ax, ay, nr_folds, nr_cores = 1, 
stripped = TRUE, p_thresh = 3000, q_thresh = p_thresh, 
tol = 1e-10, max_iterations = 100)
```

**Arguments**

<table>
<thead>
<tr>
<th>Argument</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>X</td>
<td>Numeric matrix. Vectors will be coerced to matrix with <code>as.matrix</code> (if this is possible)</td>
</tr>
<tr>
<td>Y</td>
<td>Numeric matrix. Vectors will be coerced to matrix with <code>as.matrix</code> (if this is possible)</td>
</tr>
<tr>
<td>a</td>
<td>Vector of positive integers. Denotes the numbers of joint components to consider.</td>
</tr>
<tr>
<td>ax</td>
<td>Vector of non-negative integers. Denotes the numbers of X-specific components to consider.</td>
</tr>
<tr>
<td>ay</td>
<td>Vector of non-negative integers. Denotes the numbers of Y-specific components to consider.</td>
</tr>
<tr>
<td>nr_folds</td>
<td>Positive integer. Number of folds to consider. Note: <code>kcv=N</code> gives leave-one-out CV. Note that CV with less than two folds does not make sense.</td>
</tr>
<tr>
<td>nr_cores</td>
<td>Positive integer. Number of cores to use for CV. You might want to use <code>detectCores()</code>. Defaults to 1.</td>
</tr>
<tr>
<td>stripped</td>
<td>Logical. Use the stripped version of <code>o2m</code> (usually when cross-validating)?</td>
</tr>
<tr>
<td>p_thresh</td>
<td>Integer. If X has more than <code>p_thresh</code> columns, a power method optimization is used, see <code>o2m2</code></td>
</tr>
<tr>
<td>q_thresh</td>
<td>Integer. If Y has more than <code>q_thresh</code> columns, a power method optimization is used, see <code>o2m2</code></td>
</tr>
<tr>
<td>tol</td>
<td>double. Threshold for power method iteration</td>
</tr>
<tr>
<td>max_iterations</td>
<td>Integer, Maximum number of iterations for power method</td>
</tr>
</tbody>
</table>

**Details**

This is an alternative way of cross-validating. It is proposed in `citation(OmicsPLS)`. This approach is (much) faster than the standard `crossval_o2m` approach and works fine even with two folds. For each element in `n` it looks for nx and ny that maximize the $R^2$ between T and U in the O2PLS model. This approach often yields similar integer as the standard approach. We however suggest to use the standard approach to minimize the prediction error around the found integers.
loadings

Extract the loadings from an O2PLS fit

Value
data.frame with four columns: MSE, n, nx and ny. Each row corresponds to an element in a.

Examples

```r
local({
  X = scale(jitter(tcrossprod(rnorm(100), runif(10))))
  Y = scale(jitter(tcrossprod(rnorm(100), runif(10))))
  crossval_o2m_adjR2(X, Y, a = 1:4, ax = 1:2, ay = 1:2,
    nr_folds = 5, nr_cores = 1)
})
```

loadings

Description
This function extracts loading parameters from an O2PLS fit

Usage

```r
loadings(x, ...)
```

## S3 method for class 'o2m'

```r
loadings(x, loading_name = c("Xjoint", "Yjoint", "Xorth", "Yorth"),
  subset = 0, sorted = FALSE, ...)
```

Arguments

- **x**: Object of class o2m
- **...**: For consistency
- **loading_name**: character string. One of the following: 'Xjoint', 'Yjoint', 'Xorth' or 'Yorth'.
- **subset**: subset of loading vectors to be extracted.
- **sorted**: Logical. Should the rows of the loadings be sorted according to the absolute magnitude of the first column?

Value
Loading matrix

See Also

scores.o2m

Examples

