

Package ‘cellWise’

September 23, 2025

Type Package

Version 2.5.4

Date 2025-09-23

Title Analyzing Data with Cellwise Outliers

Depends R (>= 4.0.0)

Suggests knitr, robustHD, MASS, ellipse, markdown, rospca, GSE

Imports reshape2, scales, ggplot2, matrixStats, gridExtra, robustbase,
rrcov, svd, stats, utils, shape, Rcpp (>= 0.12.10.14)

LinkingTo Rcpp, RcppArmadillo (>= 0.7.600.1.0)

Description Tools for detecting cellwise outliers and robust methods to analyze data which may contain them. Contains the implementation of the algorithms described in Rousseeuw and Van den Bossche (2018) <doi:10.1080/00401706.2017.1340909> (open access) Hubert et al. (2019) <doi:10.1080/00401706.2018.1562989> (open access), Raymaekers and Rousseeuw (2021) <doi:10.1080/00401706.2019.1677270> (open access), Raymaekers and Rousseeuw (2021) <doi:10.1007/s10994-021-05960-5> (open access), Raymaekers and Rousseeuw (2021) <doi:10.52933/jdssv.v1i3.18> (open access), Raymaekers and Rousseeuw (2022) <doi:10.1080/01621459.2023.2267777> (open access) Rousseeuw (2022) <doi:10.1016/j.ecosta.2023.01.007> (open access).
Examples can be found in the vignettes:
``DDC\_examples``, ``MacroPCA\_examples``, ``wrap\_examples``, ``transfo\_examples``,
``DI\_examples``, ``cellMCD\_examples``, ``Correspondence\_analysis\_examples``,
and ``cellwise\_weights\_examples``.

License GPL (>= 2)

LazyData No

Author Jakob Raymaekers [aut, cre],
Peter Rousseeuw [aut],
Wannes Van den Bossche [ctb],
Mia Hubert [ctb]

Maintainer Jakob Raymaekers <jakob.raymaekers@kuleuven.be>

VignetteBuilder knitr

RoxygenNote 7.1.2

NeedsCompilation yes

Repository CRAN

Date/Publication 2025-09-23 14:00:02 UTC

Contents

cellHandler	3
cellMap	4
cellMCD	7
checkDataSet	10
cwLocScat	11
data_brands	13
data_clothes	14
data_dogWalker	15
data_dposs	15
data_glass	16
data_mortality	17
data_personality_traits	17
data_philips	18
data_VOC	19
DDC	20
DDCpredict	24
DI	26
estLocScale	28
generateCorMat	30
generateData	31
ICPCA	32
MacroPCA	34
MacroPCApredict	37
outlierMap	39
plot_cellMCD	40
transfo	42
transfo_newdata	45
transfo_transformback	46
truncPC	47
unpack	48
weightedEM	49
wrap	51

Index	54
--------------	-----------

cellHandler*cellHandler algorithm*

Description

This function flags cellwise outliers in X and imputes them, if robust estimates of the center μ and scatter matrix Σ are given. When the latter are not known, as is typically the case, one can use the function [DDC](#) which only requires the data matrix X . Alternatively, the unknown center μ and scatter matrix Σ can be estimated robustly from X by the function [DI](#).

Usage

```
cellHandler(X, mu, Sigma, quant = 0.99)
```

Arguments

X	X is the input data, and must be an n by d matrix or a data frame.
μ	An estimate of the center of the data
Σ	An estimate of the covariance matrix of the data
quant	Cutoff used in the detection of cellwise outliers. Defaults to 0.99

Value

A list with components:

- `Ximp`
The imputed data matrix.
- `indcells`
Indices of the cells which were flagged in the analysis.
- `indNAs`
Indices of the NAs in the data.
- `Zres`
Matrix with standardized cellwise residuals of the flagged cells. Contains zeroes in the un-flagged cells.
- `Zres_denom`
Denominator of the standardized cellwise residuals.
- `cellPaths`
Matrix with the same dimensions as X , in which each row contains the path of least angle regression through the cells of that row, i.e. the order of the coordinates in the path (1=first, 2=second,...)

Author(s)

J. Raymaekers and P.J. Rousseeuw

References

J. Raymaekers and P.J. Rousseeuw (2020). Handling cellwise outliers by sparse regression and robust covariance. *Journal of Data Science, Statistics, and Visualisation*. doi:[10.52933/jdssv.v1i3.18](https://doi.org/10.52933/jdssv.v1i3.18)(link to open access pdf)

See Also

[DI](#)

Examples

```
mu <- rep(0, 3)
Sigma <- diag(3) * 0.1 + 0.9
X <- rbind(c(0.5, 1.0, 5.0), c(-3.0, 0.0, 1.0))
n <- nrow(X); d <- ncol(X)
out <- cellHandler(X, mu, Sigma)
Xres <- X - out$Ximp # unstandardized residual
mean(abs(as.vector(Xres - out$Zres*out$Zres_denom))) # 0
W <- matrix(rep(0,n*d),nrow=n) # weight matrix
W[out$Zres != 0] <- 1 # 1 indicates cells that were flagged
# For more examples, we refer to the vignette:
## Not run:
vignette("DI_examples")

## End(Not run)
```

cellMap

Draw a cellmap

Description

This function draws a cellmap, possibly of a subset of rows and columns of the data, and possibly combining cells into blocks. A cellmap shows which cells are missing and which ones are outlying, marking them in red for unusually large cell values and in blue for unusually low cell values. When cells are combined into blocks, the final color is the average of the colors in the individual cells.

Usage

```
cellMap(R, indcells = NULL, indrows = NULL, outrows = NULL,
  showcellvalues = NULL, D = NULL, rowlabels = NULL,
  columnlabels = NULL, mTitle = "cell map",
  rowtitle = "cases", columntitle = "variables",
  showrows = NULL, showcolumns = NULL,
  nrowsinblock = NULL, ncolumnsinblock = NULL,
  manualrowblocksizes = NULL,
  manualcolumnblocksizes = NULL,
  rowblocklabels = NULL, columnblocklabels = NULL,
  sizemain = 1.5, sizetitles = 1.2, sizerowlabels = 1,
```

```

sizecolumnlabels = 1, sizecellvalues = 1,
adjustrowlabels = 1, adjustcolumnlabels = 1,
columnangle = 90, colContrast = 1,
outlyingGrad = TRUE,
darkestColor = sqrt(qchisq(0.999, 1)),
drawCircles = FALSE, showVals = NULL, autolabel = TRUE)

```

Arguments

R	Matrix of standardized residuals of the cells (required input argument). After running DDC , DDCpredict , MacroPCA or MacroPCApredict this is typically their value \$stdResid.
indcells	Indices of flagged cells. Defaults to NULL, which flags the cells for which $ R > \sqrt{(qchisq(0.99, 1))}$.
indrows	Indices of outlying rows (if available). If not NULL, the small circle to the right of the row is filled black if the row is in this list, and white otherwise. This gets overruled if outrows is not NULL.
outrows	Outlyingness of each row (if available). If not NULL, represents the outlyingness of each row by a shade of gray in the small circle to the right of the row. This color is white for outrows below 1, and becomes fully black for outrows over 3.
showcellvalues	Takes the values "D", "R" or NULL (the default). If "R" the numerical values of the residuals in R are shown in the cellmap. If "D", the entries of the data matrix D are shown, provided the matrix D is being specified. If NULL, no entries are shown.
D	A matrix of data values, of the same dimensions as R. Default is NULL. D is only required when the data values are to be shown in the cellmap, by the option showcellvalues = "D". After running DDC or MacroPCA , D is typically their value \$remX. After running DDCpredict or MacroPCApredict it is their argument \$newX.
rowlabels	Labels of the rows of the matrix R. If NULL, these labels are taken as rownames(R), and failing that they are 1:nrow(R).
columnlabels	Labels of the columns of the matrix R. If NULL, these labels are taken as colnames(R), and failing that they are 1:ncol(R).
mTitle	Main title of the cellMap. Defaults to "cell map".
rowtitle	Title for the rows. Defaults to "cases".
columntitle	Title for the columns. Defaults to "variables".
showrows	Indices of the rows to be shown. Defaults to NULL which means all rows are shown.
showcolumns	Indices of the columns to be shown. Defaults to NULL which means all columns are shown.
nrowsinblock	How many rows are combined in a block. Defaults to NULL, which asks not to block rows. The argument nrowsinblock is overruled by the argument manualrowblocksizes when the latter is specified.

<code>ncolumnsinblock</code>	Defaults to NULL, which asks not to block columns. The argument <code>ncolumnsinblock</code> is overruled by the argument <code>manualcolumnblocksizes</code> when the latter is specified.
<code>manualrowblocksizes</code>	This allows the user to specify their own row blocks, unlike the argument <code>nrowsinblock</code> which makes all row blocks the same length. The argument takes the form <code>c(a,b,...)</code> where <code>a</code> is the length of the first block, <code>b</code> is the length of the second, and so on. The numbers <code>a,b,...</code> must be strictly positive integers, adding up to at most <code>nrow(R)</code> . They cannot all be 1, which would mean no blocking of rows. Defaults to NULL.
<code>manualcolumnblocksizes</code>	Analogous to <code>manualrowblocksizes</code> but for columns. It is allowed for one of them to be NULL while the other is not.
<code>rowblocklabels</code>	This allows the user to specify labels for the row blocks, whether obtained from <code>nrowsinblock</code> or from <code>manualrowblocksizes</code> . Defaults to NULL, and then labels will be created automatically. Will throw an error if the number of row labels does not match the number of blocks.
<code>columnblocklabels</code>	Analogous to <code>rowblocklabels</code> but for columns. It is allowed for one of them to be NULL while the other is not.
<code>sizemain</code>	Size of main title. Defaults to 1.5.
<code>sizetitles</code>	Size of row title and column title. Defaults to 1.2.
<code>sizerowlabels</code>	Size of row labels. Defaults to 1.
<code>sizecolumnlabels</code>	Size of column labels. Defaults to 1.
<code>sizecellvalues</code>	Size of values in the cells, when <code>showcellvalues = TRUE</code> . Defaults to 1.
<code>adjustrowlabels</code>	Adjust row labels: 0=left, 0.5=centered, 1=right. Defaults to 1.
<code>adjustcolumnlabels</code>	Adjust column labels: 0=left, 0.5=centered, 1=right. Defaults to 1.
<code>columnangle</code>	Angle of the column labels. Defaults to 90 so the column labels are vertical.
<code>colContrast</code>	Parameter regulating the contrast of colors, should be in $[1, 5]$. Defaults to 1.
<code>outlyingGrad</code>	If TRUE, the color is gradually adjusted in function of the outlyingness. Defaults to TRUE.
<code>darkestColor</code>	Standardized residuals whose absolute value is bigger than this will get the darkest color.
<code>drawCircles</code>	Whether or not to draw circles indicating outlyingness of rows. When both <code>indrows</code> and <code>outrows</code> are NULL, no circles are drawn.
<code>showVals</code>	old name of argument <code>showcellvalues</code> . Only for backward compatibility.
<code>autolabel</code>	obsoleted by the current mechanism for creating blocks of cells. Is only in the list for backward compatibility.

