Package 'MatrixQCvis'

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Type Package

Title Shiny-based interactive data-quality exploration for omics data

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VignetteBuilder knitr

Description Data quality assessment is an integral part of preparatory data analysis to ensure sound biological information retrieval.

We present here the MatrixQCvis package, which provides shiny-based interactive visualization of data quality metrics at the per-sample and per-feature level. It is broadly applicable to quantitative omics data types that come in matrix-like format (features x samples). It enables the detection of low-quality samples, drifts, outliers and batch effects in data sets. Visualizations include amongst others bar- and violin plots of the (count/intensity) values, mean vs standard deviation plots, MA plots, empirical cumulative distribution function (ECDF) plots, visualizations of the distances between samples, and multiple

types of dimension reduction plots. Furthermore, MatrixQCvis allows for differential expression analysis based on the limma (moderated t-tests) and proDA (Wald tests) packages. MatrixQCvis builds upon the popular Bioconductor SummarizedExperiment S4 class and enables thus the facile integration into existing workflows. The package is especially tailored towards metabolomics and proteomics mass spectrometry

data, but also allows to assess the data quality of other data types that can be represented in a SummarizedExperiment object.

Depends SummarizedExperiment (>= 1.20.0), plotly (>= 4.9.3), shiny (>= 1.6.0)

Imports ComplexHeatmap (>= 2.7.9), dplyr (>= 1.0.5), ExperimentHub (>= 2.6.0), ggplot2 (>= 3.3.3), grDevices (>= 4.1.0), Hmisc (>= 4.5-0), htmlwidgets (>= 1.5.3), impute (>= 1.65.0), imputeLCMD (>= 2.0), limma (>= 3.47.12), MASS (>= 7.3-58.1), methods (>= 4.1.0), pcaMethods (>= 1.83.0), proDA (>= 1.5.0), rlang (>= 0.4.10), rmarkdown (>= 2.7), Rtsne (>= 0.15), shinydashboard (>= 0.7.1), shinyhelper (>= 0.3.2), shinyjs (>= 2.0.0), stats (>= 4.1.0), tibble (>= 3.1.1), tidyr (>= 1.1.3), umap (>= 0.2.7.0), UpSetR (>= 1.4.0), vsn (>= 3.59.1)

Suggests BiocGenerics (>= 0.37.4), BiocStyle (>= 2.19.2), hexbin (>= 1.28.2), jpeg (>= 0.1-10), knitr (>= 1.33), statmod (>= 1.5.0), testthat (>= 3.0.2)

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barplotSamplesMeasuredMissing

Barplot of number of measured/missing features of samples

Description

barplotSamplesMeasuredMissing plots the number of measured/missing features of samples as a barplot. The function will take as input the returned tbl of samplesMeasuredMissing.

Usage

```
barplotSamplesMeasuredMissing(tbl, measured = TRUE)
```

Arguments

| tbl | tbl object |
|----------|---|
| measured | logical, should the number of measured or missing values be plotted |

Value

gg object from ggplot2

Examples

```
se <- SummarizedExperiment::SummarizedExperiment(assay = a,
    rowData = rD, colData = cD)
## create the data.frame with information on number of measured/missing
## values
tbl <- samplesMeasuredMissing(se)
## plot number of measured values
barplotSamplesMeasuredMissing(tbl, measured = TRUE)
## plot number of missing values
barplotSamplesMeasuredMissing(tbl, measured = FALSE)
```

| batchCorrectionAssay | Remove | batch | effects | from | (count/intensity) | values | of | а |
|----------------------|---------|---------|---------|------|-------------------|--------|----|---|
| | Summari | zedExpe | riment | | | | | |

Description

The function batchCorrectionAssay removes the batch effect of (count/intensity) values of a SummarizedExperiment. It uses either the removeBatchEffect function or no batch effect correction method (pass-through, none).

Usage

```
batchCorrectionAssay(
    se,
    method = c("none", "removeBatchEffect (limma)"),
    batchColumn = colnames(se@colData)
)
```

Arguments

| se | SummarizedExperiment |
|-------------|--|
| method | character, one of "none" or "removeBatchEffect" |
| batchColumn | <pre>character, one of colnames(colData(se))</pre> |

Details

The column batchColumn in colData(se) contains the information on the batch identity. Internal use in shinyQC.

If batchColumn is NULL, batchColumn is internally set to the name of the first column in colData(se) if method = "removeBatchEffect (limma)".

Value

matrix

createBoxplot

Examples

createBoxplot

Create a boxplot of (count/intensity) values per sample

Description

The function create_boxplot creates a boxplot per sample for the intensity/count values.

Usage

```
createBoxplot(
   se,
   orderCategory = colnames(colData(se)),
   title = "",
   log = TRUE,
   violin = FALSE
)
```

Arguments

| se | eq:summarizedExperiment containing the (count/intensity) values in the assay slot |
|---------------|---|
| orderCategory | character, one of colnames(colData(se)) |
| title | character or numeric of length(1) |
| log | logical, if TRUE (count/intensity) values are displayed as log values |
| violin | logical, if FALSE a boxplot is created, if TRUE a violin plot is created |

Details

Internal usage in shinyQC.