```r
loadings(o2m(scale(-2:2), scale(-2:2*4), 1, 0, 0))
```
**Description**

For (a grid of) values for \(a, nx\) and \(ny\), \texttt{loocv} estimates the prediction error using k-fold CV.

**Usage**

\[
\text{loocv}(X, Y, a = 1:2, a2 = 1, b2 = 1, \text{fitted\_model} = \text{NULL}, \\
\text{func} = \text{o2m}, \text{app\_err} = F, \text{kcv}, \text{stripped} = \text{TRUE}, \text{p\_thresh} = 3000, \\
\text{q\_thresh} = \text{p\_thresh}, \text{tol} = 1e-10, \text{max\_iterations} = 100)
\]

**Arguments**

- **X**: Numeric matrix. Vectors will be coerced to matrix with \texttt{as.matrix} (if this is possible)
- **Y**: Numeric matrix. Vectors will be coerced to matrix with \texttt{as.matrix} (if this is possible)
- **a**: Vector of integers. Contains the numbers of joint components.
- **a2**: Vector of integers. Contains the numbers of orthogonal components in \(X\).
- **b2**: Vector of integers. Contains the numbers of orthogonal components in \(Y\).
- **fitted\_model**: List. \texttt{O2PLS} model fit with \texttt{o2m}. Is used to calculate the apparent error without recalculating this fit.
- **func**: Function to fit the \texttt{O2PLS} model with. Only \texttt{o2m} and \texttt{o2m\_stripped} are supported.
- **app\_err**: Logical. Should the apparent error also be computed? Not used anymore.
- **kcv**: Integer. The value of \(k\), i.e. the number of folds. Choose \(N\) for LOO-CV.
- **stripped**: Logical. Use the stripped version of \texttt{o2m} (usually when cross-validating)?
- **p\_thresh**: Integer. If \(X\) has more than \(p\_thresh\) columns, a power method optimization is used, see \texttt{o2m2}
- **q\_thresh**: Integer. If \(Y\) has more than \(q\_thresh\) columns, a power method optimization is used, see \texttt{o2m2}
- **tol**: double. Threshold for power method iteration
- **max\_iterations**: Integer, Maximum number of iterations for power method

**Details**

Note that this function can be easily parallelized (on Windows e.g. with the \texttt{parallel} package.). The parameters \(a, a2\) and \(b2\) can be integers or vectors of integers. A for loop is used to loop over all combinations. The resulting output is a list, which is more easy to interpret if you use \texttt{array(unlist(output\_of\_loocv\$CVerr))} as in the example below. The array will have varying \(a\) along the first dimension and \(a2\) and \(b2\) along the second and third respectively. Typing \texttt{example(loocv)} (hopefully) clarifies this function.
loocv_combi

### Value

List with two numeric vectors:

- CVerr: Contains the k-fold CV estimated RMSEP
- Fiterr: Contains the apparent error

---

**loocv_combi**  
*K-fold CV based on symmetrized prediction error*

---

### Description

The prediction error of both $X\sim X\text{hat}$ and $Y\sim Y\text{hat}$ are summed. This provides a symmetrized version of loocv.

### Usage

```r
loocv_combi(X, Y, a = 1:2, a2 = 1, b2 = 1, fitted_model = NULL, 
            func = o2m, app_err = F, kcv, stripped = TRUE, p_thresh = 3000, 
            q_thresh = p_thresh, tol = 1e-10, max_iterations = 100)
```

### Arguments

- **X**: Numeric matrix. Vectors will be coerced to matrix with `as.matrix` (if this is possible)
- **Y**: Numeric matrix. Vectors will be coerced to matrix with `as.matrix` (if this is possible)
- **a**: Vector of integers. Contains the numbers of joint components.
- **a2**: Vector of integers. Contains the numbers of orthogonal components in $X$.
- **b2**: Vector of integers. Contains the numbers of orthogonal components in $Y$.
- **fitted_model**: List. O2PLS model fit with `o2m`. Is used to calculate the apparent error without recalculating this fit.
- **func**: Function to fit the O2PLS model with. Only `o2m` and `o2m_stripped` are supported.
- **app_err**: Logical. Should the apparent error also be computed? Not used anymore.
- **kcv**: Integer. The value of $k$, i.e. the number of folds. Choose $N$ for LOO-CV.
- **stripped**: Logical. Use the stripped version of `o2m` (usually when cross-validating)?
- **p_thresh**: Integer. If $X$ has more than $p\text{.thresh}$ columns, a power method optimization is used, see `o2m2`
- **q_thresh**: Integer. If $Y$ has more than $q\text{.thresh}$ columns, a power method optimization is used, see `o2m2`
- **tol**: Double. Threshold for power method iteration
- **max_iterations**: Integer. Maximum number of iterations for power method
**Details**

Note that this function can be easily parallelized (on Windows e.g. with the `parallel` package.). If there are NAs in the CVerr component, this is due to an error in the fitting.

**Value**

List with two numeric vectors:

- **CVerr**: Contains the k-fold CV estimated RMSEP
- **Fiterr**: Contains the apparent error

---

**Description**

Calculate mean squared difference

**Usage**