Author(s)

Rousseeuw P.J., Van den Bossche W.

References

Rousseeuw, P.J., Van den Bossche W. (2018). Detecting Deviating Data Cells. *Technometrics*, **60**(2), 135-145. ([link to open access pdf](#))

See Also

[DDC](#)

Examples

```
# For examples of the cellmap, we refer to the vignette:
## Not run:
vignette("DDC_examples")

## End(Not run)
```

cellMCD	<i>cellWise minimum covariance determinant estimator</i>
---------	--

Description

The cellwise minimum covariance determinant estimator computes cellwise robust estimates of the center and covariance matrix of a data set X . The algorithm guarantees a monotone decrease of an objective function, which is based on observed Gaussian log-likelihood. By default, it starts by calling [checkDataSet](#) to clean the data.

Usage

```
cellMCD(X, alpha = 0.75, quant = 0.99,
        crit = 1e-4, noCits = 100, lmin = 1e-4,
        fixedCenter = FALSE, checkPars = list())
```

Arguments

<code>X</code>	X is the input data, and must be an n by d matrix or a data frame.
<code>alpha</code>	In each column, at least $n \cdot \alpha$ cells must remain unflagged. Defaults to 75%, should not be set (much) lower.
<code>quant</code>	Determines the cutoff value to flag cells. Defaults to 0.99.
<code>crit</code>	The iteration stops when successive covariance matrices (of the standardized data) differ by less than <code>crit</code> . Defaults to $1e - 4$.
<code>noCits</code>	The maximal number of C-steps used.
<code>lmin</code>	a lower bound on the eigenvalues of the estimated covariance matrix on the standardized data. Defaults to $1e - 4$. Should not be smaller than $1e - 6$.
<code>fixedCenter</code>	if TRUE, cellMCD is fit around a fixed center in the origin.
<code>checkPars</code>	Optional list of parameters used in the call to checkDataSet . The options are:

- **coreOnly**
If TRUE, skip the execution of checkDataset. Defaults to FALSE.
- **numDiscrete**
A column that takes on numDiscrete or fewer values will be considered discrete and not retained in the cleaned data. Defaults to 5.
- **fracNA**
Only retain columns and rows with fewer NAs than this fraction. Defaults to 0.5.
- **precScale**
Only consider columns whose scale is larger than precScale. Here scale is measured by the median absolute deviation. Defaults to $1e - 12$.
- **silent**
Whether or not the function progress messages should be suppressed. Defaults to FALSE.

Details

The matrix `raw.S` in the output is the raw estimate of scatter produced by cellMCD. The final `S` is obtained from `raw.S` by rescaling such that its diagonal entries equal the squares of the univariate scales in `locsca$scale`. This reduces the bias at Gaussian data, which matters mainly for large sample sizes.

Value

A list with components:

- **mu**
the cellMCD estimate of location.
- **S**
the cellMCD estimate of scatter, after bias correction (see details).
- **W**
the cellMCD estimate of `W`, a binary matrix indicating all outlying cells as zero.
- **preds**
predictions (=conditional expectations) of the flagged cells, given the clean cells in the same row.
- **csds**
conditional standard deviations of the flagged cells, given the clean cells in the same row.
- **Ximp**
imputed data matrix.
- **Zres**
matrix of cellwise standardized residuals.
- **raw.S**
the raw cellMCD estimate of scatter, without bias correction.
- **locsca**
list containing robust locations and scales used to standardize the data before running the algorithm. The results `m`, `S`, `preds`, `Ximp` are returned in their original location/scale.

- `nosteps`
number of steps the algorithm took to converge.
- `X`
the data on which the algorithm was executed.
- `quant`
the cutoff used to flag the cells.

Author(s)

J. Raymaekers and P.J. Rousseeuw

References

J. Raymaekers and P.J. Rousseeuw (2022). The cellwise MCD estimator, Journal of the American Statistical Association, to appear. [doi:10.1080/01621459.2023.2267777](https://doi.org/10.1080/01621459.2023.2267777)(link to open access pdf)

See Also

[plot_cellMCD](#)

Examples

```
mu    <- rep(0, 3)
Sigma <- diag(3) * 0.5 + 0.5
set.seed(123)
X <- MASS::mvrnorm(1000, mu, Sigma)
X[1:5, 1] <- X[1:5, 1] + 5
X[6:10, 2] <- X[6:10, 2] - 10
X[12, 1:2] <- c(-4, 8)
colnames(X) <- c("X1", "X2", "X3")
cellMCD.out <- cellMCD(X)
cellMCD.out$mu
cov2cor(cellMCD.out$S)
cellMCD.out$W[1:15,]
cellMCD.out$Ximp[1:15,]
cellMap(cellMCD.out$Zres[1:15,])

# For more examples, we refer to the vignette:
## Not run:
vignette("cellMCD_examples")

## End(Not run)
```

checkDataSet

*Clean the dataset***Description**

This function checks the dataset X , and sets aside certain columns and rows that do not satisfy the conditions. It is used by the [DDC](#) and [MacroPCA](#) functions but can be used by itself, to clean a dataset for a different type of analysis.

Usage

```
checkDataSet(X, fracNA = 0.5, numDiscrete = 3, precScale = 1e-12, silent = FALSE,
cleanNAfirst = "automatic")
```

Arguments

<code>X</code>	X is the input data, and must be an n by d matrix or data frame.
<code>fracNA</code>	Only retain columns and rows with fewer NAs than this fraction. Defaults to 0.5.
<code>numDiscrete</code>	A column that takes on <code>numDiscrete</code> or fewer values will be considered discrete and not retained in the cleaned data. Defaults to 3.
<code>precScale</code>	Only consider columns whose scale is larger than <code>precScale</code> . Here scale is measured by the median absolute deviation. Defaults to $1e-12$.
<code>silent</code>	Whether or not the function progress messages should be printed. Defaults to FALSE.
<code>cleanNAfirst</code>	If "columns", first columns then rows are checked for NAs. If "rows", first rows then columns are checked for NAs. "automatic" checks columns first if $d \geq 5n$ and rows first otherwise. Defaults to "automatic".

Value

A list with components:

- `colInAnalysis`
Column indices of the columns used in the analysis.
- `rowInAnalysis`
Row indices of the rows used in the analysis.
- `namesNotNumeric`
Names of the variables which are not numeric.
- `namesCaseNumber`
The name of the variable(s) which contained the case numbers and was therefore removed.
- `namesNAcol`
Names of the columns left out due to too many NA's.

- `namesNArow`
Names of the rows left out due to too many NA's.
- `namesDiscrete`
Names of the discrete variables.
- `namesZeroScale`
Names of the variables with zero scale.
- `remX`
Remaining (cleaned) data after `checkDataSet`.

Author(s)

Rousseeuw P.J., Van den Bossche W.

References

Rousseeuw, P.J., Van den Bossche W. (2018). Detecting Deviating Data Cells. *Technometrics*, **60**(2), 135-145. ([link to open access pdf](#))

See Also

[DDC](#), [MacroPCA](#), [transfo](#), [wrap](#)

Examples

```
library(MASS)
set.seed(12345)
n <- 100; d = 10
A <- matrix(0.9, d, d); diag(A) = 1
x <- mvrnorm(n, rep(0,d), A)
x[sample(1:(n * d), 100, FALSE)] <- NA
x <- cbind(1:n, x)
checkedx <- checkDataSet(x)

# For more examples, we refer to the vignette:
## Not run:
vignette("DDC_examples")

## End(Not run)
```

Description

Computes different estimators of multivariate location and scatter for cellwise weighted data.

Usage

```

cwLocScat(X, W, methods = "all", lmin = 1e-3,
          crit = 1e-12, maxiter= 1000,
          initCwCov = FALSE, initEst = NULL)

```

Arguments

X	An n by d data matrix or data frame. Must be given. X is allowed to contain NA's.
W	An n by d matrix of nonnegative cellwise weights. Must be given. W is not allowed to contain NA's.
methods	either "all" or "explicit". If "explicit" only the explicit estimates <code>cwMean</code> , <code>cwCov</code> and <code>sqrtCov</code> are computed. If "all" (the default) also the cellwise MLE is carried out, yielding <code>cwMLEmu</code> and <code>cwMLEsigma</code> .
lmin	if not NULL, a lower bound on the eigenvalues of the estimated covariance matrices on the standardized data, to avoid singularity.
crit	convergence criterion of successive μ and Σ estimates in the EM algorithm.
maxiter	maximal number of iteration steps in EM.
initCwCov	if TRUE, uses the weighted mean and <code>cwCov</code> as initial estimates for the weighted EM.
initEst	if not NULL, a list with initial estimates μ of the mean, Σ of the covariance matrix, for the weighted EM. Has no effect when <code>initCwCov = TRUE</code> .

Value

A list with components:

- `cwMean`
the explicit cellwise weighted mean.
- `cwCov`
explicit cellwise weighted covariance matrix. Is asymptotically normal but not necessarily PSD (unless a nonnegative `lmin` was specified).
- `sqrtCov`
the cellwise weighted covariance matrix of Van Aelst et al (2011). Also asymptotically normal but not necessarily PSD (unless a nonnegative `lmin` was specified).
- `cwMLEmu`
the location estimate obtained by the cwMLE.
- `cwMLEsigma`
the covariance matrix obtained by the cwMLE. Is PSD when the EM algorithm converges.

