

# Package ‘NormalyzerDE’

December 16, 2025

**Title** Evaluation of normalization methods and calculation of differential expression analysis statistics

**Version** 1.28.0

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**Description** NormalyzerDE provides screening of normalization methods for LC-MS based expression data. It calculates a range of normalized matrices using both existing approaches and a novel time-segmented approach, calculates performance measures and generates an evaluation report. Furthermore, it provides an easy utility for Limma- or ANOVA- based differential expression analysis.

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---

analyzeNormalizations    *Calculate measures for normalization results*

---

## Description

This function prepares an `NormalizerEvaluationResults` object containing the evaluation measures CV (coefficient of variance), MAD (median absolute deviation), average variance, significance measures (ANOVA between condition groups) and correlation between replicates.

**Usage**

```
analyzeNormalizations(nr, categoricalAnova = FALSE)
```

**Arguments**

```
nr           Normalizer results object with calculated results.
categoricalAnova  Whether categorical or numerical (ordered) ANOVA should be calculated.
```

**Value**

Normalizer results with attached evaluation results object.

**Examples**

```
data(example_summarized_experiment)
normObj <- getVerifiedNormalizerObject("job_name", example_summarized_experiment)
normResults <- normMethods(normObj)
normResultsWithEval <- analyzeNormalizations(normResults)
```

---

`calculateANOVAPValues` *Calculates ANOVA p-values comparing the different condition groups returning a vector of resulting p-values with NA-values where too few values were present in at least one of the groups.*

---

**Description**

Calculates ANOVA p-values comparing the different condition groups returning a vector of resulting p-values with NA-values where too few values were present in at least one of the groups.

**Usage**

```
calculateANOVAPValues(methodList, sampleReplicateGroups, categoricalANOVA)
```

**Arguments**

```
methodList      List containing normalized matrices
sampleReplicateGroups  Condition header
categoricalANOVA  Whether the ANOVA should be calculated using ordered or categorical groups
```

**Value**

avgVarianceMat Matrix with average variance for each biological condition

---

|                    |                                                                                                                                                      |
|--------------------|------------------------------------------------------------------------------------------------------------------------------------------------------|
| calculateAvgMadMem | <i>Calculate average MAD (Median Absolute Deviation) for each feature in each condition and then calculates the average for each replicate group</i> |
|--------------------|------------------------------------------------------------------------------------------------------------------------------------------------------|

---

**Description**

Calculate average MAD (Median Absolute Deviation) for each feature in each condition and then calculates the average for each replicate group

**Usage**

```
calculateAvgMadMem(methodList, sampleReplicateGroups)
```

**Arguments**

|                       |                                      |
|-----------------------|--------------------------------------|
| methodList            | List containing normalized matrices. |
| sampleReplicateGroups | Condition header.                    |

**Value**

condAvgMadMat Matrix with average MAD for each biological condition.

---

|                                |                                                                                                                              |
|--------------------------------|------------------------------------------------------------------------------------------------------------------------------|
| calculateAvgReplicateVariation | <i>Calculate average variance for each feature in each condition and then calculate the average for each replicate group</i> |
|--------------------------------|------------------------------------------------------------------------------------------------------------------------------|

---

**Description**

Calculate average variance for each feature in each condition and then calculate the average for each replicate group

**Usage**

```
calculateAvgReplicateVariation(methodList, sampleReplicateGroups)
```

**Arguments**

|                       |                                      |
|-----------------------|--------------------------------------|
| methodList            | List containing normalized matrices. |
| sampleReplicateGroups | Condition header.                    |

**Value**

avgVarianceMat Matrix with average variance for each biological condition

---

|                    |                                                                                                                                                                                                                                                                                                                                                                                                         |
|--------------------|---------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| calculateContrasts | <i>Performs statistical comparisons between the supplied conditions. It uses the design matrix and data matrix in the supplied Normalyzer-Statistics object. A column is supplied specifying which of the columns in the design matrix that is used for deciding the sample groups. The comparisons vector specifies which pairwise comparisons between condition levels that are to be calculated.</i> |
|--------------------|---------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|

---

## Description

Optionally, a batch column can be specified allowing compensation for covariate variation in the statistical model. This is only compatible with a Limma-based statistical analysis.

## Usage

```
calculateContrasts(
  nst,
  comparisons,
  condCol,
  batchCol = NULL,
  splitter = "-",
  type = "limma",
  leastRepCount = 1
)

## S4 method for signature 'NormalyzerStatistics'
calculateContrasts(
  nst,
  comparisons,
  condCol,
  batchCol = NULL,
  splitter = "-",
  type = "limma",
  leastRepCount = 1
)
```

## Arguments

|               |                                                                         |
|---------------|-------------------------------------------------------------------------|
| nst           | Results evaluation object.                                              |
| comparisons   | String with comparisons for contrasts.                                  |
| condCol       | Column name in design matrix containing condition information.          |
| batchCol      | Column name in design matrix containing batch information.              |
| splitter      | Character dividing contrast conditions.                                 |
| type          | Type of statistical test (Limma or welch).                              |
| leastRepCount | Least replicates in each group to be retained for contrast calculations |

## Value

nst Statistics object with statistical measures calculated

**Examples**

```
data(example_stat_summarized_experiment)
nst <- NormalyzerStatistics(example_stat_summarized_experiment)
results <- calculateContrasts(nst, c("1-2", "2-3"), "group")
resultsBatch <- calculateContrasts(nst, c("1-2", "2-3"), "group", batchCol="batch")
```

---

|                  |                                                                                                                                                                     |
|------------------|---------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| calculateCorrSum | <i>Calculates internal correlations for each condition having at least two samples and returns a vector with correlation values corresponding to each condition</i> |
|------------------|---------------------------------------------------------------------------------------------------------------------------------------------------------------------|

---

**Description**

Calculates internal correlations for each condition having at least two samples and returns a vector with correlation values corresponding to each condition

**Usage**

```
calculateCorrSum(
  methodData,
  allReplicateGroups,
  sampleGroupsWithReplicates,
  corrType
)
```

**Arguments**

|                            |                                                            |
|----------------------------|------------------------------------------------------------|
| methodData                 | Expression data matrix                                     |
| allReplicateGroups         | Full condition header corresponding to data tables columns |
| sampleGroupsWithReplicates | Unique conditions where number of replicates exceeds one   |
| corrType                   | Type of correlation (Pearson or Spearman)                  |

**Value**

corSums

---

|                    |                                                                                                                                                                                                              |
|--------------------|--------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| calculateFeatureCV | <i>Calculate CV values for each feature. Iterates through each normalization method and calculates a matrix of CV values where each column correspond to a method and each row corresponds to a feature.</i> |
|--------------------|--------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|

---

**Description**

Calculate CV values for each feature. Iterates through each normalization method and calculates a matrix of CV values where each column correspond to a method and each row corresponds to a feature.

**Usage**

```
calculateFeatureCV(methodList)
```

**Arguments**

```
methodList      List containing normalized matrices.
sampleReplicateGroups
                  Condition header.
```

**Value**

```
methodFeatureCVMatrix Matrix with feature as rows and normalization method as columns
```

---

```
calculatePercentageAvgDiffInMat
```

*General function for calculating percentage difference of average column means in matrix*

---

**Description**

General function for calculating percentage difference of average column means in matrix

**Usage**

```
calculatePercentageAvgDiffInMat(targetMat)
```

**Arguments**

```
targetMat      Matrix for which column means should be compared
```

**Value**

```
percDiffVector Vector with percentage difference, where first element always will be 100
```

---

```
calculateReplicateCV    Calculate CV per replicate group and normalization technique
```

---

**Description**

Iterates through each normalization method and calculate average CV values per replicate group.