```r
mse(x, y = 0, na.rm = FALSE)
```

**Arguments**

- **x**: Numeric vector or matrix.
- **y**: Numeric vector or matrix. Defaults to 0.
- **na.rm**: Remove NA’s?

**Details**

Is equal to `ssq(x - y)/length(x)`. If `x` and `y` are of unequal length, the invoked minus-operator will try to make the best out of it by recycling elements of the shorter object (usually you don’t want that). In particular if `x` is an N x p matrix and `y` an N x 1 vector, `y` is subtracted from each column of `x`, and if `y=0` (default) you get the mean of `vec(x^2)`

**Value**

The mean of the squared differences elementwise.

**Examples**

```r
mse(2)
mse(1:10, 2:11) == 1
mse(matrix(rnorm(500), 100, 5), matrix(rnorm(500), 100, 5))
```
Perform O2-PLS data integration with two-way orthogonal corrections

Description

NOTE THAT THIS FUNCTION DOES NOT CENTER NOR SCALES THE MATRICES! Any normalization you will have to do yourself. It is best practice to at least center the variables though.

Usage

\[
o2m(X, Y, n, nx, ny, stripped = FALSE, p_thresh = 3000, q_thresh = p_thresh, tol = 1e-10, max_iterations = 100)
\]

Arguments

- **X**: Numeric matrix. Vectors will be coerced to matrix with `as.matrix` (if this is possible)
- **Y**: Numeric matrix. Vectors will be coerced to matrix with `as.matrix` (if this is possible)
- **n**: Integer. Number of joint PLS components. Must be positive!
- **nx**: Integer. Number of orthogonal components in X. Negative values are interpreted as 0
- **ny**: Integer. Number of orthogonal components in Y. Negative values are interpreted as 0
- **stripped**: Logical. Use the stripped version of o2m (usually when cross-validating)?
- **p_thresh**: Integer. If \(X\) has more than \(p_{thresh}\) columns, a power method optimization is used, see o2m2
- **q_thresh**: Integer. If \(Y\) has more than \(q_{thresh}\) columns, a power method optimization is used, see o2m2
- **tol**: double. Threshold for power method iteration
- **max_iterations**: Integer. Maximum number of iterations for power method

Details

If both \(nx\) and \(ny\) are zero, \(o2m\) is equivalent to PLS2 with orthonormal loadings. This is a 'slower' (in terms of memory) implementation of O2PLS, and is using `svd`, use `stripped=T` for a stripped version with less output. If either \(\text{ncol}(X) > p_{thresh}\) or \(\text{ncol}(Y) > q_{thresh}\), an alternative method is used (NIPALS) which does not store the entire covariance matrix. The squared error between iterands in the NIPALS approach can be adjusted with `tol`. The maximum number of iterations in the NIPALS approach is tuned by `max_iterations`.
Value

A list containing

- **Tt**: Joint X scores
- **W**: Joint X loadings
- **U**: Joint Y scores
- **C**: Joint Y loadings
- **E**: Residuals in X
- **f**: Residuals in Y
- **T_Yosc**: Orthogonal X scores
- **P_Yosc**: Orthogonal X loadings
- **W_Yosc**: Orthogonal X weights
- **U_Xosc**: Orthogonal Y scores
- **P_Xosc**: Orthogonal Y loadings
- **C_Xosc**: Orthogonal Y weights
- **B_U**: Regression coefficient in Tt ~ U
- **B_T**: Regression coefficient in U ~ Tt
- **H_TU**: Residuals in Tt in Tt ~ U
- **H_UT**: Residuals in U in U ~ Tt
- **X_hat**: Prediction of X with Y
- **Y_hat**: Prediction of Y with X
- **R2X**: Variation (measured with ssq) of the modeled part in X (defined by joint + orthogonal variation) as proportion of variation in X
- **R2Y**: Variation (measured with ssq) of the modeled part in Y (defined by joint + orthogonal variation) as proportion of variation in Y
- **R2Xcorr**: Variation (measured with ssq) of the joint part in X as proportion of variation in X
- **R2Ycorr**: Variation (measured with ssq) of the joint part in Y as proportion of variation in Y
- **R2X_YO**: Variation (measured with ssq) of the orthogonal part in X as proportion of variation in X
- **R2Y_XO**: Variation (measured with ssq) of the orthogonal part in Y as proportion of variation in Y
- **R2Xhat**: Variation (measured with ssq) of the predicted X as proportion of variation in X
- **R2Yhat**: Variation (measured with ssq) of the predicted Y as proportion of variation in Y

See Also

summary.o2m, plot.o2m, crossval_o2m
Examples

test_X <- scale(matrix(rnorm(100*10),100,10))
test_Y <- scale(matrix(rnorm(100*11),100,11))
# -------- Default run --------
o2m(test_X, test_Y, 3, 2, 1)
