Package 'MBQN'

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```
Title Mean/Median-balanced quantile normalization
Description Modified quantile normalization for omics or other matrix-like
     data distorted in location and scale.
License GPL-3 + file LICENSE
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Description

An exemplary matrix of a missing value (MV) pattern extracted from LFQ intensities of the protein Groups.txt dataset from PXD001584 [1].

Usage

```
example_NApattern
```

Format

A matrix of zero and ones with 1264 rows and 18 columns; 0 means MV, 1 means no MV.

Author(s)

Ariane Schad

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Source

```
https://www.ebi.ac.uk/pride/archive/projects/PXD001584
```

References

[1] Ramond E, et al. Importance of host cell arginine uptake in Francisella phagosomal escape and ribosomal protein amounts. Mol Cell Proteomics. 2015 14(4):870-881

getKminmax

Get the k largest/smallest elements

Description

Extract the k largest or smallest values and their indices for each column of a matrix.

Usage

```
getKminmax(x, k, flag = "max")
```

Arguments

x a data matrix or data frame.

k an integer specifying the number of extreme values. Must be \leq nrows(x).

flag use "min" or "max" (default) to select smallest or largest elements.

Details

Order the values of each column of x and determine the k smallest (flag = "min") or largest (flag = "max") values and their indices. NA's in the data are ignored.

Value

List with elements:

ik indices of ordered extreme values

minmax ordered extreme values.

Author(s)

Ariane Schad

References

Brombacher, E., Schad, A., Kreutz, C. (2020). Tail-Robust Quantile Normalization. BioRxiv.

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Examples

```
# Create a data matrix x \leftarrow matrix(c(5,2,3,NA,4,1,4,2,3,4,6,NA,1,3,1),ncol=3) # Get indices of the 5 largest values in each column getKminmax(x, k = 5, "max")
```

getPvalue

Calculates Pitman-Morgan variance test on two matrices

Description

Calculates Pitman-Morgan variance test on two matrices

Usage

```
getPvalue(mtx1, mtx2)
```

Arguments

mtx1 Matrix with samples in columns and features in rows mtx2 Matrix with samples in columns and features in rows

Value

Data frame with p values and statistics

Examples

```
set.seed(30)
n <- 20
m <- 20
mtx1 <- matrix(rnorm(m * n), m, n)
mtx2 <- mbqn(mtx1, FUN = "mean")
getPvalue(mtx1, mtx2)</pre>
```

mbqn

Mean/Median-balanced quantile normalization

Description

Modified quantile-normalization (QN) of a matrix, e.g., intensity values from omics data or other data sorted in columns. The modification prevents systematic flattening of features (rows) which are rank invariant (RI) or nearly rank invariant (NRI) across columns, for example features that populate mainly the tails of the intensity distribution or features that separate in intensity.

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Usage

```
mbqn(
   x,
   FUN = "mean",
   na.rm = TRUE,
   method = "limma",
   offsetmatrix = FALSE,
   verbose = FALSE
)
```

Arguments

x a data matrix, where rows represent features, e.g. of protein abundance, and

columns represent groups or samples, e.g. replicates, treatments, or conditions.

FUN a function like mean, median (default), a user defined function, or a numeric

vector of weights with length nrow(x) to balance each feature across samples. Functions can be parsed also as characters. If FUN = NULL, features are not

balanced, i.e. normal QN is used.

na.rm logical indicating to omit NAs in the computation of feature mean.

method character specifying function for computation of quantile normalization; "limma"

(default) for normalizeQuantiles() from the limma package or "preprocess-

Core" for normalize.quantiles() from the preprocessCore package.

offsetmatrix logical indicating if offset matrix should be used instead of offset vector speci-

fying offset for each row

verbose logical indicating to print messages.

Details

Balance each matrix row by substracting its feature offset computed with FUN, e.g. the median; apply quantile-normalization and add the feature means to the normalized matrix. For further details see [4]. For quantile normalization with the "limma" package see [1,2] and for the preProcessCore package see [3].