**Usage**

```
calculateReplicateCV(methodList, sampleReplicateGroups)
```

**Arguments**

```
methodList      List containing normalized matrices.
sampleReplicateGroups
                  Condition header.
```



**Value**

avgCVPerNormAndReplicates Matrix with group CVs as rows and normalization technique as columns

---

calculateSummarizedCorrelationVector

*Calculates correlation values between replicates for each condition matrix. Finally returns a matrix containing the results for all dataset*

---

**Description**

Calculates correlation values between replicates for each condition matrix. Finally returns a matrix containing the results for all dataset

**Usage**

```
calculateSummarizedCorrelationVector(
  methodlist,
  allReplicateGroups,
  sampleGroupsWithReplicates,
  corrType
)
```

**Arguments**

|                            |                                                                                      |
|----------------------------|--------------------------------------------------------------------------------------|
| methodlist                 | List containing normalized matrices for each normalization method                    |
| allReplicateGroups         | Vector with condition groups matching the columns found in the normalization methods |
| sampleGroupsWithReplicates | Unique vector with condition groups present in two or more samples                   |
| corrType                   | Type of correlation (Pearson or Spearman)                                            |

**Value**

avgCorSum Matrix with column corresponding to normalization approaches and rows corresponding to replicate group

---

|                 |                                                             |
|-----------------|-------------------------------------------------------------|
| createDirectory | <i>Create directory, or return error if already present</i> |
|-----------------|-------------------------------------------------------------|

---

**Description**

Create directory, or return error if already present

**Usage**

```
createDirectory(targetPath)
```

**Arguments**

targetPath      Path where to attempt to create directory

**Value**

None

---

detectSingleReplicate    *Detect single replicate, and assign related logical*

---

**Description**

Detect single replicate, and assign related logical

**Usage**

```
detectSingleReplicate(nds, quiet)

## S4 method for signature 'NormalizerDataset'
detectSingleReplicate(nds, quiet = FALSE)
```

**Arguments**

nds              Normalizer dataset  
quiet            Don't give non-error output

**Value**

bool on whether sample contains only one sample group

---

detectSingletonSample    *Detect single sample group*

---

**Description**

Detect single sample group

**Usage**

```
detectSingletonSample(nds, quiet)

## S4 method for signature 'NormalizerDataset'
detectSingletonSample(nds, quiet = FALSE)
```

**Arguments**

nds              Normalizer dataset.  
quiet            Only print error messages

**Value**

None

---

`elapsedSecondsBetweenSystimes`*Get number of seconds between two Sys.time() objects*

---

**Description**

Get number of seconds between two Sys.time() objects

**Usage**

```
elapsedSecondsBetweenSystimes(start, end)
```

**Arguments**

|                    |                   |
|--------------------|-------------------|
| <code>start</code> | Start-time object |
|--------------------|-------------------|

|                  |                 |
|------------------|-----------------|
| <code>end</code> | End-time object |
|------------------|-----------------|

**Value**

None

---

`example_data`*Small example dataset used to demonstrate code consistency in testing and as dummy data in the vignette.*

---

**Description**

Small example dataset used to demonstrate code consistency in testing and as dummy data in the vignette.

**Usage**

```
example_data
```

**Format**

A data frame containing annotation and expression data

---

example\_data\_only\_values

*Same data as in "example\_data", but omitting the annotation meaning that it only contains the expression data.*

---

**Description**

Same data as in "example\_data", but omitting the annotation meaning that it only contains the expression data.

**Usage**

example\_data\_only\_values

**Format**

A data frame containing expression data

---

example\_design

*Design matrix corresponding to the small example datasets.*

---

**Description**

Design matrix corresponding to the small example datasets.

**Usage**

example\_design

**Format**

A design matrix corresponding to the dataset "example\_data"

---

example\_stat\_data

*Same data as in "example\_data", but normalized and ready for statistical processing.*

---

**Description**

Same data as in "example\_data", but normalized and ready for statistical processing.

**Usage**

example\_stat\_data

**Format**

A normalized data frame ready for statistical processing

---

`example_stat_summarized_experiment`*SummarizedExperiment object prepared with design-matrix, data-matrix and annotation columns for normalized data*

---

**Description**

SummarizedExperiment object prepared with design-matrix, data-matrix and annotation columns for normalized data

**Usage**`example_stat_summarized_experiment`**Format**

An instance of the class SummarizedExperiment with stats data

---

`example_summarized_experiment`*SummarizedExperiment object prepared with design-matrix, data-matrix and annotation columns loaded for raw data*

---

**Description**

SummarizedExperiment object prepared with design-matrix, data-matrix and annotation columns loaded for raw data

**Usage**`example_summarized_experiment`**Format**

An instance of the class SummarizedExperiment

---

`example_wide_data`*Full raw NormalyzerDE matrix used for internal testing*

---

**Description**

Full raw NormalyzerDE matrix used for internal testing

**Usage**`example_wide_data`**Format**

A data table ready for analysis in NormalyzerDE

---

|                     |                                                                                            |
|---------------------|--------------------------------------------------------------------------------------------|
| example_wide_design | <i>Design matrix belonging together with example_wide_data. Used for internal testing.</i> |
|---------------------|--------------------------------------------------------------------------------------------|

---

### Description

Design matrix belonging together with example\_wide\_data. Used for internal testing.

### Usage

```
example_wide_design
```

### Format

A design table ready for analysis in NormalyzerDE

---

|              |                                                                                 |
|--------------|---------------------------------------------------------------------------------|
| filterLowRep | <i>Filter rows with lower than given number of replicates for any condition</i> |
|--------------|---------------------------------------------------------------------------------|

---

### Description

Filter rows with lower than given number of replicates for any condition

### Usage

```
filterLowRep(df, groups, leastRep = 2)
```

### Arguments

|          |                                                      |
|----------|------------------------------------------------------|
| df       | Dataframe with expression data to filter             |
| groups   | Condition groups header                              |
| leastRep | Minimum number of replicates in each group to retain |

### Value

collDesignDf Reduced design matrix

---

findLowlyVariableFeaturesCVs

*Uses a list of FDR-values to extract features with low variance in the log2-transformed dataset. This is then used to calculate the average CV for these 'lowly variable' features in each normalization approach*

---

### Description

Uses a list of FDR-values to extract features with low variance in the log2-transformed dataset. This is then used to calculate the average CV for these 'lowly variable' features in each normalization approach

### Usage

```
findLowlyVariableFeaturesCVs(referenceFDR, methodList)
```

### Arguments

|              |                                                     |
|--------------|-----------------------------------------------------|
| referenceFDR | List of FDR values used as non-normalized reference |
| methodList   | List containing normalized matrices                 |

### Value

lowVarFeaturesAverageCVs Average CV values for lowly variable features in each normalization approach

---

generateAnnotatedMatrix

*Generate an annotated data frame from statistics object*

---

### Description

Extracts key values (p-value, adjusted p-value, log2-fold change and average expression values) from an NormalyzerStatistics instance and appends these to the annotation- and data-matrices

### Usage

```
generateAnnotatedMatrix(nst, prefixSep = "_", compLabels = NULL)
```

### Arguments

|            |                                                                             |
|------------|-----------------------------------------------------------------------------|
| nst        | NormalyzerDE statistics object.                                             |
| prefixSep  | Character string for separating the prefix names from the statistics suffix |
| compLabels | Vector containing strings to use as prefix for statistical comparisons      |

### Value

outDf Annotated statistics matrix

**Examples**

```
data(example_stat_summarized_experiment)
statObj <- NormalyzerStatistics(example_stat_summarized_experiment)
statObj <- calculateContrasts(statObj, comparisons=c("1-2", "2-3"), condCol="group", type="limma")
annotDf <- generateAnnotatedMatrix(statObj)
```

---

|               |                                                                                                                                                                                                                     |
|---------------|---------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| generatePlots | <i>Generates a number of visualizations for the performance measures calculated for the normalized matrices. These contain both general measures and direct comparisons for different normalization approaches.</i> |
|---------------|---------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|

---

**Description**

They include:

**Usage**

```
generatePlots(nr, jobdir, plotRows = 3, plotCols = 4, writeAsPngs = FALSE)
```

**Arguments**

|             |                                                        |
|-------------|--------------------------------------------------------|
| nr          | Normalyzer results object.                             |
| jobdir      | Path to output directory for run.                      |
| plotRows    | Number of plot rows.                                   |
| plotCols    | Number of plot columns.                                |
| writeAsPngs | Output the report as PNG-plots instead of a single PDF |

**Details**

"Total intensity" Barplot showing the summed intensity in each sample for thelog2-transformed data

"Total missing" Barplot showing the number of missing values found in each sample for the log2-transformed data

Log2-MDS plot: MDS plot where data is reduced to two dimensions allowing inspection of the main global changes in the data

PCV - Intragroup: Mean of intragroup CV of all replicate groups

PMAD - Intragroup: Mean of intragroup median absolute deviation across replicate groups

PEV - Intragroup: Mean of intragroup pooled estimate of variance across the replicate groups

Relative PCV, PMAD and PEV compared to log2: The results from PCV, PMAD and PEV from all normalized data compared to the log2 data

Stable variables plot: 5 analysis of log2 transformed data. Thereafter, global CV of these variables is estimated from different normalized datasets. A plot of global CV of the stable variables from all datasets on the y-axis and PCV-compared to log2 on the x-axis is generated.

CV vs Raw Intensity plots: For the first replicate group in each of the normalized dataset, a plot of PCV of each variable compared to the average intensity of the variable in the replicate group is plotted.



MA plots: Plotted using the plotMA function of the limma package. The first sample in each dataset is plotted against the average of the replicate group that sample belong to.

Scatterplots: The first two samples from each dataset are plotted.