Value

gg object from ggplot2

Examples

```
## create se
a <- matrix(1:100, nrow = 10, ncol = 10,
    dimnames = list(1:10, paste("sample", 1:10)))
a[c(1, 5, 8), 1:5] <- NA
set.seed(1)
a <- a + rnorm(100)
cD <- data.frame(name = colnames(a), type = c(rep("1", 5), rep("2", 5)))
rD <- data.frame(spectra = rownames(a))
se <- SummarizedExperiment::SummarizedExperiment(assay = a,
    rowData = rD, colData = cD)
createBoxplot(se, orderCategory = "name", title = "", log = TRUE,
    violin = FALSE)
```

createDfFeature Create data frame of (count/intensity) values for a selected feature along data processing steps

Description

The function createDfFeature takes as input a list of matrices and returns the row Feature of each matrix as a column of a data.frame. The function createDfFeature provides the input for the function featurePlot.

Usage

createDfFeature(1, feature)

Arguments

| 1 | list containing matrices at different processing steps |
|---------|--|
| feature | character, element of rownames of the matrices in 1 |

Details

Internal usage in shinyQC

Value

data.frame

Examples

```
set.seed(1)
x1 <- matrix(rnorm(100), ncol = 10, nrow = 10,
    dimnames = list(paste("feature", 1:10), paste("sample", 1:10)))
x2 <- x1 + 5
x3 <- x2 + 10
l <- list(x1 = x1, x2 = x2, x3 = x3)
createDfFeature(1, "feature 1")</pre>
```

C٧

Calculate coefficient of variation

Description

The function cv calculates the coefficient of variation from columns of a matrix. The coefficients of variation are calculated according to the formula sd(y) / mean(y) * 100 with y the column values, thus, the function returns the coefficient of variation in percentage.

Usage

cv(x, name = "raw")

Arguments

| Х | matrix |
|------|--|
| name | character, the name of the returned list |

Details

The function returned a named list (the name is specified by the name argument) containing the coefficient of variation of the columns of x.

Value

list

Examples

x <- matrix(1:10, ncol = 2)
cv(x)</pre>

cvFeaturePlot

Description

The function cvFeaturePlot returns a plotly plot of coefficient of variation values. It will create a violin plot and superseded points of coefficient of variation values per list entry of 1.

Usage

cvFeaturePlot(1, lines = FALSE)

Arguments

| 1 | list containing matrices |
|-------|--------------------------|
| lines | logical |

Details

lines = TRUE will connect the points belonging to the same feature with a line. If there are less than two features, the violin plot will not be plotted. The violin plots will be ordered according to the order in 1

Value

plotly

Examples

```
x1 <- matrix(1:100, ncol = 10, nrow = 10,
    dimnames = list(paste("feature", 1:10), paste("sample", 1:10)))
x2 <- x1 + 5
x3 <- x2 + 10
1 <- list(x1 = x1, x2 = x2, x3 = x3)
cvFeaturePlot(1, lines = FALSE)
```

dimensionReduction

Dimensionality reduction with dimensionReduction methods PCA, PCoA, NMDS, UMAP and tSNE

dimensionReduction

Description

The function dimensionReduction creates a data.frame with the coordinates of the projected data (first entry of returned output). The function allows for the following projections: Principal Component Analysis (PCA), Principal Coordinates Analysis/Multidimensional Scaling (PCoA), Nonmetric Multidimensional scaling (NMDS), t-distributed stochastic neighbor embedding (tSNE), and Uniform Manifold Approximation and Projection (UMAP).

The second list entry will contains the object returned from prcomp (PCA), cmdscale (PCoA), isoMDS (NMDS), Rtsne (tSNE), or umap (UMAP).

Usage

```
dimensionReduction(
```

```
x,
type = c("PCA", "PCoA", "NMDS", "tSNE", "UMAP"),
params = list()
)
```

Arguments

| х | matrix, containing no missing values, samples are in columns and features are in rows |
|--------|--|
| type | character, specifying the type/method to use for dimensionality reduction. One of PCA, PCoA, NMDS, tSNE, or UMAP. |
| params | list, arguments/parameters given to the functions stats::prcomp, stats::dist, Rtsne::Rtsne, umap::umap |

Details

The function dimensionReduction is a wrapper around the following functions stats::prcomp (PCA), stats::cmdscale (PCoA), MASS::isoMDS (NMDS), Rtsne::Rtsne (tSNE), and umap::umap (UMAP). For the function umap::umap the method is set to naive.

Value

list, first entry contains a tbl, second entry contains the object returned from prcomp (PCA), cmdscale (PCoA), isoMDS (NMDS), Rtsne (tSNE), or umap (UMAP)

Author(s)

Thomas Naake

```
x <- matrix(rnorm(1:10000), ncol = 100)
rownames(x) <- paste("feature", 1:nrow(x))
colnames(x) <- paste("sample", 1:ncol(x))
params <- list(method = "euclidean", ## dist
    initial_dims = 10, max_iter = 100, dims = 3, perplexity = 3, ## tSNE
    min_dist = 0.1, n_neighbors = 15, spread = 1) ## UMAP</pre>
```

```
dimensionReduction(x, type = "PCA", params = params)
dimensionReduction(x, type = "PCoA", params = params)
dimensionReduction(x, type = "NMDS", params = params)
dimensionReduction(x, type = "tSNE", params = params)
dimensionReduction(x, type = "UMAP", params = params)
```

dimensionReductionPlot

Plot the coordinates from dimensionReduction values

Description

The function dimensionReductionPlot creates a dimension reduction plot. The function takes as input the tbl object obtained from the dimensionReduction function. The tbl contains transformed values by one of the dimension reduction methods.