Author(s)

P.J. Rousseeuw

References

P.J. Rousseeuw (2022). Analyzing cellwise weighted data, ArXiv:2209.12697. ([link to open access pdf](#))

See Also

[weightedEM](#), [unpack](#)

Examples

```
data("data_personality_traits")
X <- data_personality_traits$X
W <- data_personality_traits$W
fit <- cwLocScat(X, W)
fit$cwMLEiter # number of iteration steps taken
round(fit$cwMLEmu, 2)
round(fit$cwMean, 2)
round(fit$cwMLEsigma, 2)
round(fit$cwCov, 2)

# For more examples, we refer to the vignette:
## Not run:
vignette("cellwise_weights_examples")

## End(Not run)
```

data_brands

The brands dataset

Description

The brands data is a contingency table summarizing the 2014 Auto Brand Perception survey by Consumer Reports (USA), which is publicly available on <https://boraberan.wordpress.com/2016/09/22/>. The survey questioned 1578 participants on what they considered attributes of 39 different car brands.

Usage

```
data("data_brands")
```

Format

A matrix with 39 observations of 7 attributes. The attributes (columns) are Fuel Economy, Innovation, Performance, Quality, Safety, Style and Value.

Source

<https://boraberan.wordpress.com/2016/09/22/>.

References

Riani, M., Atkinson, A. C., Torti, F., Corbellini, A. (2022). Robust correspondence analysis. *Journal of the Royal Statistical Society Series C: Applied Statistics*, **71**(5), 1381–1401.

Raymaekers and Rousseeuw (2022), Challenges of cellwise outliers.

Examples

```
data(data_brands)
```

data_clothes	<i>The clothes dataset</i>
--------------	----------------------------

Description

The clothes dataset contains a contingency table of trade flows from outside the European Union into each of its 28 member states. The columns in the contingency table in Riani et al. (2022) are five different price brackets, from lowest to highest.

Usage

```
data("data_clothes")
```

Format

A matrix with 28 observations of 5 price brackets.

Source

Riani, M., Atkinson, A. C., Torti, F., Corbellini, A. (2022). Robust correspondence analysis. *Journal of the Royal Statistical Society Series C: Applied Statistics*, **71**(5), 1381–1401.

References

Raymaekers and Rousseeuw (2022), Challenges of cellwise outliers.

Examples

```
data(data_clothes)
```

data_dogWalker	<i>Dog walker dataset</i>
----------------	---------------------------

Description

A dataset containing the image sequence of a video. The sequence consists of 54 frames of 144 by 180 pixels pixels in Red/Geen/Blue (RGB) format.

Usage

```
data("data_dogWalker")
```

Format

An array of dimensions $54 \times 144 \times 180 \times 3$.

Source

<http://www.wisdom.weizmann.ac.il/~vision/SpaceTimeActions.html>

Examples

```
data("data_dogWalker")
# For more examples, we refer to the vignette:
## Not run:
vignette("Wrap_examples")

## End(Not run)
```

data_dposs	<i>DPOSS dataset</i>
------------	----------------------

Description

This is a random subset of 20'000 stars from the Digitized Palomar Sky Survey (DPOSS) described by Odewahn et al. (1998).

Usage

```
data("data_dposs")
```

Format

A matrix of dimensions 20000×21 .

References

Odewahn, S., S. Djorgovski, R. Brunner, and R. Gal (1998). Data From the Digitized Palomar Sky Survey. Technical report, California Institute of Technology.

Examples

```
data("data_dposs")
# For more examples, we refer to the vignette:
## Not run:
vignette("MacroPCA_examples")

## End(Not run)
```

data_glass

The glass dataset

Description

A dataset containing spectra with $d = 750$ wavelengths collected on $n = 180$ archeological glass samples.

Usage

```
data("data_glass")
```

Format

A data frame with 180 observations of 750 wavelengths.

Source

Lemberge, P., De Raedt, I., Janssens, K.H., Wei, F., and Van Espen, P.J. (2000). Quantitative Z-analysis of 16th-17th century archaeological glass vessels using PLS regression of EPXMA and μ -XRF data. *Journal of Chemometrics*, **14**, 751–763.

Examples

```
data("data_glass")
```

data_mortality	<i>The mortality dataset</i>
----------------	------------------------------

Description

This dataset contains the mortality by age for males in France, from 1816 to 2013 as obtained from the Human Mortality Database.

Usage

```
data("data_mortality")
```

Format

A data frame with 198 calendar years (rows) and 91 age brackets (columns).

Source

Human Mortality Database. University of California, Berkeley (USA), and Max Planck Institute for Demographic Research (Germany). Available at <https://www.mortality.org> (data downloaded in November 2015).

References

Hyndman, R.J., and Shang, H.L. (2010), Rainbow plots, bagplots, and boxplots for functional data, *Journal of Computational and Graphical Statistics*, **19**, 29–45.

Examples

```
data("data_mortality")
```

data_personality_traits	<i>The personality traits data</i>
-------------------------	------------------------------------

Description

This dataset describes personality traits of 10 persons. The variables are the 6 traits Anxiety, Agoraphobia, Arachnophobia, Adventurous, Extraversion, and Sociability.

Usage

```
data("data_personality_traits")
```

Format

The data contains a list with two elements:

- X
a 10 by 6 matrix of values describing 6 personality traits for each of the 10 participants.
- W
a 10 by 6 matrix of cellwise weights. Each weight is the inverse of the length of the support of the membership function of the fuzzy number in the original data set.

Source

G. Hesamian, and Akbari, M. G. (2019), Principal component analysis based on intuitionistic fuzzy random variables, *Computational and Applied Mathematics*, **38**(158), 1–14.

References

P.J. Rousseeuw (2022). Analyzing cellwise weighted data, ArXiv:2209.12697. ([link to open access pdf](#))

Examples

```
data(data_personality_traits)

# For the examples in Rousseeuw (2022), see:
## Not run:
vignette("cellwise_weights_examples")

## End(Not run)
```

data_philips	<i>The philips dataset</i>
--------------	----------------------------

Description

A dataset containing measurements of $d = 9$ characteristics of $n = 677$ diaphragm parts, used in the production of TV sets.

Usage

```
data("data_philips")
```

Format

A matrix with 677 rows and 9 columns.

Source

The data were provided in 1997 by Gertjan Otten and permission to analyze them was given by Herman Veraa and Frans Van Dommelen at Philips Mecoma in The Netherlands.

References

Rousseeuw, P.J., and Van Driessen, K. (1999). A fast algorithm for the Minimum Covariance Determinant estimator. *Technometrics*, **41**, 212–223.

Examples

```
data("data_philips")
```

data_VOC	<i>VOC dataset</i>
----------	--------------------

Description

This dataset contains the data on volatile organic components (VOCs) in urine of children between 3 and 10 years old. It is composed of publicly available data from the National Health and Nutrition Examination Survey (NHANES) and was analyzed in Raymaekers and Rousseeuw (2020). See below for details and references.

Usage

```
data("data_VOC")
```

Format

A matrix of dimensions 512×19 . The first 16 variables are the VOC, the last 3 are:

- SMD460: number of smokers that live in the same home as the subject
- SMD470: number of people that smoke inside the home of the subject
- RIDAGEYR: age of the subject

Note that the original variable names are kept.

Details

All of the data was collected from the NHANES website, and was part of the NHANES 2015-2016 survey. This was the most recent epoch with complete data at the time of extraction. Three datasets were matched in order to assemble this data:

- UVOC_I: contains the information on the Volative organic components in urine
- DEMO_I: contains the demographical information such as age
- SMQFAM_I: contains the data on the smoking habits of family members

The dataset was constructed as follows:

1. Select the relevant VOCs from the UVOC_I data (see column names) and transform by taking the logarithm
2. Match the subjects in the UVOC_I data with their age in the DEMO_I data
3. Select all subjects with age at most 10
4. Match the data on smoking habits with the selected subjects.

Source

<https://wwwn.cdc.gov/nchs/nhanes/Search/DataPage.aspx?Component=Laboratory&CycleBeginYear=2015>

<https://wwwn.cdc.gov/nchs/nhanes/search/datapage.aspx?Component=Demographics&CycleBeginYear=2015>

<https://wwwn.cdc.gov/nchs/nhanes/Search/DataPage.aspx?Component=Questionnaire&CycleBeginYear=2015>

References

J. Raymaekers and P.J. Rousseeuw (2020). Handling cellwise outliers by sparse regression and robust covariance. *Journal of Data Science, Statistics, and Visualisation*. doi:10.52933/jdssv.v1i3.18(link to open access pdf)

Examples

```
data("data_VOC")
# For an analysis of this data, we refer to the vignette:
## Not run:
vignette("DI_examples")

## End(Not run)
```

DDC

Detect Deviating Cells

Description

This function aims to detect cellwise outliers in the data. These are entries in the data matrix which are substantially higher or lower than what could be expected based on the other cells in its column as well as the other cells in its row, taking the relations between the columns into account. Note that this function first calls `checkDataSet` and analyzes the remaining cleaned data.

Usage

```
DDC(X, DDCpars = list())
```

Arguments

X	X is the input data, and must be an n by d matrix or a data frame.
DDCpars	A list of available options: • <code>fracNA</code> Only consider columns and rows with fewer NAs (missing values) than this fraction (percentage). Defaults to 0.5. • <code>numDiscrete</code> A column that takes on <code>numDiscrete</code> or fewer values will be considered discrete and not used in the analysis. Defaults to 3.

- **precScale**
Only consider columns whose scale is larger than `precScale`. Here scale is measured by the median absolute deviation. Defaults to $1e - 12$.
- **cleanNAfirst**
If "columns", first columns then rows are checked for NAs. If "rows", first rows then columns are checked for NAs. "automatic" checks columns first if $d \geq 5n$ and rows first otherwise. Defaults to "automatic".
- **tolProb**
Tolerance probability, with default 0.99, which determines the cutoff values for flagging outliers in several steps of the algorithm.
- **corrlim**
When trying to estimate z_{ij} from other variables h , we will only use variables h with $|\rho_{j,h}| \geq \text{corrlim}$. Variables j without any correlated variables h satisfying this are considered standalone, and treated on their own. Defaults to 0.5.
- **combinRule**
The operation to combine estimates of z_{ij} coming from other variables h : can be "mean", "median", "wmean" (weighted mean) or "wmedian" (weighted median). Defaults to wmean.
- **returnBigXimp**
If TRUE, the imputed data matrix `Ximp` in the output will include the rows and columns that were not part of the analysis (and can still contain NAs). Defaults to FALSE.
- **silent**
If TRUE, statements tracking the algorithm's progress will not be printed. Defaults to FALSE.
- **nLocScale**
When estimating location or scale from more than `nLocScale` data values, the computation is based on a random sample of size `nLocScale` to save time. When `nLocScale = 0` all values are used. Defaults to 25000.
- **fastDDC**
Whether to use the fastDDC option or not. The fastDDC algorithm uses approximations to allow to deal with high dimensions. Defaults to TRUE for $d > 750$ and FALSE otherwise.
- **standType**
The location and scale estimators used for robust standardization. Should be one of "1stepM", "mcd" or "wrap". See [estLocScale](#) for more info. Only used when `fastDDC = FALSE`. Defaults to "1stepM".
- **corrType**
The correlation estimator used to find the neighboring variables. Must be one of "wrap" (wrapping correlation), "rank" (Spearman correlation) or "gkwl" (Gnanadesikan-Kettenring correlation followed by weighting). Only used when `fastDDC = FALSE`. Defaults to "gkwl".
- **transFun**
The transformation function used to compute the robust correlations when `fastDDC = TRUE`. Can be "wrap" or "rank". Defaults to "wrap".