Q-Q plots: QQ-plots are plotted for the first sample in each normalized dataset.

Boxplots: Boxplots for all samples are plotted and colored according to the replicate grouping.

Relative Log Expression (RLE) plots: Relative log expression value plots. Ratio between the expression of the variable and the median expression of this variable across all samples. The samples should be aligned around zero. Any deviation would indicate discrepancies in the data.

Density plots: Density distributions for each sample using the density function. Can capture outliers (if single densities lies far from the others) and see if there is batch effects in the dataset (if for instance there is two clear collections of lines in the data).

MDS plots Multidimensional scaling plot using the cmdscale() function from the stats package. Is often able to show whether replicates group together, and whether there are any clear outliers in the data.

MeanSDplots Displays the standard deviation values against values ordered according to mean. If no dependency on mean is present (as is desired) a flat red line is shown.

Pearson and Spearman correlation Mean of intragroup Pearson and Spearman correlation values for each method.

Dendograms Generated using the hclust function. Data is centered and scaled prior to analysis. Coloring of replicates is done using as.phylo from the ape package.

P-value histograms Histogram plots of p-values after calculating an ANOVA between different condition groups. If no effect is present in the data a flat distribution is expected. If an effect is present a flat distribution is still expected, but with a sharp peak close to zero. If other effects are present it might indicate that the data doesn't support the assumptions of ANOVA, for instance if there are batch effects present in the data.

## Value

None

## Examples

```
data(example_summarized_experiment)
normObj <- getVerifiedNormalizerObject("job_name", example_summarized_experiment)
normResults <- normMethods(normObj)
normResultsWithEval <- analyzeNormalizations(normResults)
outputDir <- tempdir()
generatePlots(normResultsWithEval, outputDir)
```

---

|                     |                                                                                                                                                                     |
|---------------------|---------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| generateStatsReport | <i>Generate full output report plot document. Plots p-value histograms for each contrast in the NormalizerStatistics instance and writes these to a PDF report.</i> |
|---------------------|---------------------------------------------------------------------------------------------------------------------------------------------------------------------|

---

## Description

Generate full output report plot document. Plots p-value histograms for each contrast in the NormalizerStatistics instance and writes these to a PDF report.

**Usage**

```
generateStatsReport(
  nst,
  jobName,
  jobDir,
  sigThres = 0.1,
  sigThresType = "fdr",
  log2FoldThres = 0,
  plotRows = 3,
  plotCols = 4,
  writeAsPngs = FALSE
)
```

**Arguments**

|               |                                                                      |
|---------------|----------------------------------------------------------------------|
| nst           | NormalyzerDE statistics object.                                      |
| jobName       | Name of processing run.                                              |
| jobDir        | Path to output directory.                                            |
| sigThres      | Significance threshold for indicating as significant                 |
| sigThresType  | Type of significance threshold (FDR or p)                            |
| log2FoldThres | log2 fold-change required for being counted as significant           |
| plotRows      | Number of plot rows.                                                 |
| plotCols      | Number of plot columns.                                              |
| writeAsPngs   | Output the report as separate PNG files instead of a single PDF file |

**Value**

None

**Examples**

```
data(example_stat_summarized_experiment)
statObj <- NormalyzerStatistics(example_stat_summarized_experiment)
statObj <- calculateContrasts(statObj, comparisons=c("1-2", "2-3"),
  condCol="group", type="limma")
outputDir <- tempdir()
generateStatsReport(statObj, "jobName", outputDir)
```

---

|                   |                                                          |
|-------------------|----------------------------------------------------------|
| getCombinedMatrix | <i>Merge multiple dataframes using provided function</i> |
|-------------------|----------------------------------------------------------|

---

**Description**

Merge multiple dataframes using provided function

**Usage**

```
getCombinedMatrix(mList, combFunc)
```

### Arguments

|          |                                                   |
|----------|---------------------------------------------------|
| mList    | List containing dataframes of same shape          |
| combFunc | Function performing elementwise merge of matrices |

### Value

combinedMatrix A single dataframe with combined data

---

|              |                                                                    |
|--------------|--------------------------------------------------------------------|
| getIndexList | <i>Return list containing vector positions of values in string</i> |
|--------------|--------------------------------------------------------------------|

---

### Description

Return list containing vector positions of values in string

### Usage

```
getIndexList(targetVector)
```

### Arguments

targetVector

### Value

indexList List where key is condition level and values are indices for the condition

---

|                           |                                                                       |
|---------------------------|-----------------------------------------------------------------------|
| getLowCountSampleFiltered | <i>Verify that samples contain at least a lowest number of values</i> |
|---------------------------|-----------------------------------------------------------------------|

---

### Description

Verify that samples contain at least a lowest number of values

### Usage

```
getLowCountSampleFiltered(
  dataMatrix,
  groups,
  threshold = 15,
  stopIfTooFew = TRUE
)
```

### Arguments

|              |                                                              |
|--------------|--------------------------------------------------------------|
| dataMatrix   | Dataframe with processed input data.                         |
| groups       | Vector containing condition levels.                          |
| threshold    | Lowest number of allowed values in a column.                 |
| stopIfTooFew | Abort run if lower than threshold number of values in column |

**Value**

None

---

 getReplicateSortedData

*Get dataframe with raw data column sorted on replicates*


---

**Description**

Get dataframe with raw data column sorted on replicates

**Usage**

```
getReplicateSortedData(rawDataOnly, groups)
```

**Arguments**

|             |                                            |
|-------------|--------------------------------------------|
| rawDataOnly | Dataframe with unparsed input data matrix. |
| groups      | Vector containing condition levels.        |

**Value**

rawData sorted on replicate

---

 getRowNAFilterContrast

*Get contrast vector (TRUE/FALSE-values) indicating whether both at least half values are present, and each sample has at least one non-NA value*


---

**Description**

Get contrast vector (TRUE/FALSE-values) indicating whether both at least half values are present, and each sample has at least one non-NA value

**Usage**

```
getRowNAFilterContrast(dataMatrix, replicateHeader, minCount = 1)
```

**Arguments**

|                 |                                                                  |
|-----------------|------------------------------------------------------------------|
| dataMatrix      | Matrix with expression values for entities in replicate samples. |
| replicateHeader | Header showing how samples in matrix are replicated.             |
| minCount        | Minimum number of required values present in samples.            |

**Value**

Contrast vector

---

`getRTNormalizedMatrix` *Perform RT-segmented normalization by performing the supplied normalization over retention-time sliced data*

---

### Description

The function orders the retention times and steps through them using the supplied step size (in minutes). If smaller than a fixed lower boundary the window is expanded to ensure a minimum amount of data in each normalization step. An offset can be specified which can be used to perform multiple RT-segmentations with partial overlapping windows.

### Usage

```
getRTNormalizedMatrix(
  rawMatrix,
  retentionTimes,
  normMethod,
  stepSizeMinutes = 1,
  windowMinCount = 100,
  offset = 0,
  noLogTransform = FALSE
)
```

### Arguments

|                              |                                                                   |
|------------------------------|-------------------------------------------------------------------|
| <code>rawMatrix</code>       | Target matrix to be normalized                                    |
| <code>retentionTimes</code>  | Vector of retention times corresponding to <code>rawMatrix</code> |
| <code>normMethod</code>      | The normalization method to apply to the time windows             |
| <code>stepSizeMinutes</code> | Size of windows to be normalized                                  |
| <code>windowMinCount</code>  | Minimum number of values for window to not be expanded.           |
| <code>offset</code>          | Whether time window should shifted half step size                 |
| <code>noLogTransform</code>  | Don't log-transform the data                                      |

### Value

Normalized matrix

### Examples

```
data(example_data_small)
data(example_design_small)
data(example_data_only_values)
dataMat <- example_data_only_values
retentionTimes <- as.numeric(example_data[, "Average.RT"])
performCyclicLoessNormalization <- function(rawMatrix) {
  log2Matrix <- log2(rawMatrix)
  normMatrix <- limma::normalizeCyclicLoess(log2Matrix, method="fast")
  colnames(normMatrix) <- colnames(rawMatrix)
  normMatrix
}
```

```
rtNormMat <- getRTNormalizedMatrix(dataMat, retentionTimes,
performCyclicLoessNormalization, stepSizeMinutes=1, windowMinCount=100)
```

---

```
getSmoothedRTNormalizedMatrix
```

*Generate multiple RT time-window normalized matrices where one is shifted. Merge them using a specified method (mean or median) and return the result.*

---

## Description

Uses the function `getRTNormalizedMatrix` to generate multiple normalized matrices which are shifted respective to each other and finally merged into a single matrix. This could potentially reduce effect of fluctuations within individual windows.