Value

Normalized matrix

Author(s)

Ariane Schad

References

[1] Smyth, G. K., and Speed, T. P. (2003). Normalization of cDNA microarray data. Methods 31, 265–273.

Ritchie, M.E., Phipson, B., Wu, D., Hu, Y., Law, C.W., Shi, W., and Smyth, [2] G.K. (2015). limma powers differential expression analyses for RNA-sequencing and microarray studies. Nucleic Acids Research 43(7), e47.

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- [3] Bolstad B. M. (2016). preprocessCore: A collection of pre-processing functions. R package version 1.36.0. https://github.com/bmbolstad/preprocessCore
- [4] Brombacher, E., Schad, A., Kreutz, C. (2020). Tail-Robust Quantile Normalization. BioRxiv.

See Also

```
mbqnNRI(), mbqnGetNRIfeatures().
```

Examples

```
## Compute mean and median balanced quantile normalization X \leftarrow \text{matrix}(c(5,2,3,\text{NA},4,1,4,2,3,4,6,\text{NA},1,3,1),\text{ncol}=3) mbqn(X, mean) # Use arithmetic mean to center features mbqn(X, median) # Use median to center features mbqn(X, "median") ## Use user defined array of weights for averaging wt \leftarrow c(1,3,1)/5 # Weights for each sample user_array \leftarrow \text{apply}(X,1,\text{weighted.mean},\text{ wt ,na.rm =TRUE}) mbqn(X, user_array)
```

mbqnBoxplot

Combined box plot and line plot

Description

Create a box-and-whisker plot of a data matrix and plot selected features and/or additional user-defined data on top of it.

Usage

```
mbqnBoxplot(mtx, irow = NULL, vals = NULL, add.leg = TRUE, ...)
```

Arguments

mtx	a matrix or data frame.
irow	index or vector of row indices of matrix features to plot on top of the boxplot.
vals	numeric, array, matrix, or data frame of features with length $ncol(mtx)$ to plot on top of the boxplot.
add.leg	add legend to plot.
• • •	additional arguments passed to the plot functions, e.g. xlab, ylab, main, ylim, type, las.

Details

This function calls graphics::boxplot. Groups are represent by matrix columns. Selected rows/features or user-defined arrays are plot on top of the box plot. Missing values are ignored.

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Value

Figure.

Author(s)

Ariane Schad

References

Brombacher, E., Schad, A., Kreutz, C. (2020). Tail-Robust Quantile Normalization. BioRxiv.

Examples

```
## Create boxplot of quantile normalized data matrix and plot
## feature from median balanced quantile normalization on top of it.
X <- matrix(c(5,2,3,NA,4,1,4,2,3,4,6,NA,1,3,1),ncol=3) # Create data matrix
# Quantile normalization
qn.dat <- mbqn(x=X,FUN = NULL ,na.rm = TRUE)
# Median balanced quantile normalization
mbqn.dat <- mbqn(x=X,FUN = median ,na.rm = TRUE)
## Create boxplot:
plot.new()
mbqnBoxplot(qn.dat,irow = 1, vals = mbqn.dat[1,], type = "b")</pre>
```

mbqnGetIntersect

Helper function for mbqnGetThreshold

Description

Helper function for mbqnGetThreshold

Usage

```
mbqnGetIntersect(combined_qn, combined_mbqn, threshold, plot = TRUE)
```

Arguments

combined_qn Data frame containing RI, p value and statistic calculated for QN combined_mbqn Data frame containing RI, p value and statistic calculated for MBQN threshold Significance threshold for p value of Pitman-Morgan variance test

plot Boolean values if logistic regression curves that are used to calculate intersection

point should be plotted

Value

threshold value

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mbqnGetNRIfeatures

Identify rank invariant (RI) and nearly rank invariant (NRI) features

Description

Compute the rank frequency of each feature of a matrix and identify NRI/RI features.