- `nbngbrs`
When `fastDDC = TRUE`, each column is predicted from at most `nbngbrs` columns correlated to it. Defaults to 100.

Value

A list with components:

- `DDCpars`
The list of options used.
- `colInAnalysis`
The column indices of the columns used in the analysis.
- `rowInAnalysis`
The row indices of the rows used in the analysis.
- `namesNotNumeric`
The names of the variables which are not numeric.
- `namesCaseNumber`
The name of the variable(s) which contained the case numbers and was therefore removed.
- `namesNAcol`
Names of the columns left out due to too many NA's.
- `namesNArow`
Names of the rows left out due to too many NA's.
- `namesDiscrete`
Names of the discrete variables.
- `namesZeroScale`
Names of the variables with zero scale.
- `remX`
Cleaned data after `checkDataSet`.
- `locX`
Estimated location of X .
- `scaleX`
Estimated scales of X .
- `Z`
Standardized `remX`.
- `nbngbrs`
Number of neighbors used in estimation.
- `ngbrs`
Indicates neighbors of each column, i.e. the columns most correlated with it.
- `robcors`
Robust correlations.
- `robslopes`
Robust slopes.

- `deshrinkage`
The deshrinkage factor used for every connected (i.e. non-standalone) column of X .
- `Xest`
Predicted X .
- `scalestres`
Scale estimate of the residuals $X - X_{est}$.
- `stdResid`
Residuals of original X minus the estimated X_{est} , standardized by column.
- `indcells`
Indices of the cells which were flagged in the analysis.
- `Ti`
Outlyingness value of each row.
- `medTi`
Median of the T_i values.
- `madTi`
Mad of the T_i values.
- `indrows`
Indices of the rows which were flagged in the analysis.
- `indNAs`
Indices of all NA cells.
- `indall`
Indices of all cells which were flagged in the analysis plus all cells in flagged rows plus the indices of the NA cells.
- `Ximp`
Imputed X .

Author(s)

Raymaekers J., Rousseeuw P.J., Van den Bossche W.

References

- Rousseeuw, P.J., Van den Bossche W. (2018). Detecting Deviating Data Cells. *Technometrics*, **60**(2), 135-145. ([link to open access pdf](#))
- Raymaekers, J., Rousseeuw P.J. (2019). Fast robust correlation for high dimensional data. *Technometrics*, **63**(2), 184-198. ([link to open access pdf](#))

See Also

[checkDataSet](#), [cellMap](#)

Examples

```

library(MASS); set.seed(12345)
n <- 50; d <- 20
A <- matrix(0.9, d, d); diag(A) = 1
x <- mvrnorm(n, rep(0,d), A)
x[sample(1:(n * d), 50, FALSE)] <- NA
x[sample(1:(n * d), 50, FALSE)] <- 10
x[sample(1:(n * d), 50, FALSE)] <- -10
x <- cbind(1:n, x)
DDCx <- DDC(x)
cellMap(DDCx$stdResid)

# For more examples, we refer to the vignette:
## Not run:
vignette("DDC_examples")

## End(Not run)

```

DDCpredict

*DDCpredict***Description**

Based on a [DDC](#) fit on an initial (training) data set X , this function analyzes a new (test) data set X_{new} .

Usage

```
DDCpredict(Xnew, InitialDDC, DDCpars = NULL)
```

Arguments

<code>Xnew</code>	The new data (test data), which must be a matrix or a data frame. It must always be provided. Its columns (variables) should correspond to those of <code>InitialDDC\$remX</code> .
<code>InitialDDC</code>	The output of the DDC function on the initial (training) dataset. Must be provided.
<code>DDCpars</code>	The input options to be used for the prediction. By default the options of <code>InitialDDC</code> are used.

Value

A list with components:

<code>DDCpars</code>	the options used in the call, see DDC .
<code>locX</code>	the locations of the columns, from <code>InitialDDC</code> .
<code>scaleX</code>	the scales of the columns, from <code>InitialDDC</code> .
<code>Z</code>	X_{new} standardized by <code>locX</code> and <code>scaleX</code> .

nbngbrs	predictions use a combination of nbngbrs columns.
ngbrs	for each column, the list of its neighbors, from InitialDDC.
robcors	for each column, the correlations with its neighbors, from InitialDDC.
robslopes	slopes to predict each column by its neighbors, from InitialDDC.
deshrinkage	for each connected column, its deshrinkage factor used in InitialDDC.
Xest	predicted values for every cell of Xnew.
scalestres	scale estimate of the residuals (Xnew - Xest), from InitialDDC.
stdResid	columnwise standardized residuals of Xnew.
indcells	positions of cellwise outliers in Xnew.
Ti	outlyingness of rows in Xnew.
medTi	median of the Ti in InitialDDC.
madTi	mad of the Ti in InitialDDC.
indrows	row numbers of the outlying rows in Xnew.
indNAs	positions of the NA's in Xnew.
indall	positions of NA's and outlying cells in Xnew.
Ximp	Xnew where all cells in indall are imputed by their prediction.

Author(s)

Rousseeuw P.J., Van den Bossche W.

References

Hubert, M., Rousseeuw, P.J., Van den Bossche W. (2019). MacroPCA: An all-in-one PCA method allowing for missing values as well as cellwise and rowwise outliers. *Technometrics*, **61**(4), 459-473. ([link to open access pdf](#))

See Also

[checkDataSet](#), [cellMap](#), [DDC](#)

Examples

```
library(MASS)
set.seed(12345)
n <- 100; d <- 10
A <- matrix(0.9, d, d); diag(A) = 1
x <- mvrnorm(n, rep(0,d), A)
x[sample(1:(n * d), 50, FALSE)] <- NA
x[sample(1:(n * d), 50, FALSE)] <- 10
x <- cbind(1:n, x)
DDCx <- DDC(x)
xnew <- mvrnorm(50, rep(0,d), A)
xnew[sample(1:(50 * d), 50, FALSE)] <- 10
predict.out <- DDCpredict(xnew, DDCx)
cellMap(D = xnew, R = predict.out$stdResid,
```

```

columnlabels = 1:d, rowlabels = 1:50)

# For more examples, we refer to the vignette:
## Not run:
vignette("DDC_examples")

## End(Not run)

```

DI

Detection-Imputation algorithm

Description

The Detection-Imputation algorithm computes cellwise robust estimates of the center and covariance matrix of a data set X . The algorithm alternates between the detection of cellwise outliers and their imputation combined with re-estimation of the center and covariance matrix. By default, it starts by calling [checkDataSet](#) to clean the data.

Usage

```

DI(X, initEst = "DDCWcov", crit = 0.01, maxits = 10, quant = 0.99,
maxCol = 0.25, checkPars = list())

```

Arguments

<code>X</code>	X is the input data, and must be an n by d matrix or a data frame.
<code>initEst</code>	An initial estimator for the center and covariance matrix. Should be one of "DDCWcov" or "TSGS", where the latter refers to the function <code>GSE::TSGS</code> . The default option "DDCWcov" uses the proposal of Raymaekers and Rousseeuw (2020) which is much faster for increasing dimension.
<code>crit</code>	The algorithm converges when the subsequent estimates of the center and covariance matrix do not differ more than <code>crit</code> in squared Euclidean norm.
<code>maxits</code>	Maximum number of DI-iterations.
<code>quant</code>	The cutoff used to detect cellwise outliers.
<code>maxCol</code>	The maximum number of cellwise outliers allowed in a column.
<code>checkPars</code>	Optional list of parameters used in the call to checkDataSet . The options are: • <code>coreOnly</code> If TRUE, skip the execution of <code>checkDataset</code>. Defaults to FALSE • <code>numDiscrete</code> A column that takes on <code>numDiscrete</code> or fewer values will be considered discrete and not retained in the cleaned data. Defaults to 5. • <code>fracNA</code> Only retain columns and rows with fewer NAs than this fraction. Defaults to 0.15. • <code>precScale</code> Only consider columns whose scale is larger than <code>precScale</code>. Here scale is measured by the median absolute deviation. Defaults to $1e - 12$.

- **silent**
Whether or not the function progress messages should be suppressed. Defaults to FALSE.

Value

A list with components:

- **center**
The final estimate of the center of the data.
- **cov**
The final estimate of the covariance matrix.
- **nits**
Number of DI-iterations executed to reach convergence.
- **Ximp**
The imputed data.
- **indcells**
Indices of the cells which were flagged in the analysis.
- **indNAs**
Indices of the NAs in the data.
- **Zres**
Matrix with standardized cellwise residuals of the flagged cells. Contains zeroes in the unflagged cells.
- **Zres_denom**
Denominator of the standardized cellwise residuals.
- **cellPaths**
Matrix with the same dimensions as *X*, in which each row contains the path of least angle regression through the cells of that row, i.e. the order of the coordinates in the path (1=first, 2=second,...)
- **checkDataSet_out**
Output of the call to [checkDataSet](#) which is used to clean the data.

Author(s)

J. Raymaekers and P.J. Rousseeuw

References

J. Raymaekers and P.J. Rousseeuw (2020). Handling cellwise outliers by sparse regression and robust covariance. *Journal of Data Science, Statistics, and Visualisation*. doi:10.52933/jdssv.v1i3.18(link to open access pdf)

See Also

[cellHandler](#)

Examples

```
mu <- rep(0, 3)
Sigma <- diag(3) * 0.1 + 0.9
X <- MASS::mvrnorm(100, mu, Sigma)
DI.out <- DI(X)
DI.out$cov
# For more examples, we refer to the vignette:
## Not run:
vignette("DI_examples")

## End(Not run)
```

estLocScale

Estimate robust location and scale

Description

Estimate a robust location estimate and scale estimate of every column in X .