## Usage

```
getSmoothedRTNormalizedMatrix(
  rawMatrix,
  retentionTimes,
  normMethod,
  stepSizeMinutes,
  windowShifts = 2,
  windowMinCount = 100,
  mergeMethod = "mean",
  noLogTransform = FALSE
)
```

## Arguments

|                              |                                                                   |
|------------------------------|-------------------------------------------------------------------|
| <code>rawMatrix</code>       | Target matrix to be normalized                                    |
| <code>retentionTimes</code>  | Vector of retention times corresponding to <code>rawMatrix</code> |
| <code>normMethod</code>      | The normalization method to apply to the time windows             |
| <code>stepSizeMinutes</code> | Size of windows to be normalized                                  |
| <code>windowShifts</code>    | Number of frame shifts.                                           |
| <code>windowMinCount</code>  | Minimum number of features within window.                         |
| <code>mergeMethod</code>     | Layer merging approach. Mean or median.                           |
| <code>noLogTransform</code>  | Don't log transform the input                                     |

## Value

Normalized matrix

**Examples**

```

data(example_data_small)
data(example_data_only_values)
data(example_design_small)
retentionTimes <- as.numeric(example_data[, "Average.RT"])
dataMat <- example_data_only_values
performCyclicLoessNormalization <- function(rawMatrix) {
  log2Matrix <- log2(rawMatrix)
  normMatrix <- limma::normalizeCyclicLoess(log2Matrix, method="fast")
  colnames(normMatrix) <- colnames(rawMatrix)
  normMatrix
}
rtNormMat <- getSmoothedRTNormalizedMatrix(dataMat, retentionTimes,
  performCyclicLoessNormalization, stepSizeMinutes=1, windowMinCount=100,
  windowShifts=2, mergeMethod="median")

```

---

```
getVerifiedNormalizerObject
```

*Verify that input data is in correct format, and if so, return a generated NormalizerDE data object from that input data*

---

**Description**

This function performs a number of checks on the input data and provides informative error messages if the data isn't fulfilling the required format. Checks include verifying that the design matrix matches to the data matrix, that the data matrix contains valid numbers and that samples have enough values for analysis

**Usage**

```

getVerifiedNormalizerObject(
  jobName,
  summarizedExp,
  threshold = 15,
  omitSamples = FALSE,
  requireReplicates = TRUE,
  quiet = FALSE,
  noLogTransform = FALSE,
  tinyRunThres = 50
)

```

**Arguments**

|                   |                                                       |
|-------------------|-------------------------------------------------------|
| jobName           | Name of ongoing run.                                  |
| summarizedExp     | Summarized experiment input object                    |
| threshold         | Minimum number of features.                           |
| omitSamples       | Automatically omit invalid samples from analysis.     |
| requireReplicates | Require there to be at least to samples per condition |
| quiet             | Don't print output messages during processing         |
| noLogTransform    | Don't log-transform the provided data                 |
| tinyRunThres      | If less features in run, a limited run is performed   |

**Value**

Normalyzer data object representing verified input data.

**Examples**

```
data(example_summarized_experiment)
normObj <- getVerifiedNormalyzerObject("job_name", example_summarized_experiment)
```

---

|                   |                                                                                                                                                                                |
|-------------------|--------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| getWidenedRTRange | <i>Pick datapoints before and after window until a minimum number is reached Expects the start and end retention times to match actual retention times present in the data</i> |
|-------------------|--------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|

---

**Description**

Pick datapoints before and after window until a minimum number is reached Expects the start and end retention times to match actual retention times present in the data

**Usage**

```
getWidenedRTRange(
  rtStart,
  rtEnd,
  minimumDatapoints,
  retentionTimes,
  allowTooWideData = FALSE
)
```

**Arguments**

|                   |                                          |
|-------------------|------------------------------------------|
| rtStart           | Original retention time start point      |
| rtEnd             | Original retention time end point        |
| minimumDatapoints | Required number of datapoints to fulfill |
| retentionTimes    | Vector with all retention times          |

**Value**

Vector with start and end of new RT range



---

**globalIntensityNormalization**

*The normalization divides the intensity of each variable in a sample with the sum of intensities of all variables in the sample and multiplies with the median of sum of intensities of all variables in all samples. The normalized data is then log2-transformed.*

---

**Description**

The normalization divides the intensity of each variable in a sample with the sum of intensities of all variables in the sample and multiplies with the median of sum of intensities of all variables in all samples. The normalized data is then log2-transformed.

**Usage**

```
globalIntensityNormalization(rawMatrix, noLogTransform = FALSE)
```

**Arguments**

rawMatrix      Target matrix to be normalized  
noLogTransform   Assumes no need for log transformation

**Value**

Normalized and log-transformed matrix

**Examples**

```
data(example_data_only_values_small)
normMatrix <- globalIntensityNormalization(example_data_only_values)
```

---

|          |                                     |
|----------|-------------------------------------|
| loadData | <i>Load raw data into dataframe</i> |
|----------|-------------------------------------|

---

**Description**

General function which allows specifying different types of input data including "proteios", "maxquant-pep" (peptide output from MaxQuant) and "maxquantprot" (protein output from MaxQuant) formats.

**Usage**

```
loadData(dataPath, inputFormat = "default")
```

**Arguments**

dataPath      File path to design matrix.  
inputFormat   If input is given in standard NormalyzerDE format, Proteios format or in MaxQuant protein or peptide format

**Value**

rawData Raw data loaded into data frame

**Examples**

```
## Not run:  
df <- loadData("data.tsv")  
  
## End(Not run)
```

---

|            |                                       |
|------------|---------------------------------------|
| loadDesign | <i>Load raw design into dataframe</i> |
|------------|---------------------------------------|

---

**Description**

Takes a design path, loads the matrix and ensures that the sample column is in character format and that the group column is in factor format.

**Usage**

```
loadDesign(designPath, sampleCol = "sample", groupCol = "group")
```

**Arguments**

|            |                                                     |
|------------|-----------------------------------------------------|
| designPath | File path to design matrix.                         |
| sampleCol  | Column name for column containing sample names.     |
| groupCol   | Column name for column containing condition levels. |

**Value**

designMatrix Design data loaded into data frame

**Examples**

```
## Not run:  
df <- loadDesign("design.tsv")  
  
## End(Not run)
```

---

|                     |                                                                 |
|---------------------|-----------------------------------------------------------------|
| loadRawDataFromFile | <i>Try reading raw Normalyzer matrix from provided filepath</i> |
|---------------------|-----------------------------------------------------------------|

---

### Description

Try reading raw Normalyzer matrix from provided filepath

### Usage

```
loadRawDataFromFile(inputPath)
```

### Arguments

|           |                          |
|-----------|--------------------------|
| inputPath | Path to Normalyzer data. |
|-----------|--------------------------|

### Value

Table containing raw data from input file.

---

|                   |                                                                                                                                                                                                                                                                         |
|-------------------|-------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| meanNormalization | <i>Intensity of each variable in a given sample is divided by the mean of sum of intensities of all variables in the sample and then multiplied by the mean of sum of intensities of all variables in all samples. The normalized data is then transformed to log2.</i> |
|-------------------|-------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|

---

### Description

Intensity of each variable in a given sample is divided by the mean of sum of intensities of all variables in the sample and then multiplied by the mean of sum of intensities of all variables in all samples. The normalized data is then transformed to log2.

### Usage

```
meanNormalization(rawMatrix, noLogTransform = FALSE)
```

### Arguments

|                |                                        |
|----------------|----------------------------------------|
| rawMatrix      | Target matrix to be normalized         |
| noLogTransform | Assumes no need for log transformation |

### Value

Normalized and log-transformed matrix

### Examples

```
data(example_data_only_values_small)
normMatrix <- meanNormalization(example_data_only_values)
```

---

|                     |                                                                                                                                                                                                                                                                           |
|---------------------|---------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| medianNormalization | <i>Intensity of each variable in a given sample is divided by the median of intensities of all variables in the sample and then multiplied by the mean of median of sum of intensities of all variables in all samples. The normalized data is then log2-transformed.</i> |
|---------------------|---------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|

---

### Description

Intensity of each variable in a given sample is divided by the median of intensities of all variables in the sample and then multiplied by the mean of median of sum of intensities of all variables in all samples. The normalized data is then log2-transformed.