Usage

```
dimensionReductionPlot(
  tbl,
  se,
  highlight = c("none", colnames(se@colData)),
  explainedVar = NULL,
  x_coord,
  y_coord,
  height = 600,
  interactive = TRUE
)
```

Arguments

| tbl | tbl as obtained by the function dimensionReduction |
|--------------|--|
| se | SummarizedExperiment |
| highlight | character, one of "none" or colnames(se@colData) |
| explainedVar | NULL or named numeric, if numeric explainedVar contains the explained variance per principal component (names of explainedVar corresponds to the principal components) |
| x_coord | character, column name of tbl that stores x coordinates |
| y_coord | character, column name of tbl that stores y coordinates |
| height | numeric, specifying the height of the plot (in pixels) |
| interactive | logical(1), if TRUE dimensionReductionPlot will return a plotly object, if FALSE dimensionReductionPlot will return a gg object |

distSample

Details

The function dimensionReductionPlot is a wrapper for a ggplot/ggplotly expression.

Value

plotly or gg

Author(s)

Thomas Naake

Examples

library(SummarizedExperiment)

distSample

Create a heatmap using distance information between samples

Description

The function distSample creates a heatmap from a distance matrix created by the function distShiny. The heatmap is annotated by the column specified by the label column in colData(se).

Usage

distSample(d, se, label = "name", title = "raw", ...)

Arguments

| d | matrix containing distances, obtained from distShiny |
|-------|--|
| se | SummarizedExperiment |
| label | character, refers to a column in colData(se) |
| title | character |
| | further arguments passed to ComplexHeatmap::Heatmap |

Details

Internal use in shinyQC

Value

Heatmap object from ComplexHeatmap

Examples

distShiny

Create distance matrix from numerical matrix

Description

The function distShiny takes as an input a numerical matrix or data.frame and returns the distances between the rows and columns based on the defined method (e.g. euclidean distance).

Usage

```
distShiny(x, method = "euclidean")
```

Arguments

| Х | matrix or data.frame with samples in columns and features in rows |
|--------|---|
| method | character, method for distance calculation |

Details

Internal use in shinyQC.

driftPlot

Value

matrix

Examples

```
driftPlot
```

Plot the trend line for aggregated values

Description

The function driftPlot aggregates the (count/intensity) values from the assay() slot of a SummarizedExperiment by the median or sum of the (count/intensity) values. driftPlot then visualizes these aggregated values and adds a trend line (using either LOESS or a linear model) from (a subset of) the aggregated values. The subset is specified by the arguments category and level.

Usage

```
driftPlot(
    se,
    aggregation = c("median", "sum"),
    category = colnames(colData(se)),
    orderCategory = colnames(colData(se)),
    level = c("all", unique(colData(se)[, category])),
    method = c("loess", "lm")
)
```

Arguments

| se | SummarizedExperiment |
|---------------|---|
| aggregation | character, type of aggregation of (count/intensity) values |
| category | character, column of colData(se) |
| orderCategory | character, column of colData(se) |
| level | character, from which samples should the LOESS curve be calculated, either "all" or one of the levels of the selected columns of colData(se) ("category") |
| method | character, either "loess" or "lm" |

Details

The x-values are sorted according to the orderCategory argument: The levels of the corresponding column in colData(se) are pasted with the sample names (in the column name) and factorized. Internal usage in shinyQC.

Value

gg object from ggplot2

Examples

ECDF

Create ECDF plot of a sample against a reference

Description

The function ECDF creates a plot of the empirical cumulative distribution function of a specified sample and an outgroup (reference). The reference is specified by the group argument. The row-wise (feature) mean values of the reference are calculated after excluding the specified sample.

Usage

ECDF(se, sample = colnames(se), group = c("all", colnames(colData(se))))

Arguments

| se | SummarizedExperiment object |
|--------|--|
| sample | character, name of the sample to compare against the group |
| group | <pre>character, either "all" or one of colnames(colData(se))</pre> |

Details

Internal use in shinyQC.

The function ECDF uses the ks.test function from stats to perform a two-sample Kolmogorov-Smirnov test. The Kolmogorov-Smirnov test is run with the alternative "two.sided" (null hypothesis is that the true distribution function of the sample is equal to the hypothesized distribution function of the group).

The exact argument in ks.test is set to NULL, meaning that an exact p-value is computed if the product of the sample sizes is less than 10000 of sample and group. Otherwise, asymptotic distributions are used whose approximations might be inaccurate in low sample sizes.

explVar

Value

gg object from ggplot2

Examples

Retrieve the explained variance for each principal component (PCA) or axis (PCoA)

Description

The function explVar calculates the proportion of explained variance for each principal component (PC, type = "PCA") and axis (type = "PCA").