Usage

```
mbqnGetNRIfeatures(x, low_thr = 0.5, method = NULL, verbose = TRUE)
```

Arguments

x a data matrix. Rows represent features, e.g. protein abundances; columns repre-

sent samples.

low_thr a value between [0 1]. Features with RI frequency >=low_thr are considered as

NRI/RI; default 0.5.

method character specifying function for computation of quantile normalization; "limma"

(default) for normalizeQuantiles() from the limma package or "preprocess-

Core" for normalize.quantiles() from the preprocessCore package.

verbose logical indicating to print messages.

Details

Quantile normalize the data matrix and sort ranks. Determine the maximum frequency of equal rank across all columns for each feature. Features with maximum frequency above the user-defined threhold are declared as nearly rank invariant.

Value

A list with elements:

p a matrix with the rank invariance frequencies ri. freq and the sample coverage

sample.coverage for all detected RI/NRI features

max_p maximum rank invariance frequency in percent

ip index of feature with maximum rank invariance frequency

nri table of the rank invariance frequencies in percent for each NRI/RI feature

var0_feature indices of features with zero sample variance after QN low_thr threshold used for NRI/RI detection from RI frequency.

Author(s)

Ariane Schad

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References

Brombacher, E., Schad, A., Kreutz, C. (2020). Tail-Robust Quantile Normalization. BioRxiv.

See Also

mbqnPlotRI() for visualization of detected NRI/RI features.

Examples

```
## Check data matrix for RI and NRI features
set.seed(1234)
x <- mbqnSimuData("omics.dep")
RI <- mbqnGetNRIfeatures(x, low_thr = 0.5, verbose = FALSE)
mbqnPlotRI(RI)</pre>
```

mbqnGetThreshold

Calculates the rank invariance threshold from which on MBQN should be used instead of 'classical' QN

Description

Calculates the rank invariance threshold from which on MBQN should be used instead of 'classical' ON

Usage

```
mbqnGetThreshold(mtx, meanMedian = "mean", plot = TRUE)
```

Arguments

mtx Matrix with samples in columns and features in rows

meanMedian Offset function for the MBQN calculation

plot Boolean values if logistic regression curves that are used to calculate intersection

point should be plotted

Value

threshold value

```
set.seed(30)
n <- 20
m <- 20
mtx <- matrix(rnorm(m * n), m, n)
mbqnGetThreshold(mtx)</pre>
```

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mbqnNRI

Selective mean/median-balanced quantile normalization

Description

Quantile normalization of a data matrix where rank invariant (RI)/nearly rank invariant (NRI) rows/features or other user-selected rows are normalized by the mean/median-balanced quantile normalization.

Usage

```
mbqnNRI(
    x,
    FUN = "mean",
    na.rm = TRUE,
    method = NULL,
    low_thr = 0.5,
    index = NULL,
    offsetmatrix = FALSE,
    verbose = TRUE
)
```

Arguments

x	a data matrix, where rows represent features, e.g. of protein abundance, and columns represent groups or samples, e.g. replicates, treatments, or conditions.
FUN	a function like mean, median (default), a user defined function, or a numeric vector of weights with length $nrow(x)$ to balance each feature across samples. Functions can be parsed also as characters. If $FUN = NULL$, features are not balanced, i.e. normal QN is used.
na.rm	logical indicating to omit NAs in the computation of feature mean.
method	character specifying function for computation of quantile normalization; "limma" (default) for normalizeQuantiles() from the limma package or "preprocess-Core" for normalize.quantiles() from the preprocessCore package.
low_thr	a value between [0 1]. Features with RI frequency >=low_thr are considered as NRI/RI; default 0.5.
index	an integer or a vector integers specifying the indices of selected rows.
offsetmatrix	logical indicating if offset matrix should be used instead of offset vector specifying offset for each row
verbose	logical indicating to print messages.