### Usage

```
medianNormalization(rawMatrix, noLogTransform = FALSE)
```

### Arguments

rawMatrix      Target matrix to be normalized  
noLogTransform   Assumes no need for log transformation

### Value

Normalized and log-transformed matrix

### Examples

```
data(example_data_only_values_small)
normMatrix <- medianNormalization(example_data_only_values)
```

---

|            |                                          |
|------------|------------------------------------------|
| normalyzer | <i>NormalyzerDE pipeline entry point</i> |
|------------|------------------------------------------|

---

### Description

This function is the main execution point for the normalization part of the NormalyzerDE analysis pipeline. When executed it performs the following steps:

### Usage

```
normalyzer(
  jobName,
  designPath = NULL,
  dataPath = NULL,
  experimentObj = NULL,
  outputDir = ".",
  forceAllMethods = FALSE,
  omitLowAbundSamples = FALSE,
  sampleAbundThres = 5,
  tinyRunThres = 50,
```

```

    requireReplicates = TRUE,
    normalizeRetentionTime = TRUE,
    plotRows = 3,
    plotCols = 4,
    zeroToNA = FALSE,
    sampleColName = "sample",
    groupColName = "group",
    inputFormat = "default",
    skipAnalysis = FALSE,
    quiet = FALSE,
    noLogTransform = FALSE,
    writeReportAsPngs = FALSE,
    rtStepSizeMinutes = 1,
    rtWindowMinCount = 100,
    rtWindowShifts = 1,
    rtWindowMergeMethod = "mean"
)

```

## Arguments

|                        |                                                                                                                                                                              |
|------------------------|------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| jobName                | Give the current run a name.                                                                                                                                                 |
| designPath             | Path to file containing design matrix.                                                                                                                                       |
| dataPath               | Specify an output directory for generated files. Defaults to current working directory.                                                                                      |
| experimentObj          | SummarizedExperiment object, can be provided as input as alternative to 'designPath' and 'dataPath'                                                                          |
| outputDir              | Directory where results folder is created.                                                                                                                                   |
| forceAllMethods        | Debugging function. Run all normalizations even if they aren't in the recommended range of number of values                                                                  |
| omitLowAbundSamples    | Automatically remove samples with fewer non-NA values compared to threshold given by sampleAbundThres. Will otherwise stop with error message if such sample is encountered. |
| sampleAbundThres       | Threshold for omitting low-abundant samples. Is by default set to 15.                                                                                                        |
| tinyRunThres           | If total number of features is less than this, a limited run is performed.                                                                                                   |
| requireReplicates      | Require multiple samples per condition to pass input validation.                                                                                                             |
| normalizeRetentionTime | Perform normalizations over retention time.                                                                                                                                  |
| plotRows               | Number of plot-rows in output documentation.                                                                                                                                 |
| plotCols               | Number of plot-columns in output documentation.                                                                                                                              |
| zeroToNA               | Convert zero values to NA.                                                                                                                                                   |
| sampleColName          | Column name in design matrix containing sample IDs.                                                                                                                          |
| groupColName           | Column name in design matrix containing condition IDs.                                                                                                                       |
| inputFormat            | Type of input format.                                                                                                                                                        |
| skipAnalysis           | Only perform normalization steps.                                                                                                                                            |

**quiet** Omit status messages printed during run.  
**noLogTransform** Don't log-transform the input.  
**writeReportAsPngs** Output the evaluation report as PNG files instead of a single PDF  
**rtStepSizeMinutes** Retention time normalization window size.  
**rtWindowMinCount** Minimum number of datapoints in each retention-time segment.  
**rtWindowShifts** Number of layered retention time normalized windows.  
**rtWindowMergeMethod** Merge approach for layered retention time windows.

## Details

1: Loads the data matrix containing expression values and optional annotations, as well as the design matrix containing the experimental setup 2: Performs input data verification to validate that the data is in correct format. This step captures many common formatting errors. It returns an instance of the `NormalyzerDataset` class representing the unprocessed data. 3: Calculate a range of normalizations for the dataset. The result is provided as a `NormalyzerResults` object containing the resulting data matrices from each normalization. 4: Analyze the normalizations and generate performance measures for each of the normalized datasets. This result is provided as a `NormalyzerEvaluationResults` object. 5: Output the matrices containing the normalized datasets to files. 6: Generate visualizations overviewing the performance measures and write them to a PDF report.

## Value

None

## Examples

```
## Not run:
data_path <- system.file(package="NormalyzerDE", "extdata", "tiny_data.tsv")
design_path <- system.file(package="NormalyzerDE", "extdata", "tiny_design.tsv")
out_dir <- tempdir()
normalyzer(
  jobName="my_jobname",
  designPath=design_path,
  dataPath=data_path,
  outputDir=out_dir)
normalyzer(
  "my_jobname",
  designMatrix="design.tsv",
  "data.tsv",
  outputDir="path/to/output",
  normalizeRetentionTime=TRUE,
  retentionTimeWindow=2)
normalyzer(
  "my_jobname",
  designMatrix="design.tsv",
  "data.tsv",
  outputDir="path/to/output",
  inputFormat="maxquantprot")

## End(Not run)
```

---

|                   |                                                                             |
|-------------------|-----------------------------------------------------------------------------|
| NormalyzerDataset | <i>Represents raw input data together with basic annotation information</i> |
|-------------------|-----------------------------------------------------------------------------|

---

## Description

Takes a job name, a data matrix, a design matrix as well as specification of the group and sample columns in the design matrix. Provides the basic representation of a dataset in the NormalyzerDE normalization part.

## Usage

```
NormalyzerDataset(
  jobName,
  designMatrix,
  rawData,
  annotationData,
  sampleNameCol,
  groupNameCol,
  tinyRunThres = 50,
  quiet = FALSE
)
```

```
NormalyzerDataset(
  jobName,
  designMatrix,
  rawData,
  annotationData,
  sampleNameCol,
  groupNameCol,
  tinyRunThres = 50,
  quiet = FALSE
)
```

## Arguments

|                |                                                                                                                                                                    |
|----------------|--------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| jobName        | Name of the NormalyzerDE processing run                                                                                                                            |
| designMatrix   | Matrix containing sample conditions                                                                                                                                |
| rawData        | Matrix containing raw input data                                                                                                                                   |
| annotationData | Matrix containing annotation information for each input feature. Is expected to contain the same number of rows as the data but can contain any number of features |
| sampleNameCol  | Name of column in design matrix containing sample information                                                                                                      |
| groupNameCol   | Name of column in design matrix containing condition information                                                                                                   |
| tinyRunThres   | If fewer features than this is present in the input a limited run will be performed to avoid some steps requiring a more extensive number of features.             |
| quiet          | If set to TRUE no information messages will be printed                                                                                                             |

## Value

nds Generated NormalyzerDataset instance

**Slots**

jobName Name of the job represented by the dataset.  
 rawData Matrix with raw values.  
 sampleNameCol Name column for sample.  
 groupNameCol Name column for groups.  
 designMatrix Data frame containing design.  
 sampleNames Vector containing sample names.  
 filterrawdata Reduced raw data matrix where low abundance rows are removed  
 sampleReplicateGroups Vector with sample replicate information  
 samplesGroupsWithReplicates Vector with replicated sample replicate information  
 annotationValues Annotation part of original dataframe.  
 retentionTimes Vector of retention time values.  
 singleReplicateRun Conditional whether run is single replicate.

---

 normalyzerDE

*NormalyzerDE differential expression*


---

**Description**

Performs differential expression analysis on a normalization matrix. This command executes a pipeline processing the data and generates an annotated normalization matrix and a report containing p-value histograms for each of the performed comparisons.

**Usage**

```

normalyzerDE(
  jobName,
  comparisons,
  designPath = NULL,
  dataPath = NULL,
  experimentObj = NULL,
  outputDir = ".",
  logTrans = FALSE,
  type = "limma",
  sampleCol = "sample",
  condCol = "group",
  batchCol = NULL,
  techRepCol = NULL,
  leastRepCount = 1,
  quiet = FALSE,
  sigThres = 0.1,
  sigThresType = "fdr",
  log2FoldThres = 0,
  writeReportAsPngs = FALSE
)

```



**Arguments**

|                   |                                                                                                                                                                                 |
|-------------------|---------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| jobName           | Name of job                                                                                                                                                                     |
| comparisons       | Character vector containing target contrasts. If comparing condA with condB, then the vector would be c("condA-condB")                                                          |
| designPath        | File path to design matrix                                                                                                                                                      |
| dataPath          | File path to normalized matrix                                                                                                                                                  |
| experimentObj     | SummarizedExperiment object, can be provided as input as alternative to 'designPath' and 'dataPath'                                                                             |
| outputDir         | Path to output directory                                                                                                                                                        |
| logTrans          | Log transform the input (needed if providing non-logged input)                                                                                                                  |
| type              | Type of statistical comparison, "limma", "limma_intensity" or "welch", where "limma_intensity" allows the prior to be fit according to intensity rather than using a flat prior |
| sampleCol         | Design matrix column header for column containing sample IDs                                                                                                                    |
| condCol           | Design matrix column header for column containing sample conditions                                                                                                             |
| batchCol          | Provide an optional column for inclusion of possible batch variance in the model                                                                                                |
| techRepCol        | Design matrix column header for column containing technical replicates                                                                                                          |
| leastRepCount     | Minimum required replicate count                                                                                                                                                |
| quiet             | Omit status messages printed during run                                                                                                                                         |
| sigThres          | Significance threshold use for illustrating significant hits in diagnostic plots                                                                                                |
| sigThresType      | Type of significance threshold, "fdr" or "p". "fdr" is strongly recommended (Benjamini-Hochberg corrected p-values)                                                             |
| log2FoldThres     | Fold-size cutoff for being considered significant in diagnostic plots                                                                                                           |
| writeReportAsPngs | Output report as separate PNG files instead of a single PDF                                                                                                                     |