Usage

explVar(d, type = c("PCA", "PCoA"))

Arguments

| d | prcomp or list from cmdscale |
|------|-----------------------------------|
| type | character, one of "PCA" or "PCoA" |

Details

explVar uses the function prcomp from the stats package to retrieve the explained standard deviation per PC (type = "PCA") and the function cmdscale from the stats package to retrieve the explained variation based on eigenvalues per Axis (type = "PCoA").

Value

numeric vector with the proportion of explained variance for each PC or Axis

Author(s)

Thomas Naake

Examples

```
x <- matrix(1:100, nrow = 10, ncol = 10,
    dimnames = list(1:10, paste("sample", 1:10)))
set.seed(1)
x <- x + rnorm(100)
## run for PCA
pca <- dimensionReduction(x = x,
    params = list(center = TRUE, scale = TRUE), type = "PCA")[[2]]
explVar(d = pca, type = "PCA")
## run for PCoA
pcoa <- dimensionReduction(x = x,
    params = list(method = "euclidean"), type = "PCA")[[2]]
explVar(d = pcoa, type = "PCA")
```

```
extractComb
```

Obtain the features that are present in a specified set

Description

The function extractComb extracts the features that match a combination depending if the features was measured or missing. The function will return the sets that match the combination, thus, the function might be useful when answering questions about which features are measured/missing under certain combinations (e.g. sample types or experimental conditions).

Usage

```
extractComb(se, combination, measured = TRUE, category = "type")
```

Arguments

| se | SummarizedExperiment |
|-------------|--|
| combination | character, refers to factors in category |
| measured | logical |
| category | character, corresponding to a column name in colData(se) |

Details

The function extractComb uses the make_comb_mat function from ComplexHeatmap package. Presence is defined by a feature being measured in at least one sample of a set. Absence is defined by a feature with only missing values (i.e. no measured values) of a set.

Value

character

featurePlot

Examples

featurePlot

Create a plot of (count/intensity) values over the samples

Description

The function featurePlot creates a plot of (count/intensity) values for different data processing steps (referring to columns in the data.frame) over the different samples (referring to rows in the data.frame).

Usage

featurePlot(df)

Arguments

df data.frame

Details

Internal usage in shinyQC.

Value

gg object from ggplot2

```
set.seed(1)
x1 <- matrix(rnorm(100), ncol = 10, nrow = 10,
    dimnames = list(paste("feature", 1:10), paste("sample", 1:10)))
x2 <- x1 + 5
x3 <- x2 + 10
l <- list(x1 = x1, x2 = x2, x3 = x3)
df <- createDfFeature(1, "feature 1")
featurePlot(df)</pre>
```

histFeature

Description

The function $\verb+histFeature$ creates a histogram with the number of measured/missing values per feature.

Usage

```
histFeature(x, measured = TRUE, ...)
```

Arguments

| Х | matrix containing intensities. Missing values are encoded as NA. |
|----------|---|
| measured | logical, should the measured values (measured = TRUE) or missing values (measured = FALSE) be taken |
| | additional parameters passed to geom_histogram, e.g. binwidth. |

Value

plotly object from ggplotly

Examples

histFeatureCategory Histogram of features per sample type

Description

The function histFeatureCategory creates histogram plots for each sample type in se.

Usage

```
histFeatureCategory(se, measured = TRUE, category = "type", ...)
```

hist_sample

Arguments

| se | SummarizedExperiment, the assay slot contains the intensity values per sample. Missing values are encoded as NA. |
|----------|---|
| measured | logical, should the measured values (measured = TRUE) or missing values (measured = FALSE) be taken |
| category | character, corresponding to a column in colData(se) |
| | additional parameters passed to geom_histogram, e.g. binwidth. |

Value

plotly object from ggplotly

Examples

histFeatureCategory(se, measured = TRUE, category = "type")

hist_sample

Plot a histogram of the number of a category

Description

hist_sample plots the number of a category (e.g. sample types) as a histogram. It use the returned tbl from hist_sample_num.

Usage

hist_sample(tbl, category = "type")

Arguments

| tbl | tbl as returned by hist_sample_num |
|----------|-------------------------------------|
| category | character, x-axis label of the plot |

Value

gg object from ggplot2

Examples

hist_sample_num *Return the number of a category*

Description

hist_sample_num returns the number of a category (e.g. sample types) as a tbl. The function will retrieve first the column category in colData(se). The function will return a tbl containing the numerical values of the quantities.

Usage

```
hist_sample_num(se, category = "type")
```

Arguments

| se | SummarizedExperiment object |
|----------|---|
| category | character, corresponding to a column in colData(se) |

Value

tbl

Examples

hoeffDPlot

```
rowData = rD, colData = cD)
hist_sample_num(se, category = "type")
```

hoeffDPlot

Create a plot from a list of Hoeffding's D values

Description

The function hoeffDPlot creates via ggplot a violin plot per factor, a jitter plot of the data points and (optionally) connects the points via lines. hoeffDPlot uses the plotly package to make the figure interactive.

Usage

hoeffDPlot(df, lines = TRUE)

Arguments

| df | data.frame containing one or multiple columns containing the Hoeffding's D statistics |
|-------|---|
| lines | logical, should points belonging to the same sample be connected |
| 22000 | 1081041, should points coronging to the sample of connected |

Details

The function hoeffDPlot will create the violin plot and jitter plot according to the specified order given by the colnames of df. hoeffDPlot will thus internally refactor the colnames of the supplied data.frame according to the order of the colnames.