# -------- Stripped version --------
o2m(test_X, test_Y, 3, 2, 1, stripped = TRUE)
# -------- High dimensional version --------
o2m(test_X, test_Y, 3, 2, 1, p_thresh = 1)
# -------- High D and stripped version --------
o2m(test_X, test_Y, 3, 2, 1, stripped = TRUE, p_thresh = 1)
# -------- Now with more iterations --------
o2m(test_X, test_Y, 3, 2, 1, stripped = TRUE, p_thresh = 1, max_iterations = 1e6)

---

OmicsPLS

Data integration with O2PLS: Two-Way Orthogonal Partial Least Squares

Description

OmicsPLS software is described in (el Bouhaddani et al, 2018, BMC Bioinformatics). This is based on work of (Trygg & Wold, 2003). Includes the O2PLS fit, some misc functions and some cross-validation tools.

Model and assumptions

Note that the rows of $X$ and $Y$ are the subjects and columns are variables. The number of columns may be different, but the subjects should be the same in both datasets.

The O2PLS model (Trygg & Wold, 2003) decomposes two datasets $X$ and $Y$ into three parts.

- 1. A joint part, representing the relationship between $X$ and $Y$
- 2. An orthogonal part, representing the unrelated latent variation in $X$ and $Y$ separately.
- 3. A noise part capturing all residual variation.

See also the corresponding paper (el Bouhaddani et al, 2018).

Fitting

The O2PLS fit is done with o2m. For data $X$ and $Y$ you can run o2m(X, Y, n, nx, ny) for an O2PLS fit with $n$ joint and $nx, ny$ orthogonal components. See the help page of o2m for more information on parameters. There are four ways to obtain an O2PLS fit, depending on the dimensionality.

- For the not-too-high dimensional case, you may use o2m with default parameters. E.g. o2m(X, Y, n, nx, ny).
• In case you don’t want the fancy output, but only the parameters, you may add `stripped = TRUE` to obtain a stripped version of `o2m` which avoids calculating and storing some matrices. E.g. `o2m(X, Y, n, nx, ny, stripped=TRUE).

• For high dimensional cases defined by `ncol(X) > p_thresh` and `ncol(Y) > q_thresh` a Power-Method approach is used which avoids storing large matrices. E.g. `o2m(X, Y, n, nx, ny, p_thresh=3000, q_thresh=3000)`. The thresholds are by default both at 3000 variables.

• If you want a stripped version in the high dimensional case, add `stripped=TRUE`. E.g. `o2m(X, Y, n, nx, ny, stripped=TRUE, p_thresh=3000, q_thresh=3000).

Obtaining results

After fitting an O2PLS model, by running e.g. `fit = o2m(X, Y, n, nx, ny)`, the results can be visualised. Use `plot(fit, ...)` to plot the desired loadings with/without ggplot2. Use `summary(fit, ...)` to see the relative explained variances in the joint/orthogonal parts. Also plotting the joint scores `fit$T_t, fit$U` and orthogonal scores `fit$T_Yosc, fit$U_Xosc` are of help.

Cross-validating

Determining the number of components `n, nx, ny` is an important task. For this we have two methods. See `citation("OmicsPLS")` for our proposed approach for determining the number of components, implemented in `crossval_o2m_adjR2`!

- Cross-validation (CV) is done with `crossval_o2m` and `crossval_o2m_adjR2`, both have built in parallelization which relies on the parallel package. Usage is something like `crossval_o2m(X, Y, a, ax, ay)` where `a, ax, ay` are vectors of integers. See the help pages. `kcv` is the number of folds, with `kcv = nrow(X)` for Leave-One-Out CV.

- For `crossval_o2m_adjR2` the same parameters are to be specified. This way of cross-validating is (potentially much) faster than the standard approach.

Misc

Also some handy tools are available

- `orth(X)` is a function to obtain an orthogonalized version of a matrix or vector `X`.
- `ssq(X)` is a function to calculate the sum of squares (or squared Frobenius norm) of `X`. See also `vnorm` for calculating the norm of each column in `X`.
- `mse(x, y)` returns the mean squared difference between two matrices/vectors. By default `y = 0`.

Citation

If you use the OmicsPLS R package in your research, please cite the corresponding software paper:


The bibtex entry can be obtained with command `citation("OmicsPLS")`. Thank You!

The original paper proposing O2PLS is

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Maintainer: Said el Bouhaddani (<s.elbouhaddani@umcutrecht.nl>).

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**orth**

**Orthogonalize a matrix**

**Description**

Orthogonalize a matrix

**Usage**