**Details**

When executed, it performs the following steps:

1: Read the data and the design matrices into dataframes. 2: Generate an instance of the NormalyzerStatistics class representing the data and their statistical comparisons. 3: Optionally reduce technical replicates in both the data matrix and the design matrix 4: Calculate statistical contrasts between supplied groups 5: Generate an annotated version of the original dataframe where columns containing statistical key measures have been added 6: Write the table to file 7: Generate a PDF report displaying p-value histograms for each calculated contrast

**Value**

None

**Examples**

```
data_path <- system.file(package="NormalyzerDE", "extdata", "tiny_data.tsv")
design_path <- system.file(package="NormalyzerDE", "extdata", "tiny_design.tsv")
out_dir <- tempdir()
normalyzerDE(
  jobName="my_jobname",
  comparisons=c("4-5"),
```

```

designPath=design_path,
dataPath=data_path,
outputDir=out_dir,
condCol="group")

```

---

## NormalyzerEvaluationResults

*Representation of evaluation results by calculating performance measures for an an NormalyzerResults instance*

---

### Description

Contains the resulting information from the processing which subsequently can be used to generate the quality assessment report.

### Usage

```
NormalyzerEvaluationResults(nr)
```

```
NormalyzerEvaluationResults(nr)
```

### Arguments

nr                      NormalyzerResults object

### Value

nds Generated NormalyzerEvaluationResults instance

### Slots

avgcvmem   Average coefficient of variance per method

avgcvmempdiff   Percentage difference of mean coefficient of variance compared to log2-transformed data

featureCVPerMethod   CV calculated per feature and normalization method.

avgmadmem   Average median absolute deviation

avgmadmempdiff   Percentage difference of median absolute deviation compared to log2-transformed data

avgvarmem   Average variance per method

avgvarmempdiff   Percentage difference of mean variance compared to log2-transformed data

lowVarFeaturesCVs   List of 5 for log2-transformed data

lowVarFeaturesCVsPercDiff   Coefficient of variance for least variable entries

anovaP   ANOVA calculated p-values

repCorPear   Within group Pearson correlations

repCorSpear   Within group Spearman correlations

**Examples**

```
data(example_summarized_experiment)
normObj <- getVerifiedNormalizerObject("job_name", example_summarized_experiment)
normResults <- normMethods(normObj)
normEval <- NormalizerEvaluationResults(normResults)
```

---

|                   |                                                                                   |
|-------------------|-----------------------------------------------------------------------------------|
| NormalizerResults | <i>Representation of the results from performing normalization over a dataset</i> |
|-------------------|-----------------------------------------------------------------------------------|

---

**Description**

It is linked to a NormalizerDataset instance representing the raw data which has been processed. After performing evaluation it also links to an instance of NormalizerEvaluationResults representing the results from the evaluation.

**Usage**

```
NormalizerResults(nds)
```

```
NormalizerResults(nds)
```

**Arguments**

|     |                          |
|-----|--------------------------|
| nds | NormalizerDataset object |
|-----|--------------------------|

**Value**

nr Prepared NormalizerResults object

**Slots**

normalizations SummarizedExperiment object containing calculated normalization results

nds Normalizer dataset representing run data

ner Normalizer evaluation results for running extended normalizations

**Examples**

```
data(example_summarized_experiment)
normObj <- getVerifiedNormalizerObject("job_name", example_summarized_experiment)
emptyNormResults <- NormalizerResults(normObj)
```

---

|                      |                                                                                |
|----------------------|--------------------------------------------------------------------------------|
| NormalyzerStatistics | <i>Class representing a dataset for statistical processing in NormalyzerDE</i> |
|----------------------|--------------------------------------------------------------------------------|

---

### Description

Is initialized with an annotation matrix, a data matrix and a design data frame. This object can subsequently be processed to generate statistical values and in turn used to write a full matrix with additional statistical information as well as a graphical report of the comparisons.

### Usage

```
NormalyzerStatistics(experimentObj, logTrans = FALSE)
```

```
NormalyzerStatistics(experimentObj, logTrans = FALSE)
```

### Arguments

|               |                                                                                          |
|---------------|------------------------------------------------------------------------------------------|
| experimentObj | Instance of SummarizedExperiment containing matrix and design information as column data |
| logTrans      | Whether the input data should be log transformed                                         |

### Value

nds Generated NormalyzerStatistics instance

### Slots

|                   |                                                              |
|-------------------|--------------------------------------------------------------|
| annotMat          | Matrix containing annotation information                     |
| dataMat           | Matrix containing (normalized) expression data               |
| filteredDataMat   | Filtered matrix with low-count rows removed                  |
| designDf          | Data frame containing design conditions                      |
| filteringContrast | Vector showing which entries are filtered (due to low count) |
| pairwiseCompsP    | List with P-values for pairwise comparisons                  |
| pairwiseCompsFdr  | List with FDR-values for pairwise comparisons                |
| pairwiseCompsAve  | List with average expression values                          |
| pairwiseCompsFold | List with log2 fold-change values for pairwise comparisons   |
| contrasts         | Spot for saving vector of last used contrasts                |
| condCol           | Column containing last used conditions                       |
| batchCol          | Column containing last used batch conditions                 |

### Examples

```
data(example_stat_summarized_experiment)
nst <- NormalyzerStatistics(example_stat_summarized_experiment)
```

normMethods

*Perform normalizations on Normalyzer dataset***Description**

Perform normalizations on Normalyzer dataset

**Usage**

```
normMethods(
  nds,
  forceAll = FALSE,
  normalizeRetentionTime = TRUE,
  quiet = FALSE,
  rtStepSizeMinutes = 1,
  rtWindowMinCount = 100,
  rtWindowShifts = 1,
  rtWindowMergeMethod = "mean",
  noLogTransform = FALSE
)
```

**Arguments**

|                        |                                                                                                              |
|------------------------|--------------------------------------------------------------------------------------------------------------|
| nds                    | Normalyzer dataset object.                                                                                   |
| forceAll               | Force all methods to run despite not qualifying for thresholds.                                              |
| normalizeRetentionTime | Perform retention time based normalization methods.                                                          |
| quiet                  | Prevent diagnostic output                                                                                    |
| rtStepSizeMinutes      | Retention time normalization window size.                                                                    |
| rtWindowMinCount       | Minimum number of datapoints in each retention-time segment.                                                 |
| rtWindowShifts         | Number of layered retention time normalized windows.                                                         |
| rtWindowMergeMethod    | Merge approach for layered retention time windows.                                                           |
| noLogTransform         | Per default NormalyzerDE performs a log-transformation on the input data. If not needed, specify this option |

**Value**

Returns Normalyzer results object with performed analyzes assigned as attributes

**Examples**

```
data(example_summarized_experiment)
normObj <- getVerifiedNormalyzerObject("job_name", example_summarized_experiment)
normResults <- normMethods(normObj)
```

---

```
performCyclicLoessNormalization
```

*Cyclic Loess normalization*

---

**Description**

Log2 transformed data is normalized by Loess method using the function "normalizeCyclicLoess". Further information is available for the function "normalizeCyclicLoess" in the Limma package.

**Usage**

```
performCyclicLoessNormalization(rawMatrix, noLogTransform = FALSE)
```

**Arguments**

rawMatrix      Target matrix to be normalized  
noLogTransform Assumes no need for log transformation

**Value**

Normalized matrix

**Examples**

```
data(example_data_only_values_small)  
normMatrix <- performCyclicLoessNormalization(example_data_only_values)
```

---

```
performGlobalRLRNormalization
```

*Global linear regression normalization*

---

**Description**

Log2 transformed data is normalized by robust linear regression using the function "rlm" from the MASS package.