Value

gg object from ggplot2

```
## normalized values
se_n <- se
assay(se_n) <- normalizeAssay(a, "sum")
tbl_n <- MAvalues(se_n, log = FALSE, group = "all")
hd_n <- hoeffDValues(tbl_n, "normalized")

df <- data.frame(raw = hd_r, normalized = hd_n)
hoeffDPlot(df, lines = TRUE)
hoeffDPlot(df, lines = FALSE)</pre>
```

hoeffDValues

Create values of Hoeffding's D statistics from M and A values

Description

The function creates and returns Hoeffding's D statistics values from MA values.

In case sample_n is set to a numerical value (e.g. 10000), a random subset containing sample_n is taken to calculate Hoeffding's D values to speed up the calculation. In case there are less features than sample_n, all features are taken.

Usage

```
hoeffDValues(tbl, name = "raw", sample_n = NULL)
```

Arguments

| tbl | tibble, as obtained from the function MAvalues |
|----------|--|
| name | character(1), name of the returned list |
| sample_n | numeric(1), number of features (subset) to be taken for calculation of Hoeffd- ing's D values |

Details

The function uses the function hoeffd from the Hmisc package to calculate the values.

Value

named list with Hoeffding's D values per sample

Examples

imputeAssay

```
cD <- data.frame(name = colnames(a), type = c(rep("1", 5), rep("2", 5)))
rD <- data.frame(spectra = rownames(a))</pre>
se <- SummarizedExperiment::SummarizedExperiment(assay = a,</pre>
    rowData = rD, colData = cD)
tbl <- MAvalues(se)</pre>
hoeffDValues(tbl, "raw")
## normalized values
se_n <- se
assay(se_n) <- normalizeAssay(a, "sum")</pre>
tbl_n <- MAvalues(se_n, group = "all")</pre>
hoeffDValues(tbl_n, "normalized")
## transformed values
se_t <- se
assay(se_t) <- transformAssay(a, "log")</pre>
tbl_t <- MAvalues(se_t, group = "all")</pre>
hoeffDValues(tbl_t, "transformed")
```

imputeAssay

Impute missing values in a matrix

Description

The function impute imputes missing values based on one of the following principles: Bayesian missing value imputation (BPCA), k-nearest neighbor averaging (kNN), Malimum likelihood-based imputation method using the EM algorithm (MLE), replacement by the smallest non-missing value in the data (Min), replacement by the minimal value observed as the q-th quantile (MinDet, default q = 0.01), and replacement by random draws from a Gaussian distribution centred to a minimal value (MinProb).

Usage

```
imputeAssay(a, method = c("BPCA", "kNN", "MLE", "Min", "MinDet", "MinProb"))
```

Arguments

| а | matrix with samples in columns and features in rows |
|--------|---|
| method | character, one of "BPCA", "kNN", "MLE", "Min", "MinDet", or "MinProb" |

Details

BPCA wrapper for pcaMethods::pca with methods = "bpca". BPCA is a missing at random (MAR) imputation method.

kNN wrapper for impute::impute.knn with k = 10, rowmax = 0.5, colmax = 0.5, maxp = 1500. kNN is a MAR imputation method.

MLE wrapper for imputeLCMD::impute.MAR with method = "MLE", model.selector = 1/imputeLCMD::impute.wrapper.M MLE is a MAR imputation method.

Min imputes the missing values by the observed minimal value of x. Min is a missing not at random (MNAR) imputation method.

MinDet is a wrapper for imputeLCMD::impute.MinDet with q = 0.01. MinDet performs the imputation using a deterministic minimal value approach. The missing entries are replaced with a minimal value, estimated from the q-th quantile from each sample. MinDet is a MNAR imputation method.

MinProb is a wrapper for imputeLCMD: : impute.MinProb with q = 0.01 and tune.sigma = 1. MinProb performs the imputation based on random draws from a Gaussion distribution with the mean set to the minimal value of a sample. MinProb is a MNAR imputation method.

Value

matrix

Examples

```
a <- matrix(1:100, nrow = 10, ncol = 10,
    dimnames = list(1:10, paste("sample", 1:10)))
a[c(1, 5, 8), 1:5] <- NA
imputeAssay(a, method = "kNN")
imputeAssay(a, method = "Min")
imputeAssay(a, method = "MinDet")
imputeAssay(a, method = "MinProb")
```

MAplot

Create a MA plot

Description

The function creates a 2D histogram of M and A values.

Usage

```
MAplot(
   tbl,
   group = c("all", colnames(tbl)),
   plot = c("all", unique(tbl[["name"]]))
)
```

Arguments

| tbl | tibble containing the M and A values, as obtained from the MAvalues function |
|-------|--|
| group | character, one of colnames(colData(se)) (se used in MAvalues) or "all" |
| plot | character, one of colData(se)\$name (se used in MAvalues) or "all" |

MAvalues

Details

MAplot returns a 2D hex histogram instead of a classical scatterplot due to computational reasons and better visualization of overlaying points. The argument plot specifies the sample (refering to colData(se)\$name) to be plotted. If plot = "all", MA values for all samples will be plotted (samples will be plotted in facets). If the number of features (tbl\$Features) is below 1000, points will be plotted (via geom_points), otherwise hexagons will be plotted (via geom_hex).