Details

Selected rows and/or rows with rank invariance frequency >=threshold are normalized with the mean/median-balanced quantile normalization. Remaining rows are quantile normalized without mean balancing.

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Value

Normalized matrix.

Author(s)

Ariane Schad

References

Brombacher, E., Schad, A., Kreutz, C. (2020). Tail-Robust Quantile Normalization. BioRxiv.

See Also

```
mbqn(), mbqnGetNRIfeatures().
```

Examples

```
## Quantile normalize a data matrix where ## nearly rank invariant (NRI) features are balanced X <- matrix(c(5,2,3,NA,4,1,4,2,3,4,6,NA,1,3,1),ncol=3) mbqnNRI(X, median,low_thr = 0.5) # Balance NRI features selected by threshold mbqnNRI(X, median, index = c(1,2)) # Balance selected features
```

mbqnPlotAll

Plot RI/NRI feature frequencies and normalized/unnormalized features

Description

Check data matrix for rank invariant (RI) and nearly rank invariant (NRI) features/rows across samples and visualize result for different normalizations.

Usage

```
mbqnPlotAll(
    x,
    FUN = NULL,
    low_thr = 0.5,
    show_nri_only = FALSE,
    verbose = TRUE,
    ...
)
```

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Arguments

x a data matrix. Rows represent features, e.g. protein abundances; columns repre-

sent samples.

FUN a function like mean, median (default), a user defined function, or a numeric

vector of weights with length nrow(x) to balance each feature across samples. Functions can be parsed also as characters. If FUN = NULL, features are not

balanced, i.e. normal QN is used.

low_thr a value between [0 1]. Features with RI frequency >=low_thr are considered as

NRI/RI; default 0.5.

show_nri_only logical indicating to display only the RI/NRI detection graph.

verbose logical indicating to print messages.

... additional plot arguments passed to mbqnBoxplot, and mbqnPlotRI.

Details

Rank data and check if lower and upper intensity tails are dominated by few features. Apply quantile normalization without and with mean-balancing and check the standard deviation of normalized features located in the tails.

Value

A set of figures that display the detected RI/NRI features and a list with elements:

p a matrix with the rank invariance frequencies ri.freq and the sample coverage

sample.coverage for all detected RI/NRI features

max_p maximum rank invariance frequency in percent

ip index of feature with maximum rank invariance frequency

nri table of the rank invariance frequencies in percent for each NRI/RI feature

var0_feature indices of features with zero sample variance after QN.

Author(s)

Ariane Schad

References

Brombacher, E., Schad, A., Kreutz, C. (2020). Tail-Robust Quantile Normalization. BioRxiv.

See Also

mbqnPlotRI() and mbqnBoxplot() for the generation of figures, and mbqn() for normalization.

```
## Check data matrix for RI and NRI features
X <- matrix(c(5,2,3,NA,4,1,4,2,3,4,6,NA,1,3,1),ncol=3)
mbqnPlotAll(X, mean, low_thr = 0.5)</pre>
```

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mbqnPlotRI	Plot frequency of detected RI/NRI features	

Description

Plot rank invariance frequency and feature coverage of detected RI and NRI features

Usage

```
mbqnPlotRI(obj, verbose = FALSE, ...)
```

Arguments

obj list object of RI frequencies from mbqnGetNRIfeatures().

verbose logical indicating to run function quietly.

... additional arguments (cex, cex.lab, cex.axis, cex.main) passed to the plot func-

tion.

Details

Graphical output of the NRI/RI identification results from mbqnGetNRIfeatures(). For each detected NRI/RI feature, plot the feature index against the RI frequencies together with the RI frequency detection threshold and print the sample coverage.

Value

Figure

Author(s)

Ariane Schad

References

Brombacher, E., Schad, A., Kreutz, C. (2020). Tail-Robust Quantile Normalization. BioRxiv.