Usage

```
estLocScale(X, type = "wrap", precScale = 1e-12,
center = TRUE, alpha = 0.5, nLocScale = 25000, silent = FALSE)
```

Arguments

X	The input data. It must be an n by d matrix or a data frame.
type	The type of estimators used. One of: • "1stepM": The location is the 1-step M-estimator with the biweight psi function. The scale estimator is the 1-step M-estimator using a Huber rho function with $b = 2.5$. • "mcd": the location is the weighted univariate MCD estimator with cutoff $\sqrt{(qchisq(0.975, 1))}$. The scale is the corresponding weighted univariate MCD estimator, with a correction factor to make it approximately unbiased at gaussian data. • "wrap": Starting from the initial estimates corresponding to option "mcd", the location is the 1-step M-estimator with the wrapping psi function with $b = 1.5$ and $c = 4$. The scale estimator is the same as in option "mcd". Defaults to "wrap".
precScale	The precision scale used throughout the algorithm. Defaults to $1e - 12$.
center	Whether or not the data has to be centered before calculating the scale. Not in use for type = "mcd". Defaults to TRUE.

alpha	The value of α in the univariate mcd, must be between 0.5 and 1. The subsetsize is $h = \lceil \alpha n \rceil$. Only used for <code>type = "mcd"</code> . Defaults to $\alpha = 0.5$.
nLocScale	If <code>nLocScale</code> $< n$, <code>nLocScale</code> observations are sampled to compute the location and scale. This speeds up the computation if n is very large. When <code>nLocScale</code> = 0 all observations are used. Defaults to <code>nLocScale</code> = 25000.
silent	Whether or not a warning message should be printed when very small scales are found. Defaults to FALSE.

Value

A list with components:

- `loc`
A vector with the estimated locations.
- `scale`
A vector with the estimated scales.

Author(s)

Raymaekers, J. and Rousseeuw P.J.

References

Raymaekers, J., Rousseeuw P.J. (2019). Fast robust correlation for high dimensional data. *Technometrics*, **63**(2), 184-198. ([link to open access pdf](#))

See Also

[wrap](#)

Examples

```
library(MASS)
set.seed(12345)
n = 100; d = 10
X = mvrnorm(n, rep(0, 10), diag(10))
locScale = estLocScale(X)
# For more examples, we refer to the vignette:
## Not run:
vignette("wrap_examples")

## End(Not run)
```

generateCorMat	<i>Generates correlation matrices</i>
----------------	---------------------------------------

Description

This function generates correlation matrices frequently used in simulation studies.

Usage

```
generateCorMat(d, corrType = "ALYZ", CN = 100, seed = NULL)
```

Arguments

d	The dimension of the correlation matrix. The resulting matrix is $d \times d$.
corrType	The type of correlation matrix to be generated. Should be one of: "ALYZ": Generates a correlation matrix as in Agostinelli et. al (2015). "A09": Generates the correlation matrix defined by $\rho_{jh} = (-0.9)^{ h-j }$. Note that the option "ALYZ" produces a randomly generated correlation matrix.
CN	Condition number of the correlation matrix. Only used for corrType = "ALYZ".
seed	Seed used in set.seed before generating the correlation matrix. Only relevant for corrType = "ALYZ".

Value

A $d \times d$ correlation matrix of the given type.

Author(s)

J. Raymaekers and P.J. Rousseeuw

References

C. Agostinelli, Leung, A., Yohai, V. J., and Zamar, R. H. (2015). Robust Estimation of Multivariate Location and Scatter in the Presence of Cellwise and Casewise Contamination. *Test*, 24, 441-461.

Rousseeuw, P.J., Van den Bossche W. (2018). Detecting Deviating Data Cells. *Technometrics*, **60**(2), 135-145. ([link to open access pdf](#))

J. Raymaekers and P.J. Rousseeuw (2020). Handling cellwise outliers by sparse regression and robust covariance. *Arxiv: 1912.12446*. ([link to open access pdf](#))

See Also

[generateData](#)

Examples

```
d      <- 5
Sigma <- generateCorMat(d, corrType = "ALYZ", seed = 1)
Sigma
```

generateData	<i>Generates artificial datasets with outliers</i>
--------------	--

Description

This function generates multivariate normal datasets with several possible types of outliers. It is used in several simulation studies. For a detailed description, see the referenced papers.

Usage

```
generateData(n, d, mu, Sigma, perout, gamma,
             outlierType = "casewise", seed = NULL)
```

Arguments

n	The number of observations
d	The dimension of the data.
mu	The center of the clean data.
Sigma	The covariance matrix of the clean data. Could be obtained from generateCorMat .
outlierType	The type of contamination to be generated. Should be one of: • "casewise": Generates point contamination in the direction of the last eigenvector of Sigma. • "cellwisePlain": Generates cellwise contamination by randomly replacing a number of cells by gamma. • "cellwiseStructured": Generates cellwise contamination by first randomly sampling contaminated cells, after which for each row, they are replaced by a multiple of the smallest eigenvector of Sigma restricted to the dimensions of the contaminated cells. • "both": combines "casewise" and "cellwiseStructured".
perout	The percentage of generated outliers. For outlierType = "casewise" this is a fraction of rows. For outlierType = "cellwisePlain" or outlierType = "cellwiseStructured", a fraction of perout cells are replaced by contaminated cells. For outlierType = "both", a fraction of 0.5*perout of rowwise outliers is generated, after which the remaining data is contaminated with a fraction of 0.5*perout outlying cells.
gamma	How far outliers are from the center of the distribution.
seed	Seed used to generate the data.

Value

A list with components:

- `X`
The generated data matrix of size $n \times d$.
- `indcells`
A vector with the indices of the contaminated cells.
- `indrows`
A vector with the indices of the rowwise outliers.

Author(s)

J. Raymaekers and P.J. Rousseeuw

References

- C. Agostinelli, Leung, A., Yohai, V. J., and Zamar, R. H. (2015). Robust Estimation of Multivariate Location and Scatter in the Presence of Cellwise and Casewise Contamination. *Test*, 24, 441-461.
- Rousseeuw, P.J., Van den Bossche W. (2018). Detecting Deviating Data Cells. *Technometrics*, **60**(2), 135-145. ([link to open access pdf](#))
- J. Raymaekers and P.J. Rousseeuw (2020). Handling cellwise outliers by sparse regression and robust covariance. *Arxiv: 1912.12446*. ([link to open access pdf](#))

See Also

[generateCorMat](#)

Examples

```
n      <- 100
d      <- 5
mu     <- rep(0, d)
Sigma  <- diag(d)
perout <- 0.1
gamma  <- 10
data <- generateData(n, d, mu, Sigma, perout, gamma, outlierType = "cellwisePlain", seed = 1)
pairs(data$X)
data$indcells
```

Description

This function carries out classical PCA when the data may contain missing values, by an iterative algorithm. It is based on a Matlab function from the Missing Data Imputation Toolbox v1.0 by A. Folch-Fortuny, F. Arteaga and A. Ferrer.

Usage

```
ICPCA(X, k, scale = FALSE, maxiter = 20, tol = 0.005,
      tolProb = 0.99, distprob = 0.99)
```

Arguments

X	the input data, which must be a matrix or a data frame. It may contain NA's. It must always be provided.
k	the desired number of principal components
scale	a value indicating whether and how the original variables should be scaled. If scale=FALSE (default) or scale=NULL no scaling is performed (and a vector of 1s is returned in the \$scaleX slot). If scale=TRUE the variables are scaled to have a standard deviation of 1. Alternatively scale can be a function like mad, or a vector of length equal to the number of columns of x. The resulting scale estimates are returned in the \$scaleX slot of the output.
maxiter	maximum number of iterations. Default is 20.
tol	tolerance for iterations. Default is 0.005.
tolProb	tolerance probability for residuals. Defaults to 0.99.
distprob	probability determining the cutoff values for orthogonal and score distances. Default is 0.99.

Value

A list with components:

scaleX	the scales of the columns of X.
k	the number of principal components.
loadings	the columns are the k loading vectors.
eigenvalues	the k eigenvalues.
center	vector with the fitted center.
covmatrix	estimated covariance matrix.
It	number of iteration steps.
diff	convergence criterion.
X.NAimp	data with all NA's imputed.
scores	scores of X.NAimp.
OD	orthogonal distances of the rows of X.NAimp.
cutoffOD	cutoff value for the OD.
SD	score distances of the rows of X.NAimp.
cutoffSD	cutoff value for the SD.
highOD	row numbers of cases whose OD is above cutoffOD.
highSD	row numbers of cases whose SD is above cutoffSD.
residScale	scale of the residuals.
stdResid	standardized residuals. Note that these are NA for all missing values of X.
indcells	indices of cellwise outliers.

Author(s)

Wannes Van Den Bossche

References

Folch-Fortuny, A., Arteaga, F., Ferrer, A. (2016). Missing Data Imputation Toolbox for MATLAB. *Chemometrics and Intelligent Laboratory Systems*, **154**, 93-100.

Examples

```
library(MASS)
set.seed(12345)
n <- 100; d <- 10
A <- diag(d) * 0.1 + 0.9
x <- mvrnorm(n, rep(0,d), A)
x[sample(1:(n * d), 100, FALSE)] <- NA
ICPCA.out <- ICPCA(x, k = 2)
plot(ICPCA.out$scores)
```

MacroPCA

MacroPCA

Description

This function performs the MacroPCA algorithm, which can deal with Missing values and Cellwise and Rowwise Outliers. Note that this function first calls [checkDataSet](#) and analyzes the remaining cleaned data.