Value

gg object from ggplot2

Examples

MAvalues

Create values (M and A) for MA plot

Description

The function MAvalues will create MA values as input for the function MAplot and hoeffDValues. M and A are specified relative to specified samples which is determined by the group argument. In case of group == "all", all samples (expect the specified one) are taken for the reference calculation. In case of group != "all" will use the samples belonging to the same group given in colnames(colData(se)) expect the specified one.

Usage

```
MAvalues(se, log2 = TRUE, group = c("all", colnames(colData(se))))
```

Arguments

| se | SummarizedExperiment |
|------|--|
| log2 | logical, specifies if values are log2-transformed prior to calculating M and A values. If the values are already transformed, log2 should be set to FALSE. If $log2 = TRUE$ and if there are values in assay(se) that are 0, the log2 values are calculated by log2(assay(se) + 1) |

character, either "all" or one of colnames(colData(se))

Value

group

tbl with columns Feature, name (sample name), A, M and additional columns of colData(se)

Examples

measuredCategory Obtain the number of measured intensities per sample type

Description

The function measuredCategory creates a tbl with the number of measured values per feature. 0 means that there were only missing values (NA) for the feature and sample type. measuredCategory will return a tbl where columns are the unique sample types and rows are the features as in assay(se).

Usage

```
measuredCategory(se, measured = TRUE, category = "type")
```

Arguments

| se | SummarizedExperiment |
|----------|---|
| measured | logical, should the measured values (measured = TRUE) or missing values (measured = FALSE) be taken |
| category | character, corresponds to a column name in colData(se) |

Details

measuredCategory is a helper function.

Value

matrix with number of measured/missing features per category type

mosaic

Examples

mosaic

Mosaic plot for two factors in colData(se)

Description

The function mosaic creates a mosaic plot of two factors from an SummarizedExperiment object. The columns f1 and f2 are taken from colData(se).

Usage

mosaic(se, f1, f2)

Arguments

| se | SummarizedExperiment object |
|----|---|
| f1 | character, f1 is one of the column names in colData(se) |
| f2 | character, f2 is one of the column names in colData(se) |

Details

Code partly taken from https://stackoverflow.com/questions/21588096/pass-string-to-facet-grid-ggplot2

Value

gg object from ggplot2

```
type = c(rep("1", 5), rep("2", 5)),
 cell_type = c("A", "B"))
rD <- data.frame(spectra = rownames(a))
se <- SummarizedExperiment::SummarizedExperiment(assay = a,
 rowData = rD, colData = cD)
mosaic(se, "cell_type", "type")
```

normalizeAssay Normalize a data sets (reduce technical sample effects)

Description

The function normalizeAssay performs normalization by sum of the (count/intensity) values per sample or quantile division per sample or by quantile normalization (adjusting the distributions that they become identical in statistical distributions). The divisor for quantile division (e.g., the 75 argument. Quantile normalization is performed by using the normalizeQuantiles function from limma.

Usage

```
normalizeAssay(
    a,
    method = c("none", "sum", "quantile division", "quantile"),
    probs = 0.75
)
```

Arguments

| а | matrix with samples in columns and features in rows |
|--------|--|
| method | character, one of "none", "sum", "quantile division", "quantile" |
| probs | <pre>numeric, ranging between [0, 1). probs is used as the divisor for quantile division in method = "quantile division"</pre> |

Details

Internal usage in shinyQC. If method is set to "none", the object x is returned as is (pass-through). If probs is NULL, probs is internally set to "name" if method = "quantile division".

Value

matrix

permuteExplVar

Examples

permuteExplVar Permute the expression values and retrieve the explained variance

Description

The function permuteExplVar determines the explained variance of the permuted expression matrix (x). It is used to determine the optimal number of PCs for tSNE.

Usage

permuteExplVar(x, n = 10, center = TRUE, scale = TRUE, sample_n = NULL)

Arguments

| х | matrix or data.frame, samples in columns and features in rows |
|----------|---|
| n | numeric, number of permutation rounds |
| center | logical, passed to the function explVar |
| scale | logical, passed to the function explVar |
| sample_n | numeric(1), number of features (subset) to be taken for calculation of permuted explained variance, the top sample_n varying values based on their standard deviation will be taken |

Details

For the input of tSNE, typically, we want to reduce the initial number of dimensions linearly with PCA (used as the initial_dims arguments in the Rtsne funtion). The reduced data set is used for feeding into tSNE. By plotting the percentage of variance explained by the Princical Components (PCs) we can estimate how many PCs we keep as input into tSNE. However, if we select too many PCs, noise will be included as input to tSNE; if we select too few PCs we might loose the important data structures. To get a better understanding how many PCs to include, randomization will be employed and the observed variance will be compared to the permuted variance.

Value

matrix with explained variance

Author(s)

Thomas Naake

Examples

```
x <- matrix(1:100, nrow = 10, ncol = 10,
    dimnames = list(1:10, paste("sample", 1:10)))
permuteExplVar(x = x, n = 10, center = TRUE, scale = TRUE, sample_n = NULL)
```

plotCV

Plot CV values

Description

The function plotCV displays the coefficient of variation values of set of values supplied in a data.frame object. The function will create a plot using the ggplot2 package and will print the values in the different columns in different colors.