```r
orth(X, X_true = NULL, type = c("QR", "SVD"))
```

**Arguments**

- `X` Numeric vector or matrix.
- `X_true` (optional) A ‘true’ matrix/vector. Used to correct the sign of the orthonormalized `X` if QR is used. Only the first column is corrected.
- `type` A character or numeric. Should be one of "QR" or "SVD".

**Details**

Choosing type='QR' uses a QR decomposition of `X` to produce orthonormal columns. For type=='SVD' it uses an SVD decomposition. The columns are corrected for sign.

**Value**

An orthogonalized representation of `X`

**Examples**

```r
orth(c(3,4))
round(crossprod(orth(matrix(rnorm(500),100,5))),4)
orth(matrix(1:9,3,3),type='QR')[,1] - orth(1:3); orth(matrix(1:9,3,3),type='SVD')[,1] - orth
```
**plot.o2m**  
*Plot one or two loading vectors for class o2m*

**Description**  
This function plots one or two loading vectors, by default with ggplot2.

**Usage**  
```r  
## S3 method for class 'o2m'  
plot(x, loading_name = c("Xjoint", "Yjoint", "Xorth",  
"Yorth"), i = 1, j = NULL, use_ggplot2 = TRUE,  
label = c("number", "colnames"), ...)  
```

**Arguments**  
- `x`: An O2PLS fit, with class 'o2m'
- `loading_name`: character string. One of the following: 'Xjoint', 'Yjoint', 'Xorth' or 'Yorth'.
- `i`: Integer. First component to be plotted.
- `j`: NULL (default) or Integer. Second component to be plotted.
- `use_ggplot2`: Logical. Default is TRUE. If FALSE, the usual plot device will be used.
- `label`: Character, either 'number' or 'colnames'. The first option prints numbers, the second prints the colnames
- `...`: Further arguments to `geom_text`, such as size, col, alpha, etc.

**Value**  
If `use_ggplot2` is TRUE a ggplot2 object. Else NULL.

**See Also**  
- `summary.o2m`

**predict.o2m**  
*Predicts X or Y*

**Description**  
Predicts X or Y based on new data on Y or X

**Usage**  
```r  
## S3 method for class 'o2m'  
predict(object, newdata, XorY = c("X", "Y"), ...)  
```
Arguments

object List. Should be of class o2m.
newdata New data, which one of X or Y is specified in XorY.
XorY Character specifying whether newdata is X or Y.
... For compatibility

Details

Prediction is done after correcting for orthogonal parts.

Value

Predicted Data

Examples

predict(o2m(scale(1:10), scale(1:10), 1, 0, 0), newdata = scale(1:5), XorY = "X")

---

print.cvo2m Cross-validate procedure for O2PLS

Description

Cross-validate procedure for O2PLS

Usage

## S3 method for class 'cvo2m'
print(x, include_matrix = FALSE, ...)

Arguments

x List of class "cvo2m", produced by crossval_o2m.
include_matrix Logical. Should the 3-d array with Prediction errors also be printed.
... For consistency.
**print.o2m**  
*Print function for O2PLS.*

**Description**

This function is the print method for an O2PLS fit.

**Usage**

```r
## S3 method for class 'o2m'
print(x, ...)```

**Arguments**

- `x`  
  An O2PLS fit (an object of class o2m)
- `...`  
  For consistency

---

**rmsep**  
*Root MSE of Prediction*

**Description**

Calculates the Root MSE of prediction on test data. Only tested to work inside `loocv`.