Usage

```
MacroPCA(X, k = 0, MacroPCApars = NULL)
```

Arguments

- | | |
|--------------|---|
| X | X is the input data, and must be an n by d matrix or a data frame. It must always be provided. |
| k | k is the desired number of principal components. If $k = 0$ or $k = \text{NULL}$, the algorithm will compute the percentage of explained variability for k upto kmax and show a scree plot, and suggest to choose a value of k such that the cumulative percentage of explained variability is at least 80%. |
| MacroPCApars | <p>A list of available options detailed below. If MacroPCApars = NULL the defaults below are used.</p> <ul style="list-style-type: none"> • DDCpars
A list with parameters for the first step of the MacroPCA algorithm (for the complete list see the function DDC). Default is NULL. |

- **kmax**
The maximal number of principal components to compute. Default is `kmax = 10`. If `k` is provided `kmax` does not need to be specified, unless `k` is larger than 10 in which case you need to set `kmax` high enough.
- **alpha**
This is the coverage, i.e. the fraction of rows the algorithm should give full weight. Alpha should be between 0.50 and 1, the default is 0.50.
- **scale**
A value indicating whether and how the original variables should be scaled. If `scale = FALSE` or `scale = NULL` no scaling is performed (and a vector of 1s is returned in the `$scaleX` slot). If `scale = TRUE` (default) the data are scaled by a 1-step M-estimator of scale with the Tukey biweight weight function to have a robust scale of 1. Alternatively `scale` can be a vector of length equal to the number of columns of `x`. The resulting scale estimates are returned in the `$scaleX` slot of the MacroPCA output.
- **maxdir**
The maximal number of random directions to use for computing the outlyingness of the data points. Default is `maxdir = 250`. If the number n of observations is small all $n * (n - 1) / 2$ pairs of observations are used.
- **distprob**
The quantile determining the cutoff values for orthogonal and score distances. Default is 0.99.
- **silent**
If `TRUE`, statements tracking the algorithm's progress will not be printed. Defaults to `FALSE`.
- **maxiter**
Maximum number of iterations. Default is 20.
- **tol**
Tolerance for iterations. Default is 0.005.
- **center**
if `NULL`, MacroPCA will compute the center. If a vector with d components, this center will be used.
- **bigOutput**
whether to compute and return `NAimp`, `Cellimp` and `Fullimp`. Defaults to `TRUE`.

Value

A list with components:

MacroPCApars	the options used in the call.
remX	Cleaned data after <code>checkDataSet</code> .
DDC	results of the first step of MacroPCA. These are needed to run <code>MacroPCApredict</code> on new data.
scaleX	the scales of the columns of <code>X</code> . When <code>scale = FALSE</code> these are all 1.
k	the number of principal components.

loadings	the columns are the k loading vectors.
eigenvalues	the k eigenvalues.
center	vector with the center.
alpha	alpha from the input.
h	h (computed from alpha).
It	number of iteration steps.
diff	convergence criterion.
X.NAimp	data with all NA's imputed by MacroPCA.
scores	scores of X.NAimp.
OD	orthogonal distances of the rows of X.NAimp.
cutoffOD	cutoff value for the OD.
SD	score distances of the rows of X.NAimp.
cutoffSD	cutoff value for the SD.
highOD	row numbers of cases whose OD is above cutoffOD.
highSD	row numbers of cases whose SD is above cutoffSD.
residScale	scale of the residuals.
stdResid	standardized residuals. Note that these are NA for all missing values of X.
indcells	indices of cellwise outliers.
NAimp	various results for the NA-imputed data.
Cellimp	various results for the cell-imputed data.
Fullimp	various result for the fully imputed data.

Author(s)

Rousseeuw P.J., Van den Bossche W.

References

Hubert, M., Rousseeuw, P.J., Van den Bossche W. (2019). MacroPCA: An all-in-one PCA method allowing for missing values as well as cellwise and rowwise outliers. *Technometrics*, **61**(4), 459-473. ([link to open access pdf](#))

See Also

[checkDataSet](#), [cellMap](#), [DDC](#)

Examples

```
library(MASS)
set.seed(12345)
n <- 50; d <- 10
A <- matrix(0.9, d, d); diag(A) = 1
x <- mvrnorm(n, rep(0,d), A)
x[sample(1:(n * d), 50, FALSE)] <- NA
```

```
x[sample(1:(n * d), 50, FALSE)] <- 10
MacroPCA.out <- MacroPCA(x, 2)
cellMap(MacroPCA.out$stdResid)

# For more examples, we refer to the vignette:
## Not run:
vignette("MacroPCA_examples")

## End(Not run)
```

MacroPCApredict

*MacroPCApredict***Description**

Based on a [MacroPCA](#) fit of an initial (training) data set X , this function analyzes a new (test) data set X_{new} .

Usage

```
MacroPCApredict(Xnew, InitialMacroPCA, MacroPCApars = NULL)
```

Arguments

<code>Xnew</code>	The new data (test data), which must be a matrix or a data frame. It must always be provided. Its columns (variables) should correspond to those of <code>InitialMacroPCA\$remX</code> .
<code>InitialMacroPCA</code>	The output of the <code>MacroPCA</code> function on the initial (training) dataset. Must be provided.
<code>MacroPCApars</code>	The input options to be used for the prediction. By default the options of <code>InitialMacroPCA</code> are used. For the complete list of options see the function MacroPCA .

Value

A list with components:

<code>MacroPCApars</code>	the options used in the call.
<code>DDC</code>	result of <code>DDCpredict</code> which is the first step of <code>MacroPCApredict</code> . See the function DDCpredict .
<code>scaleX</code>	the scales of the columns of X .
<code>k</code>	the number of principal components.
<code>loadings</code>	the columns are the k loading vectors.
<code>eigenvalues</code>	the k eigenvalues.
<code>center</code>	vector with the fitted center.

It	number of iteration steps.
diff	convergence criterion.
Xnew.NAimp	Xnew with all NA's imputed by MacroPCA.
scores	scores of Xnew.NAimp.
OD	orthogonal distances of the rows of Xnew.NAimp.
cutoffOD	cutoff value for the OD.
SD	score distances of the rows of Xnew.NAimp.
cutoffSD	cutoff value for the SD.
highOD	row numbers of cases in Xnew.NAimp whose OD is above cutoffOD.
highSD	row numbers of cases in Xnew.NAimp whose SD is above cutoffSD.
residScale	scale of the residuals.
stdResid	standardized residuals. Note that these are NA for all missing values of Xnew.
indcells	indices of cellwise outliers.
NAimp	various results for the NA-imputed Xnew.
Cellimp	various results for the cell-imputed Xnew.
Fullimp	various result for the fully imputed Xnew.

Author(s)

Rousseeuw P.J., Van den Bossche W.

References

Hubert, M., Rousseeuw, P.J., Van den Bossche W. (2019). MacroPCA: An all-in-one PCA method allowing for missing values as well as cellwise and rowwise outliers. *Technometrics*, **61**(4), 459-473. ([link to open access pdf](#))

See Also

[checkDataSet](#), [cellMap](#), [DDC](#), [DDCpredict](#), [MacroPCA](#)

Examples

```
library(MASS)
set.seed(12345)
n <- 50; d <- 10
A <- matrix(0.9, d, d); diag(A) = 1
x <- mvrnorm(n, rep(0,d), A)
x[sample(1:(n * d), 50, FALSE)] <- NA
x[sample(1:(n * d), 50, FALSE)] <- 10
MacroPCA.out <- MacroPCA(x, 2)
xnew <- mvrnorm(25, rep(0,d), A)
xnew[sample(1:(25 * d), 12, FALSE)] <- 10
predict.out <- MacroPCApredict(xnew, MacroPCA.out)
cellMap(predict.out$stdResid)
```

```
# For more examples, we refer to the vignette:
## Not run:
vignette("MacroPCA_examples")

## End(Not run)
```

outlierMap	<i>Plot the outlier map.</i>
------------	------------------------------

Description

The outlier map is a diagnostic plot for the output of [MacroPCA](#).

Usage

```
outlierMap(res,title="Robust PCA",col="black", pch=16,labelOut=TRUE,id=3,
xlim = NULL, ylim = NULL, cex = 1, cex.main=1.2, cex.lab=NULL, cex.axis=NULL)
```

Arguments

res	A list containing the orthogonal distances (OD), the score distances (SD) and their respective cut-offs (cutoffOD and cutoffSD). Can be the output of MacroPCA , rospca::robpc , rospca::rospca .
title	Title of the plot, default is "Robust PCA".
col	Colour of the points in the plot, this can be a single colour for all points or a vector or list specifying the colour for each point. The default is "black".
pch	Plotting characters or symbol used in the plot, see points for more details. The default is 16 which corresponds to filled circles.
labelOut	Logical indicating if outliers should be labelled on the plot, default is TRUE.
id	Number of OD outliers and number of SD outliers to label on the plot, default is 3.
xlim	Optional argument to set the limits of the x-axis.
ylim	Optional argument to set the limits of the y-axis.
cex	Optional argument determining the size of the plotted points. See plot.default for details.
cex.main	Optional argument determining the size of the main title. See plot.default for details.
cex.lab	Optional argument determining the size of the labels. See plot.default for details.
cex.axis	Optional argument determining the size of the axes. See plot.default for details.

Details

The outlier map contains the score distances on the x-axis and the orthogonal distances on the y-axis. To detect outliers, cut-offs for both distances are shown, see Hubert et al. (2005).

Author(s)

P.J. Rousseeuw

References

Hubert, M., Rousseeuw, P. J., and Vanden Branden, K. (2005). ROBPCA: A New Approach to Robust Principal Component Analysis. *Technometrics*, **47**, 64-79.

See Also[MacroPCA](#)**Examples**

```
# empty for now
```

plot_cellMCD	<i>Draw plots based on the cellwise minimum covariance determinant estimator cellMCD</i>
--------------	--

Description

Function for making plots based on the output of [cellMCD](#).