Usage

plotCV(df)

Arguments

df

data.frame containing one or multiple columns containing the coefficients of variation

Details

Internal usage in shinyQC.

Value

gg object from ggplot2

Examples

```
x1 <- matrix(1:10, ncol = 2)
x2 <- matrix(11:20, ncol = 2)
x3 <- matrix(21:30, ncol = 2)
x4 <- matrix(31:40, ncol = 2)
## calculate cv values
cv1 <- cv(x1, "x1")
cv2 <- cv(x2, "x2")
cv3 <- cv(x3, "x3")
cv4 <- cv(x4, "x4")</pre>
```

df <- data.frame(cv1, cv2, cv3, cv4)
plotCV(df)</pre>

plotPCALoadings

Description

The function plotPCALoadings creates a loadings plot of the features.

Usage

plotPCALoadings(tbl, x_coord, y_coord)

Arguments

| tbl | tbl as obtained by the function dimensionReduction |
|---------|---|
| x_coord | character, column name of tbl that stores \boldsymbol{x} coordinates |
| y_coord | $\ensuremath{character}$, column name of tbl that stores y coordinates |

Details

The function takes as input the output of the function tblPlotPCALoadings. It uses the ggplotly function from plotly to create an interactive plotly plot.

Value

plotly

Author(s)

Thomas Naake

```
x <- matrix(rnorm(1:10000), ncol = 100)
rownames(x) <- paste("feature", 1:nrow(x))
colnames(x) <- paste("sample", 1:ncol(x))
params <- list(method = "euclidean", ## dist
    initial_dims = 10, max_iter = 100, dims = 3, perplexity = 3, ## tSNE
    min_dist = 0.1, n_neighbors = 15, spread = 1) ## UMAP
tbl <- tblPCALoadings(x, params)
plotPCALoadings(tbl, x_coord = "PC1", y_coord = "PC2")</pre>
```

plotPCAVar

Description

The function plotPCAVar plots the explained variance (in y-axis against the principal components for the measured and permuted values.

Usage

plotPCAVar(var_x, var_perm = NULL)

Arguments

| var_x | numeric (named numeric vector) |
|----------|---|
| var_perm | ${\tt matrix} \ with \ the \ explained \ variance \ obtained \ by \ permutation \ (function \ permute {\tt ExplVar})$ |

Details

The argument var_perm is optional and visualization of permuted values can be omitted by setting var_perm = NULL.

Value

gg object from ggplot

Author(s)

Thomas Naake

```
x <- matrix(1:100, ncol = 10)
pca <- dimensionReduction(x = x, params = list(center = TRUE, scale = TRUE),
    type = "PCA")[[2]]
var_x <- explVar(d = pca, type = "PCA")
var_perm <- permuteExplVar(x = x, n = 100, center = TRUE, scale = TRUE)
plotPCAVar(var_x = var_x, var_perm = var_perm)</pre>
```

plotPCAVarPvalue Plot p-values for the significance of principal components

Description

The function plotPCAVarPvalue plots the p-values of significances of principal components. Using the visual output, the optimal number of principal components can be selected.

Usage

plotPCAVarPvalue(var_x, var_perm)

Arguments

| var_x | numeric, measured variances |
|----------|---|
| var_perm | matrix, variances obtained by permutation |

Details

Internal usage in shinyQC.

Value

gg object from ggplot

Author(s)

Thomas Naake

```
x <- matrix(1:100, ncol = 10)
pca <- dimensionReduction(x = x, params = list(center = TRUE, scale = TRUE),
    type = "PCA")[[2]]
var_x <- explVar(d = pca, type = "PCA")
var_perm <- permuteExplVar(x = x, n = 100, center = TRUE, scale = TRUE)
plotPCAVarPvalue(var_x = var_x, var_perm = var_perm)</pre>
```

```
samplesMeasuredMissing
```

Create tibble containing number of measured/missing features of samples

Description

samplesMeasuredMissing returns a tbl with the number of measured/missing features of samples. The function will take as input a SummarizedExperiment object and will access its assay() slot

Usage

```
samplesMeasuredMissing(se)
```

Arguments

se SummarizedExperiment object

Value

tbl with number of measured/missing features per sample

Examples

samplesMeasuredMissing(se)

shinyQC

Description

The shiny application allows to explore -omics data sets especially with a focus on quality control. shinyQC gives information on the type of samples included (if this was previously specified within the SummarizedExperiment object). It gives information on the number of missing and measured values across features and across sets (e.g. quality control samples, control, and treatment groups, only displayed for SummarizedExperiment objects that contain missing values).

shinyQC includes functionality to display (count/intensity) values across samples (to detect drifts in intensity values during the measurement), to display mean-sd plots, MA plots, ECDF plots, and distance plots between samples. shinyQC includes functionality to perform dimensionality reduction (currently limited to PCA, PCoA, NMDS, tSNE, and UMAP). Additionally, it includes functionality to perform differential expression analysis (currently limited to moderated t-tests and the Wald test).