**Usage**

```r
rmsep(Xtst, Ytst, fit, combi = FALSE)
```

**Arguments**

- `Xtst`  
  Numeric vector or matrix.
- `Ytst`  
  Numeric vector or matrix.
- `fit`  
  o2m fit (on data without Xtst and Ytst).
- `combi`  
  Logical. Should the symmetrized MSE be used, i.e. add both MSEs. Not yet implemented, but see `rmsep_combi`

**Details**

This function is the building block for `loocv`, as it produced the prediction error for test (left out) data.

**Value**

Mean squares difference between predicted Y and true Y.
**rmsep_combi**  
*Symmetrized root MSE of Prediction*

**Description**
Calculates the symmetrized root MSE of prediction on test data. *Expected* to work in combination with loocv.

**Usage**

\[
\text{rmsep_combi}(X_{\text{tst}}, Y_{\text{tst}}, \text{fit})
\]

**Arguments**
- **Xtst**
  - Numeric vector or matrix.
- **Ytst**
  - Numeric vector or matrix.
- **fit**
  - o2m fit (on data without \(X_{\text{tst}}\) and \(Y_{\text{tst}}\)).

**Details**
This function is the building block for loocv, as it produced the prediction error for test (left out) data.
This is a symmetrized version of rmsep and is used by loocv. The prediction error of both \(X_{\text{tst}}\) and \(Y_{\text{tst}}\) are calculated and summed. Whether this is a good idea depends: If \(X\) and \(Y\) have similar meanings (LC-MS versus MALDI) this is a good thing to do. If the two matrices do not have similar meanings, (Metabolomics versus Transcriptomics) then you may want to not sum up the two prediction errors or include weights in the sum.

**Value**
Mean squares difference between predicted \(Y\) and true \(Y\)

**scores**  
*Extract the scores from an O2PLS fit*

**Description**
This function extracts score matrices from an O2PLS fit

**Usage**

\[
\text{scores}(x, \ldots)
\]

## S3 method for class 'o2m'
scores(x, which_part = c("Xjoint", "Yjoint", "Xorth", "Yorth"), subset = 0, \ldots)
ssq

Arguments

x Object of class o2m
...
which_part character string. One of the following: 'Xjoint', 'Yjoint', 'Xorth' or 'Yorth'.
subset subset of scores vectors to be extracted.

Value

Scores matrix

See Also

loadings.o2m

Examples

scores(o2m(scale(-2:2),scale(-2:2*4),1,0,0))

ssq Calculate Sum of Squares

Description

Calculate Sum of Squares

Usage

ssq(X)

Arguments

x Numeric vector or matrix.

Details

This is the Frobenius norm of X.

Value

The sum of squared elements of X

Examples

ssq(tcrossprod(1:5))
ssq(rnorm(1e5))/1e5
**summary.o2m**  
*Summary of an O2PLS fit*

---

**Description**

Until now only variational summary given by the R2's is outputted

**Usage**

```r
## S3 method for class 'o2m'
summary(object, digits = 3, ...)
```

**Arguments**

- `object`  
  List. Should be of class o2m.
- `digits`  
  Integer, number of digits.
- `...`  
  For compatibility

**Value**

List with R2 values.

**See Also**

`plot.o2m`

**Examples**

```r
summary(o2m(scale(-2:2),scale(-2:2*4),1,0,0))
```

---

**vnorm**  
*Norm of a vector or columns of a matrix*

---

**Description**

Norm of a vector or columns of a matrix

**Usage**

```r
vnorm(x)
```

**Arguments**

- `x`  
  Numeric vector or matrix.
**vnorm**

**Value**

(columnwise) Euclidian norm of \( x \)

**Examples**

\[
\text{vnorm(orth(1:5))} \\
\text{vnorm(matrix(1:9,3,3))^2 - colSums(matrix(1:9,3)^2)}
\]