**Usage**

```
performGlobalRLRNormalization(rawMatrix, noLogTransform = FALSE)
```

**Arguments**

rawMatrix      Target matrix to be normalized  
noLogTransform Assumes no need for log transformation

**Value**

Normalized matrix

**Examples**

```
data(example_data_only_values_small)
normMatrix <- performGlobalRLRNormalization(example_data_only_values)
```

---

```
performNoNormalization
```

*Do no normalization (For debugging purposes)*

---

**Description**

Do no normalization (For debugging purposes)

**Usage**

```
performNoNormalization(rawMatrix)
```

**Arguments**

rawMatrix      Target matrix to be normalized

**Value**

Normalized matrix

---

```
performNormalizations    Main function for executing normalizations
```

---

**Description**

Main function for executing normalizations

**Usage**

```
performNormalizations(
  nr,
  forceAll = FALSE,
  rtNorm = FALSE,
  rtStepSizeMinutes = 1,
  rtWindowMinCount = 100,
  rtWindowShifts = 1,
  rtWindowMergeMethod = "median",
  noLogTransform = FALSE,
  quiet = FALSE
)

## S4 method for signature 'NormalyzerResults'
performNormalizations(
  nr,
  forceAll = FALSE,
```

```

rtNorm = FALSE,
rtStepSizeMinutes = 1,
rtWindowMinCount = 100,
rtWindowShifts = 1,
rtWindowMergeMethod = "median",
noLogTransform = FALSE,
quiet = FALSE
)

```

### Arguments

|                     |                                                                                         |
|---------------------|-----------------------------------------------------------------------------------------|
| nr                  | Normalyzer results object.                                                              |
| forceAll            | Ignore dataset size limits and run all normalizations (only meant for testing purposes) |
| rtNorm              | Perform retention time based normalizations                                             |
| rtStepSizeMinutes   | Retention time normalization window size.                                               |
| rtWindowMinCount    | Minimum number of datapoints in each retention-time segment.                            |
| rtWindowShifts      | Number of layered retention time normalized windows.                                    |
| rtWindowMergeMethod | Merge approach for layered retention time windows.                                      |
| noLogTransform      | Prevent log-transforming input                                                          |
| quiet               | Don't show regular output messages                                                      |

### Value

nr NormalyzerDE results object

---

performQuantileNormalization

*Quantile normalization is performed by the function "normalize.quantiles" from the package preprocessCore.*

---

### Description

It makes the assumption that the data in different samples should originate from an identical distribution. It does this by generating a reference distribution and then scaling the other samples accordingly.

### Usage

```
performQuantileNormalization(rawMatrix, noLogTransform = FALSE)
```

### Arguments

|                |                                        |
|----------------|----------------------------------------|
| rawMatrix      | Target matrix to be normalized         |
| noLogTransform | Assumes no need for log transformation |



**Value**

Normalized matrix

**Examples**

```
data(example_data_only_values_small)
normMatrix <- performQuantileNormalization(example_data_only_values)
```

---

```
performSMADNormalization
```

*Median absolute deviation normalization Normalization subtracts the median and divides the data by the median absolute deviation (MAD).*

---

**Description**

Median absolute deviation normalization Normalization subtracts the median and divides the data by the median absolute deviation (MAD).

**Usage**

```
performSMADNormalization(rawMatrix, noLogTransform = FALSE)
```

**Arguments**

rawMatrix      Target matrix to be normalized  
noLogTransform Assumes no need for log transformation

**Value**

Normalized matrix

**Examples**

```
data(example_data_only_values_small)
normMatrix <- performSMADNormalization(example_data_only_values)
```

---

```
performVSNNormalization
```

*Log2 transformed data is normalized using the function "justvsnn" from the VSN package.*

---

**Description**

The VSN (Variance Stabilizing Normalization) attempts to transform the data in such a way that the variance remains nearly constant over the intensity spectrum

**Usage**

```
performVSNNormalization(rawMatrix)
```

**Arguments**

rawMatrix            Target matrix to be normalized

**Value**

Normalized matrix

**Examples**

```
data(example_data_only_values_small)
normMatrix <- performVSNNormalization(example_data_only_values)
```

---

|             |                                                                               |
|-------------|-------------------------------------------------------------------------------|
| plotBoxPlot | <i>Boxplots showing distribution of values after different normalizations</i> |
|-------------|-------------------------------------------------------------------------------|

---

**Description**

Boxplots showing distribution of values after different normalizations

**Usage**

```
plotBoxPlot(nr, currentLayout, pageno)
```

**Arguments**

nr                    Normalyzer results object.  
currentLayout        Layout used for document.  
pageno                Current page number.

**Value**

None

---

|                     |                                                                |
|---------------------|----------------------------------------------------------------|
| plotComparisonVenns | <i>If multiple comparisons - Show overlap in Venn diagrams</i> |
|---------------------|----------------------------------------------------------------|

---

**Description**

If multiple comparisons - Show overlap in Venn diagrams

**Usage**

```
plotComparisonVenns(
  nst,
  jobName,
  currentLayout,
  pageno,
  sigThres = 0.1,
  sigThresType = "fdr",
  log2FoldThres = 0,
  maxContrasts = 4
)
```

**Arguments**

|               |                                                    |
|---------------|----------------------------------------------------|
| nst           | NormalyzerDE statistics object.                    |
| jobName       | Name of processing run.                            |
| currentLayout | Layout used for document.                          |
| pageno        | Current page number.                               |
| sigThres      | Cutoff value for significance threshold            |
| sigThresType  | Type of significance cutoff                        |
| log2FoldThres | Log2-fold based cutoff threshold                   |
| maxContrasts  | Maximum contrasts to show pairwise comparisons for |

**Value**

None

---

|                 |                                                                                                                                                                                             |
|-----------------|---------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| plotContrastPCA | <i>Show in a PCA plot what samples are compared in statistical contrast<br/>This is useful to understand what conditions are compared and for<br/>checking for outliers in the contrast</i> |
|-----------------|---------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|

---

**Description**

Show in a PCA plot what samples are compared in statistical contrast This is useful to understand what conditions are compared and for checking for outliers in the contrast

**Usage**

```
plotContrastPCA(nst, jobName, currentLayout, pageno, pcs = c(1, 2))
```

**Arguments**

|               |                                 |
|---------------|---------------------------------|
| nst           | NormalyzerDE statistics object. |
| jobName       | Name of processing run.         |
| currentLayout | Layout used for document.       |
| pageno        | Current page number.            |
| pcs           | Principal components to show.   |

**Value**

None

---

|                    |                                                                                                                       |
|--------------------|-----------------------------------------------------------------------------------------------------------------------|
| plotContrastPHists | <i>Takes an NormalyzerStatistics instance and generates and prints a p-value histogram for each onto the viewport</i> |
|--------------------|-----------------------------------------------------------------------------------------------------------------------|

---

**Description**

Takes an NormalyzerStatistics instance and generates and prints a p-value histogram for each onto the viewport

**Usage**

```
plotContrastPHists(nst, jobName, currentLayout, pageno)
```

**Arguments**

|               |                                 |
|---------------|---------------------------------|
| nst           | NormalyzerDE statistics object. |
| jobName       | Name of processing run.         |
| currentLayout | Layout used for document.       |
| pageno        | Current page number.            |

**Value**

None

---

|                 |                                                 |
|-----------------|-------------------------------------------------|
| plotCorrelation | <i>Visualize within-replicates correlations</i> |
|-----------------|-------------------------------------------------|

---

**Description**

Visualize within-replicates correlations

**Usage**

```
plotCorrelation(nr, currentLayout, pageno)
```

**Arguments**

|               |                            |
|---------------|----------------------------|
| nr            | Normalyzer results object. |
| currentLayout | Layout used for document.  |
| pageno        | Current page number.       |

**Value**

None

---

|                   |                                                                                                                                  |
|-------------------|----------------------------------------------------------------------------------------------------------------------------------|
| plotCVvsIntensity | <i>Plots page displaying coefficient of variance (CV) against raw intensity for features across the performed normalizations</i> |
|-------------------|----------------------------------------------------------------------------------------------------------------------------------|

---

**Description**

Plots page displaying coefficient of variance (CV) against raw intensity for features across the performed normalizations

**Usage**

```
plotCVvsIntensity(nr, currentLayout, pageno)
```

**Arguments**

|               |                            |
|---------------|----------------------------|
| nr            | Normalyzer results object. |
| currentLayout | Layout used for document.  |
| pageno        | Current page number.       |

**Value**

None

---

|                 |                                                 |
|-----------------|-------------------------------------------------|
| plotDendrograms | <i>Visualize dendrogram grouping of samples</i> |
|-----------------|-------------------------------------------------|

---

**Description**

Visualize dendrogram grouping of samples

**Usage**

```
plotDendrograms(nr, currentLayout, pageno)
```

**Arguments**

|               |                            |
|---------------|----------------------------|
| nr            | Normalyzer results object. |
| currentLayout | Layout used for document.  |
| pageno        | Current page number.       |