Usage

```
shinyQC(se, app_server = FALSE)
```

Arguments

| se | SummarizedExperiment object (can be omitted) |
|------------|---|
| app_server | logical (set to TRUE if run under a server environment) |

Details

rownames(se) should be set to the corresponding name of features, while colnames(se) should be set to the sample IDs. rownames(se) and colnames(se) are not allowed to be NULL. colnames(se), colnames(assay(se)) and rownames(colData(se)) all have to be identical.

shinyQC allows to subset the supplied SummarizedExperiment object.

On exit of the shiny application, the (subsetted) SummarizedExperiment object is returned with information on the processing steps (normalization, transformation, batch correction and imputation). The object will only returned if app_server = FALSE and if the function call is assigned to an object, e.g. tmp <- shinyQC(se).

If the se argument is omitted the app will load an interface that allows for data upload.

Value

shiny application, SummarizedExperiment upon exiting the shiny application

Author(s)

Thomas Naake

Examples

sumDistSample

Plot the sum of distances to other samples

Description

The function sumDistSample creates a plot showing the sum of distance of a sample to other samples.

Usage

```
sumDistSample(d, title = "raw")
```

Arguments

| d | matrix containing distances, obtained from distShiny |
|-------|--|
| title | character specifying the title to be added to the plot |

Value

gg object from ggplot2

Examples

tblPCALoadings

Description

The function tblPCALoadings returns a tibble with loadings values for the features (row entries) in x.

Usage

tblPCALoadings(x, params)

Arguments

| x | matrix, containing no missing values |
|--------|--|
| params | list, arguments/parameters given to the function stats::prcomp |

Details

The function tblPCALoadings accesses the list entry rotation of the prcomp object.

Value

tbl

Author(s)

Thomas Naake

```
set.seed(1)
x <- matrix(rnorm(1:10000), ncol = 100)
rownames(x) <- paste("feature", 1:nrow(x))
colnames(x) <- paste("sample", 1:ncol(x))
params <- list(method = "euclidean", ## dist
    initial_dims = 10, max_iter = 100, dims = 3, perplexity = 3, ## tSNE
    min_dist = 0.1, n_neighbors = 15, spread = 1) ## UMAP
tblPCALoadings(x, params)</pre>
```

transformAssay

Description

The function transformAssay transforms the (count/intensity) values of a matrix. It uses either log, log2, variance stabilizing normalisation (vsn) or no transformation method (pass-through, none). The object x has the samples in the columns and the features in the rows.

Usage

```
transformAssay(a, method = c("none", "log", "log2", "vsn"), .offset = 1)
```

Arguments

| а | matrix with samples in columns and features in rows |
|---------|---|
| method | character, one of "none", "log", "log2" or "vsn" |
| .offset | numeric(1), offset to add when method set to "log" or "log2" and a contains values of 0, default to 1 |

Details

Internal use in shinyQC.

Value

matrix

Description

The function upsetCategory displays the frequency of measured values per feature with respect to class/sample type to assess difference in occurrences. Internally, the measured values per sample are obtained via the measuredCategory function: this function will access the number of measured/missing values per category and feature. From this, a binary tbl will be created specifying if the feature is present/missing, which will be given to the upset function from the UpSetR package.

Usage

```
upsetCategory(se, category = colnames(colData(se)), measured = TRUE)
```

Arguments

| se | SummarizedExperiment, containing the intensity values in assay(se), missing values are encoded by NA |
|----------|--|
| category | character, corresponding to a column in colData(se) |
| measured | logical, should the measured values (measured = TRUE) or missing values (measured = FALSE) be taken |

Details

Presence is defined by a feature being measured in at least one sample of a set.

Absence is defined by a feature with only missing values (i.e. no measured values) of a set.

Value

upset plot

```
volcanoPlot
```

Description

The function ComplexHeatmap creates a volcano plot. On the y-axis the -log10(p-values) are displayed, while on the x-axis the fold changes/differences are displayed. The output of the function differs depending on the type parameter. For type == "ttest", the fold changes are plotted; for type == "proDA", the differences are plotted.

Usage

```
volcanoPlot(df, type = c("ttest", "proDA"))
```

Arguments

| df | data.frame as received from topTable (ttest) or test_diff (proDA) |
|------|---|
| type | character |

Details

Internal use in shinyQC.

Value

plotly

```
## create se
a <- matrix(1:100, nrow = 10, ncol = 10,
             dimnames = list(1:10, paste("sample", 1:10)))
a[c(1, 5, 8), 1:5] <- NA
set.seed(1)
a <- a + rnorm(100)
a_i <- imputeAssay(a, method = "MinDet")</pre>
cD <- data.frame(sample = colnames(a),</pre>
    type = c(rep("1", 5), rep("2", 5)))
rD <- data.frame(spectra = rownames(a))</pre>
se <- SummarizedExperiment::SummarizedExperiment(assay = a,</pre>
                                  rowData = rD, colData = cD)
se_i <- SummarizedExperiment::SummarizedExperiment(assay = a_i,</pre>
                                   rowData = rD, colData = cD)
## create model and contrast matrix
modelMatrix_expr <- stats::formula("~ 0 + type")</pre>
contrast_expr <- "type1-type2"</pre>
modelMatrix <- model.matrix(modelMatrix_expr, data = colData(se))</pre>
contrastMatrix <- limma::makeContrasts(contrasts = contrast_expr,</pre>
```

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