See Also

mbqnGetNRIfeatures() for detection of NRI/RI features.

```
## Check data matrix for RI and NRI features
x <- mbqnSimuData("omics.dep")
RI <- mbqnGetNRIfeatures(x, low_thr = 0.5, verbose = FALSE)
mbqnPlotRI(RI)</pre>
```

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mbqnSimuData

Generate a random/structured data matrix

Description

Generate a random data matrix with or without proteomics, log-transformed feature intensity-like properties.

Usage

```
mbqnSimuData(model = "rand", nrow = NULL, ncol = NULL, show.fig = FALSE)
```

Arguments

model character indicating one of the three different type of models: "rand"(default)

a Gaussian random matrix of size nrow x ncol (default 1000 x 10), "omics" a Gaussian random matrix of size 1264 x 18 that mimics intensity profiles and missing values as present in real data, and "omics.dep" is the same as "omics"

but with an additional single, differentially expressed RI feature.

nrow number of rows of data matrix (only for model = "rand").

ncol number of columns of data matrix (only for model = "rand").

show.fig logical inidicating whether data properties are plot to figure (only for model =

"omics" and model = "omics.dep").

Details

For model "rand", each matrix element is drawn from a standard normal distribution N(0,1). For model "omics", the matrix elements of each row are drawn from a Gaussian distribution $N(\mu_i, \sigma_i^2)$ where the mean and standard deviation itself are drawn Gaussian distributions, i.e. $\sigma_i \ N(0, 0.0625)$ and $\mu_i \ N(28,4)$. About 35\ to the missing value pattern present in real protein LFQ intensities. For model "omics.dep", a single differentially epxressed RI feature is stacked on top of the matrix from model "omics".

Value

matrix of size nrow x ncol.

Author(s)

Ariane Schad

References

Brombacher, E., Schad, A., Kreutz, C. (2020). Tail-Robust Quantile Normalization. BioRxiv.

See Also

example_NApattern() for description of missing value pattern.

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Examples

```
mbqnSimuData(model = "rand")
mbqnSimuData(model = "rand", 2000,6)
set.seed(1234)
mbqnSimuData(model = "omics")
set.seed(1111)
mbqnSimuData(model = "omics.dep")
```

mbqnSimuDistortion

Perturbation of sample mean and scale

Description

mbqnSimuDistortion adds a random perturbation of mean and scale to each column of a matrix.

Usage

```
mbqnSimuDistortion(x, s.mean = 0.05, s.scale = 0.01)
```

Arguments

x a matrix or data frame.

s.mean scatter of relative change of mean.

s. scale scatter of realtive change in scale, i.e. 0.01 corresponds to 1 percent.

Details

Shift and scale the sample mean and standard deviation of a matrix. The perturbation of center and scale relative to mean and standard deviation of each sample are drawn from a Gaussian distribution $|N(0,\sigma^2)|$ with $\sigma_m ean = s$. mean and $\sigma_s cale = s$. scale, respectively.

Value

List with:

x.mod perturbed matrix

mx.offset numeric array of shifts of the sample means

mx.scale numeric array of relative scales of the sample standard deviations.

Author(s)

Ariane Schad

References

Brombacher, E., Schad, A., Kreutz, C. (2020). Tail-Robust Quantile Normalization. BioRxiv.

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See Also

```
mbqnSimuData() for data generation.
```

Examples

```
set.seed(1234)
x <- mbqnSimuData("omics.dep")
df <- mbqnSimuDistortion(x)</pre>
```

one Sided Test

Recalulate p value from two-sided to one-sided

Description

Recalulate p value from two-sided to one-sided

Usage

```
oneSidedTest(sign_value, z_value)
```

Arguments

sign_value P value from two-sided significance test z_value Z value from two-sided significance test

Value

P value from one sided significance test

truncateDecimals

Truncate float to defined number of decimal values

Description

Truncate float to defined number of decimal values

Usage

```
truncateDecimals(x, digits = 2)
```

Arguments

x float

digits Number of decimal values

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Value

Truncated number

```
x <- 2.567836
truncateDecimals(x, 3)</pre>
```

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