Usage

```
plot_cellMCD(cellout, type = "Zres/X", whichvar = NULL,
             horizvar = NULL, vertivar = NULL,
             hband = NULL, vband = NULL, drawellipse = T,
             opacity = 0.5, identify = FALSE,
             ids = NULL, labelpoints = T, vl原因 = FALSE,
             clines = TRUE, main = NULL,
             xlab = NULL, ylab = NULL, xlim = NULL,
             ylim = NULL, cex = 1, cex.main = 1.2,
             cex.txt = 0.8, cex.lab = 1, line = 2.0)
```

Arguments

cellout	output of function cellMCD
type	type of diagnostic plot. Should be one of "index", "Zres/X", "Zres/pred", "X/pred", or "bivariate".
whichvar	number or name of the variable to be plotted. Not applicable when type = "bivariate".
horizvar	number or name of the variable to be plotted on the horizontal axis. Only when type = "bivariate".
vertivar	number or name of the variable to be plotted on the vertical axis. Only when type = "bivariate".

hband	whether to draw a horizontal tolerance band. TRUE or FALSE. NULL yields TRUE when type is "index", "Zres/X", or "Zres/pred".
vband	whether to draw a vertical tolerance band. TRUE or FALSE. NULL yields TRUE when type is "Zres/X" or "Zres/pred".
drawellipse	whether to draw a 99% tolerance ellipse. Only for type = "bivariate".
opacity	opacity of the plotted points: 1 is fully opaque, less is more transparent.
identify	if TRUE, identify cases by mouseclick, then Esc.
ids	vector of case numbers to be emphasized (colored red) in the plot. If NULL or of length zero, none are emphasized.
labelpoints	if TRUE, labels the points in ids by their row name in X.
vlines	for the points in ids, draw dashed vertical lines from their standardized residual to 0 when type is "index", "Zres/X", or "Zres/pred". Draws dashed vertical lines to the diagonal when type = "X/pred". Can be TRUE or FALSE, default is FALSE.
clines	only for type == "bivariate". If TRUE, draws a red connecting line from each point in ids to its imputed point, shown in blue.
main	main title of the plot. If NULL, it is constructed automatically from the arguments.
xlab	overriding label for x-axis, unless NULL.
ylab	overriding label for y-axis, unless NULL.
xlim	overriding limits of horizontal axis.
ylim	overriding limits of vertical axis.
cex	size of plotted points.
cex.main	size of the main title.
cex.lab	size of the axis labels.
cex.txt	size of the point labels.
line	distance of axis labels to their axis.

Value

NULL, unless identify = TRUE. Then a list with components:

- ids
the case number(s) that were identified
- coords
coordinates of all points in the plot.

Author(s)

J. Raymaekers and P.J. Rousseeuw

References

J. Raymaekers and P.J. Rousseeuw (2022). The cellwise MCD estimator, Journal of the American Statistical Association, to appear. doi:[10.1080/01621459.2023.2267777](https://doi.org/10.1080/01621459.2023.2267777)(link to open access pdf)

See Also[cellMCD](#)**Examples**

```

mu <- rep(0, 3)
Sigma <- diag(3) * 0.5 + 0.5
set.seed(123)
X <- MASS::mvrnorm(1000, mu, Sigma)
X[1:5, 1] <- X[1:5, 1] + 5
X[6:10, 2] <- X[6:10, 2] - 10
X[12, 1:2] <- c(-4, 8)
cellMCD.out <- cellMCD(X)
plot_cellMCD(cellMCD.out, type="bivariate",
             horizvar=1, vertivar=2, ids=c(1:10,12))

# For more examples, we refer to the vignette:
## Not run:
vignette("cellMCD_examples")

## End(Not run)

```

transfo

*Robustly fit the Box-Cox or Yeo-Johnson transformation***Description**

This function uses reweighted maximum likelihood to robustly fit the Box-Cox or Yeo-Johnson transformation to each variable in a dataset. Note that this function first calls [checkDataSet](#) to ensure that the variables to be transformed are not too discrete.

Usage

```

transfo(X, type = "YJ", robust = TRUE,
        standardize = TRUE,
        quant = 0.99, nbsteps = 2, checkPars = list())

```

Arguments

- | | |
|------|--|
| X | A data matrix of dimensions n x d. Its columns are the variables to be transformed. |
| type | The type of transformation to be fit. Should be one of: <ul style="list-style-type: none"> • "BC": Box-Cox power transformation. Only works for strictly positive variables. If this type is given but a variable is not strictly positive, the function stops with a message about that variable. • "YJ" Yeo-Johnson power transformation. The data may have positive as well as negative values. |

	• "bestObj" for strictly positive variables both BC and YJ are run, and the solution with lowest objective is kept. On the other variables YJ is run.
robust	if TRUE the Reweighted Maximum Likelihood method is used, which first computes a robust initial estimate of the transformation parameter lambda. If FALSE the classical ML method is used.
standardize	whether to standardize the variables before and after the power transformation. See Details below.
quant	quantile for determining the weights in the reweighting step (ignored when robust=FALSE).
nbsteps	number of reweighting steps (ignored when robust=FALSE).
checkPars	Optional list of parameters used in the call to checkDataSet . The options are: • coreOnly If TRUE, skip the execution of checkDataset. Defaults to FALSE • numDiscrete A column that takes on numDiscrete or fewer values will be considered discrete and not retained in the cleaned data. Defaults to 5. • precScale Only consider columns whose scale is larger than precScale. Here scale is measured by the median absolute deviation. Defaults to $1e - 12$. • silent Whether or not the function progress messages should be printed. Defaults to FALSE.

Details

In case standardize = TRUE, the variables is standardized before and after transformation. For BC the variable is divided by its median before transformation. For YJ and robust = TRUE this subtracts its median and divides by its mad (median absolute deviation) before transformation. For YJ and robust = FALSE this subtracts the mean and divides by the standard deviation before transformation. For the standardization after the transformation, the classical mean and standard deviation are used in case robust = FALSE. If robust = TRUE, the mean and standard deviation are calculated robustly on a subset of inliers.

Value

A list with components:

- lambdahats
the estimated transformation parameter for each column of X.
- Y
A matrix in which each column is the transformed version of the corresponding column of X. The transformed version includes pre- and post-standardization if standardize=TRUE.
- muhat
The estimated location of each column of Y.
- sigmahat
The estimated scale of each column of Y.

- **weights**
The final weights from the reweighting.
- **ttypes**
The type of transform used in each column.
- **objective**
Value of the (reweighted) maximum likelihood objective function.
- values of [checkDataSet](#), unless `coreOnly` is TRUE.

Author(s)

J. Raymaekers and P.J. Rousseeuw

References

J. Raymaekers and P.J. Rousseeuw (2021). Transforming variables to central normality. *Machine Learning*. doi:[10.1007/s10994021059605](#)(link to open access pdf)

See Also

[transfo_newdata](#), [transfo_transformback](#)

Examples

```
# find Box-Cox transformation parameter for lognormal data:
set.seed(123)
x <- exp(rnorm(1000))
transfo.out <- transfo(x, type = "BC")
# estimated parameter:
transfo.out$lambdahat
# value of the objective function:
transfo.out$objective
# the transformed variable:
transfo.out$Y
# the type of transformation used:
transfo.out$ttypes
# qqplot of the transformed variable:
qqnorm(transfo.out$Y); abline(0,1)

# For more examples, we refer to the vignette:
## Not run:
vignette("transfo_examples")

## End(Not run)
```

transfo_newdata	<i>Transform variables based on the output of transfo.</i>
-----------------	--

Description

Based on the output of [transfo](#), transform the variables using Yeo-Johnson and/or Box-Cox transformations with the previously estimated parameters and standardization.

Usage

```
transfo_newdata(Xnew, transfo.out)
```

Arguments

Xnew	A data matrix with d columns, which contain the variables to be transformed. The number of columns and their names must be the same as those of the original data on which transfo was run. The number of rows may be different.
transfo.out	The output of a call to transfo .

Value

Returns a matrix with transformed variables.

Author(s)

J. Raymaekers and P.J. Rousseeuw

References

J. Raymaekers and P.J. Rousseeuw (2021). Transforming variables to central normality. *Machine Learning*. doi:[10.1007/s10994021059605](https://doi.org/10.1007/s10994021059605)(link to open access pdf)

See Also

[transfo](#)

Examples

```
set.seed(123); tempraw <- matrix(rnorm(2000), ncol = 2)
tempx <- cbind(tempraw[, 1], exp(tempraw[, 2]))
tempy <- 0.5 * tempraw[, 1] + 0.5 * tempraw[, 2] + 1
x <- tempx[1:900, ]
y <- tempy[1:900]
tx.out <- transfo(x, type = "best0bj")
tx.out$ttypes
tx.out$lbmdahats
tx <- tx.out$Y
lm.out <- lm(y ~ tx)
summary(lm.out)
```

```
xnew <- tempx[901:1000, ]
xtnew <- transfo_newdata(xnew, tx.out)
yhatnew <- tcrossprod(lm.out$coefficients, cbind(1, xtnew))
plot(tempy[901:1000], yhatnew); abline(0, 1)
```

transfo_transformback *Backtransform variables based on the output of [transfo](#).*

Description

Based on the output of [transfo](#), backtransform the variables to their original shape through the inverse Yeo-Johnson and/or Box-Cox transformations with the previously estimated parameters and standardization.

Usage

```
transfo_transformback(Ynew, transfo.out)
```

Arguments

Ynew	A data matrix with d columns, which contain the variables to be backtransformed. The number of columns must be the same as the output Y of the run of transfo on the original data. The number of rows may be different.
transfo.out	The output of a call to transfo .

Value

Returns a matrix with backtransformed variables.

Author(s)

J. Raymaekers and P.J. Rousseeuw

References

J. Raymaekers and P.J. Rousseeuw (2021). Transforming variables to central normality. *Machine Learning*. doi:[10.1007/s10994021059605](https://doi.org/10.1007/s10994021059605)(link to open access pdf)

See Also

[transfo](#)

Examples

```
set.seed(123); x <- matrix(rnorm(2000), ncol = 2)
y <- sqrt(abs(0.3 * x[, 1] + 0.5 * x[, 2] + 4))
ty.out <- transfo(y, type = "BC")
ty.out$lambdahats
ty <- ty.out$Y
lm.out <- lm(ty ~ x)
yhat <- transfo_transformback(lm.out$fitted.values, ty.out)
plot(y, yhat); abline(0, 1)
```

truncPC

Classical Principal Components by truncated SVD.

Description

Similar usage to `robustbase::classPC` except for the new argument `ncomb` which is the desired number of components. Only this many PC's are computed in order to save computation time. Makes use of `propack.svd` of package **svd**.

Usage

```
truncPC(X, ncomp = NULL, scale = FALSE, center = TRUE,
        signflip = TRUE, via.svd = NULL, scores = FALSE)
```

Arguments

<code>X</code>	a numeric matrix.
<code>ncomp</code>	the desired number of components (if not specified, all components are computed).
<code>scale</code>	logical, or numeric vector for scaling the columns.
<code>center</code>	logical or numeric vector for centering the matrix.
<code>signflip</code>	logical indicating if the signs of the loadings should be flipped such that the absolutely largest value is always positive.
<code>via.svd</code>	dummy argument for compatibility with <code>classPC</code> calls, will be ignored.
<code>scores</code>	logical indicating whether or not scores should be returned.

Value

A list with components:

<code>rank</code>	the (numerical) matrix rank of <code>X</code> , i.e. an integer number between 0 and <code>min(dim(x))</code> .
<code>eigenvalues</code>	the <code>k</code> eigenvalues, proportional to the variances, where <code>k</code> is the rank above.
<code>loadings</code>	the loadings, a $d \times k$ matrix.
<code>scores</code>	if the <code>scores</code> argument was <code>TRUE</code> , the $n \times k$ matrix of scores.
<code>center</code>	a vector of means, unless the <code>center</code> argument was <code>FALSE</code> .
<code>scale</code>	a vector of column scales, unless the <code>scale</code> argument was <code>false</code> .

Author(s)

P.J. Rousseeuw

See Also[classPC](#)**Examples**

```
library(MASS)
set.seed(12345)
n <- 100; d <- 10
A <- diag(d) * 0.1 + 0.9
x <- mvrnorm(n, rep(0,d), A)
truncPCA.out <- truncPC(x, ncomp = 2, scores = TRUE)
plot(truncPCA.out$scores)
```

unpack

*Unpacks cellwise weighted data***Description**

This function transforms a dataset X with cellwise weights W to an extended data matrix U with the same number of columns but more rows, and containing more NA's. Its rows have the case weights v .

Usage

```
unpack(X, W)
```

Arguments

X	An n by d data matrix or data frame. Must be given. X is allowed to contain NA's.
W	An n by d matrix of nonnegative cellwise weights. Must be given. W is not allowed to contain NA's.

Value

A list with components:

- U
unpacked data matrix, with the same columns as X but typically more rows.
- v
vector with the rowwise (=casewise) weights of U .

Author(s)

P.J. Rousseeuw

References

P.J. Rousseeuw (2023). Analyzing cellwise weighted data. *Econometrics and Statistics*, appeared online. [doi:10.1016/j.ecosta.2023.01.007](https://doi.org/10.1016/j.ecosta.2023.01.007)(link to open access pdf)

See Also

[weightedEM](#), [cwLocScat](#)

Examples

```
X <- matrix(c(2.8, 5.3, 4.9, 7.4,
              2.3, 5.7, 4.3, 7.2,
              2.5, 5.1, 4.4, 7.6), nrow = 3, byrow = TRUE)
W <- matrix(c(0.8, 1.0, 0.3, 0.4,
              0.3, 0.5, 0.9, 0.5,
              1.0, 0.6, 0, 0.7), nrow = 3, byrow = TRUE)
rownames(X) <- rownames(W) <- c("A", "B", "C")
colnames(X) <- colnames(W) <- c("V1", "V2", "V3", "V4")
X
W
out <- unpack(X, W)
cbind(out$U, out$v)

# For more examples, we refer to the vignette:
## Not run:
vignette("cellwise_weights_examples")

## End(Not run)
```

 weightedEM

Estimates location and scatter on incomplete data with case weights

Description

Carries out a rowwise weighted EM algorithm to estimate mu and Sigma of incomplete Gaussian data.

Usage

```
weightedEM(X, w=NULL, lmin=NULL, crit=1e-4,
            maxiter=1000, initEst=NULL, computeloglik=F)
```

Arguments

<code>X</code>	n by d data matrix or data frame.
<code>w</code>	vector with n nonnegative rowwise (casewise) weights. If NULL, all weights are set to 1 so an unweighted EM is carried out.
<code>lmin</code>	if not NULL, a lower bound on the eigenvalues of the estimated EM covariance matrix on the standardized data, to avoid singularity.
<code>crit</code>	convergence criterion of successive mu and Sigma estimates.
<code>maxiter</code>	maximal number of iteration steps.
<code>initEst</code>	if not NULL, a list with initial estimates μ of the mean, Σ of the covariance matrix.
<code>computeloglik</code>	if TRUE, the log(likelihood) is computed in every step and reported. Default is FALSE to save computation time.

Value

A list with components:

- `mu`
the estimated location vector.
- `Sigma`
the estimated covariance matrix.
- `impX`
the imputed data matrix.
- `niter`
the number of iteration steps taken.
- `loglikhd`
vector with the total log(likelihood) at every iteration step. When `computeloglik = FALSE` this array contains NA's.

Author(s)

P.J. Rousseeuw

References

P.J. Rousseeuw (2023). Analyzing cellwise weighted data. *Econometrics and Statistics*, appeared online. doi:[10.1016/j.ecosta.2023.01.007](https://doi.org/10.1016/j.ecosta.2023.01.007)(link to open access pdf)

See Also

[unpack](#), [cwLocScat](#)

Examples

```

Sigma <- matrix(0.7, 3, 3); diag(Sigma) <- 1
set.seed(12345); X <- MASS::mvrnorm(1000, rep(0, 3), Sigma)
X[1, 3] <- X[2, 2] <- X[3, 1] <- X[4, 1] <- X[5, 2] <- NA
w <- runif(1000, 0, 1) # rowwise weights
out <- weightedEM(X, w, crit = 1e-12, computeloglik = TRUE)
out$niter # number of iteration steps taken
plot(1:out$niter, out$loglikhd[1:out$niter], type = 'l',
     lty = 1, col = 4, xlab = 'step', ylab = 'log(likelihood)',
     main = 'log(likelihood) of weighted EM iterations')
out$mu # estimated center
round(out$Sigma, 6) # estimated covariance matrix
head(X) # the data has NA's
head(out$impX) # imputed data, has no NA's

# For more examples, we refer to the vignette:
## Not run:
vignette("cellwise_weights_examples")

## End(Not run)

```

wrap

Wrap the data.

Description

Transforms multivariate data X using the wrapping function with $b = 1.5$ and $c = 4$. By default, it starts by calling [checkDataSet](#) to clean the data and [estLocScale](#) to estimate the location and scale of the variables in the cleaned data, yielding the vectors $(\hat{\mu}_1, \dots, \hat{\mu}_d)$ and $(\hat{\sigma}_1, \dots, \hat{\sigma}_d)$ where d is the number of variables. Alternatively, the user can specify such vectors in the arguments `locX` and `scaleX`. In either case, the data cell x_{ij} containing variable j of case i is transformed to

$$y_{ij} = \hat{\mu}_j - b_j + \hat{\sigma}_j * \psi((x_{ij} - \hat{\mu}_j)/\hat{\sigma}_j)/a_j$$

in which a_j and b_j are such that for any fixed j the average of y_{ij} equals $\hat{\mu}_j$ and the standard deviation of y_{ij} equals $\hat{\sigma}_j$.

Usage

```

wrap(X, locX = NULL, scaleX = NULL, precScale = 1e-12,
     imputeNA = TRUE, checkPars = list())

```

Arguments

<code>X</code>	the input data. It must be an n by d matrix or a data frame.
<code>locX</code>	The location estimates of the columns of the input data X . Must be a vector of length d .

scaleX	The scale estimates of the columns of the input data X. Must be a vector of length d .
precScale	The precision scale used throughout the algorithm. Defaults to $1e - 12$
imputeNA	Whether or not to impute the NAs with the location estimate of the corresponding variable. Defaults to TRUE.
checkPars	Optional list of parameters used in the call to checkDataSet . The options are: • <code>coreOnly</code> If TRUE, skip the execution of checkDataset. Defaults to FALSE • <code>numDiscrete</code> A column that takes on numDiscrete or fewer values will be considered discrete and not retained in the cleaned data. Defaults to 5. • <code>precScale</code> Only consider columns whose scale is larger than precScale. Here scale is measured by the median absolute deviation. Defaults to $1e - 12$. • <code>silent</code> Whether or not the function progress messages should be printed. Defaults to FALSE.

Value

A list with components:

- `Xw`
The wrapped data.
- `colInWrap`
The column numbers of the variables which were wrapped. Variables which were filtered out by [checkDataSet](#) (because of a (near) zero scale for example), will not appear in this output.
- `loc`
The location estimates for all variables used for wrapping.
- `scale`
The scale estimates for all variables used for wrapping.

Author(s)

Raymaekers, J. and Rousseeuw P.J.

References

Raymaekers, J., Rousseeuw P.J. (2019). Fast robust correlation for high dimensional data. *Technometrics*, **63**(2), 184-198. ([link to open access pdf](#))

See Also

[estLocScale](#)

Examples

```
library(MASS)
set.seed(12345)
n <- 100; d <- 10
X <- mvrnorm(n, rep(0, 10), diag(10))
locScale <- estLocScale(X)
Xw <- wrap(X, locScale$loc, locScale$scale)$Xw
# For more examples, we refer to the vignette:
## Not run:
vignette("wrap_examples")

## End(Not run)
```

Index

cellHandler, [3](#), [27](#)
cellMap, [4](#), [23](#), [25](#), [36](#), [38](#)
cellMCD, [7](#), [40](#), [42](#)
checkDataSet, [7](#), [10](#), [20](#), [23](#), [25–27](#), [34](#), [36](#),
[38](#), [42–44](#), [51](#), [52](#)
classPC, [48](#)
cwLocScat, [11](#), [49](#), [50](#)

data_brands, [13](#)
data_clothes, [14](#)
data_dogWalker, [15](#)
data_dposs, [15](#)
data_glass, [16](#)
data_mortality, [17](#)
data_personality_traits, [17](#)
data_philips, [18](#)
data_VOC, [19](#)
DDC, [3](#), [5](#), [7](#), [10](#), [11](#), [20](#), [24](#), [25](#), [34](#), [36](#), [38](#)
DDCpredict, [5](#), [24](#), [37](#), [38](#)
DI, [3](#), [4](#), [26](#)

estLocScale, [21](#), [28](#), [51](#), [52](#)

generateCorMat, [30](#), [31](#), [32](#)
generateData, [30](#), [31](#)

ICPCA, [32](#)

MacroPCA, [5](#), [10](#), [11](#), [34](#), [37–40](#)
MacroPCApredict, [5](#), [37](#)

outlierMap, [39](#)

plot.default, [39](#)
plot_cellMCD, [9](#), [40](#)

transfo, [11](#), [42](#), [45](#), [46](#)
transfo_newdata, [44](#), [45](#)
transfo_transformback, [44](#), [46](#)
truncPC, [47](#)

unpack, [13](#), [48](#), [50](#)

weightedEM, [13](#), [49](#), [49](#)
wrap, [11](#), [29](#), [51](#)