**Value**

None

---

|             |                                                                       |
|-------------|-----------------------------------------------------------------------|
| plotDensity | <i>Density plots showing value distributions after normalizations</i> |
|-------------|-----------------------------------------------------------------------|

---

**Description**

Density plots showing value distributions after normalizations

**Usage**

```
plotDensity(nr, currentLayout, pageno)
```

**Arguments**

|               |                            |
|---------------|----------------------------|
| nr            | Normalyzer results object. |
| currentLayout | Layout used for document.  |
| pageno        | Current page number.       |

**Value**

None

---

|               |                                                                   |
|---------------|-------------------------------------------------------------------|
| plotFrontPage | <i>Generate first page in output report and write to viewport</i> |
|---------------|-------------------------------------------------------------------|

---

**Description**

Generate first page in output report and write to viewport

**Usage**

```
plotFrontPage(currentjob, currentFont)
```

**Arguments**

|             |                                |
|-------------|--------------------------------|
| currentjob  | Name of current run.           |
| currentFont | Font used for output document. |

**Value**

None

---

|        |                                                                                                                                                                                                             |
|--------|-------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| plotMA | <i>Produces a page containing expression vs. fold-change figures (MA plots) The visualized fold is between the first sample in each group and the average of the replicate to which that sample belongs</i> |
|--------|-------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|

---

**Description**

Produces a page containing expression vs. fold-change figures (MA plots) The visualized fold is between the first sample in each group and the average of the replicate to which that sample belongs

**Usage**

```
plotMA(nr, currentLayout, pageno)
```

**Arguments**

|               |                            |
|---------------|----------------------------|
| nr            | Normalyzer results object. |
| currentLayout | Layout used for document.  |
| pageno        | Current page number.       |

**Value**

None

---

|         |                                                                   |
|---------|-------------------------------------------------------------------|
| plotMDS | <i>MDS plots showing grouping of samples after normalizations</i> |
|---------|-------------------------------------------------------------------|

---

**Description**

MDS plots showing grouping of samples after normalizations

**Usage**

```
plotMDS(nr, currentLayout, pageno)
```

**Arguments**

|               |                            |
|---------------|----------------------------|
| nr            | Normalyzer results object. |
| currentLayout | Layout used for document.  |
| pageno        | Current page number.       |

**Value**

None

---

|            |                                                                             |
|------------|-----------------------------------------------------------------------------|
| plotMeanSD | <i>Visualize standard deviation over (expression?) for different values</i> |
|------------|-----------------------------------------------------------------------------|

---

**Description**

Visualize standard deviation over (expression?) for different values

**Usage**

```
plotMeanSD(nr, currentLayout, pageno)
```

**Arguments**

|               |                            |
|---------------|----------------------------|
| nr            | Normalyzer results object. |
| currentLayout | Layout used for document.  |
| pageno        | Current page number.       |

**Value**

None

---

|           |                                                                            |
|-----------|----------------------------------------------------------------------------|
| plotPHist | <i>Generate P-histograms for ANOVA calculated after each normalization</i> |
|-----------|----------------------------------------------------------------------------|

---

**Description**

Generate P-histograms for ANOVA calculated after each normalization

**Usage**

```
plotPHist(nr, currentLayout, pageno)
```

**Arguments**

|               |                            |
|---------------|----------------------------|
| nr            | Normalyzer results object. |
| currentLayout | Layout used for document.  |
| pageno        | Current page number.       |

**Value**

None



---

|        |                                                                                                                                                                           |
|--------|---------------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| plotQQ | <i>Produces page showing QQ-plots for the first sample for each normalization method. This plot can be used to assess whether the data follows a normal distribution.</i> |
|--------|---------------------------------------------------------------------------------------------------------------------------------------------------------------------------|

---

### Description

Produces page showing QQ-plots for the first sample for each normalization method. This plot can be used to assess whether the data follows a normal distribution.

### Usage

```
plotQQ(nr, currentLayout, pageno)
```

### Arguments

|               |                            |
|---------------|----------------------------|
| nr            | Normalyzer results object. |
| currentLayout | Layout used for document.  |
| pageno        | Current page number.       |

### Value

None

---

|                                    |                                                                                                                                                                                                                                                                                                                                                                                                      |
|------------------------------------|------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| plotReplicateVarAndStableVariables | <i>Write figures displaying pooled coefficient of variance, median absolute deviation and pooled estimate of variance percentage compared to log2-transformed and stable variables plot displaying CV of stable variables against pooled CV measure. The stable variables are calculated by an ANOVA comparison across sample conditions and selecting features with the least clear difference.</i> |
|------------------------------------|------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|

---

### Description

Write figures displaying pooled coefficient of variance, median absolute deviation and pooled estimate of variance percentage compared to log2-transformed and stable variables plot displaying CV of stable variables against pooled CV measure. The stable variables are calculated by an ANOVA comparison across sample conditions and selecting features with the least clear difference.

### Usage

```
plotReplicateVarAndStableVariables(nr, currentLayout, pageno)
```

### Arguments

|               |                            |
|---------------|----------------------------|
| nr            | Normalyzer results object. |
| currentLayout | Layout used for document.  |
| pageno        | Current page number.       |

**Value**

None

---

|                       |                                                                                                                                                                                                                      |
|-----------------------|----------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| plotReplicateVariance | <i>Generate normalization replicate variance summary by displaying CV (coefficient of variance), MAD (mean of intragroup median absolute deviation) and PEV (Pooled Estimate of Variance) as mean of intragroups</i> |
|-----------------------|----------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|

---

**Description**

Generate normalization replicate variance summary by displaying CV (coefficient of variance), MAD (mean of intragroup median absolute deviation) and PEV (Pooled Estimate of Variance) as mean of intragroups

**Usage**

```
plotReplicateVariance(nr, currentLayout, pageno)
```

**Arguments**

|               |                            |
|---------------|----------------------------|
| nr            | Normalyzer results object. |
| currentLayout | Layout used for document.  |
| pageno        | Current page number.       |

**Value**

None

---

|         |                                                                      |
|---------|----------------------------------------------------------------------|
| plotRLE | <i>Boxplots showing relative log expression after normalizations</i> |
|---------|----------------------------------------------------------------------|

---

**Description**

Boxplots showing relative log expression after normalizations

**Usage**

```
plotRLE(nr, currentLayout, pageno)
```

**Arguments**

|               |                            |
|---------------|----------------------------|
| nr            | Normalyzer results object. |
| currentLayout | Layout used for document.  |
| pageno        | Current page number.       |

**Value**

None

---

plotSampleMappingPage *Write page with sample mapping*

---

**Description**

Write page with sample mapping

**Usage**

```
plotSampleMappingPage(nr, currentFont, currentLayout, currentjob, pageno)
```

**Arguments**

|               |                            |
|---------------|----------------------------|
| nr            | Normalyzer results object. |
| currentLayout | Layout used for document.  |
| pageno        | Current page number.       |

**Value**

None

---

plotSampleOutlierSummary  
*Write page containing sample summary of intensities, missing values  
and MDS plot to the viewport*

---

**Description**

Write page containing sample summary of intensities, missing values and MDS plot to the viewport

**Usage**

```
plotSampleOutlierSummary(nr, currentLayout, pageno)
```

**Arguments**

|               |                            |
|---------------|----------------------------|
| nr            | Normalyzer results object. |
| currentLayout | Layout used for document.  |
| pageno        | Current page number.       |

**Value**

None

---

|             |                                                                                                                                                |
|-------------|------------------------------------------------------------------------------------------------------------------------------------------------|
| plotScatter | <i>Produces page containing scatter plot plotting the first two samples from each dataset against each other for each normalization method</i> |
|-------------|------------------------------------------------------------------------------------------------------------------------------------------------|

---

### Description

Produces page containing scatter plot plotting the first two samples from each dataset against each other for each normalization method

### Usage

```
plotScatter(nr, currentLayout, pageno)
```

### Arguments

|               |                            |
|---------------|----------------------------|
| nr            | Normalyzer results object. |
| currentLayout | Layout used for document.  |
| pageno        | Current page number.       |

### Value

None

---

|                |                                                                                       |
|----------------|---------------------------------------------------------------------------------------|
| plotSigScatter | <i>Takes an NormalyzerStatistics instance and generates and prints a volcano plot</i> |
|----------------|---------------------------------------------------------------------------------------|

---

### Description

Takes an NormalyzerStatistics instance and generates and prints a volcano plot

### Usage

```
plotSigScatter(
  nst,
  jobName,
  currentLayout,
  pageno,
  type = "Volcano",
  sigThres = 0.1,
  sigThresType = "fdr",
  log2FoldThres = 0
)
```

**Arguments**

|               |                                            |
|---------------|--------------------------------------------|
| nst           | NormalyzerDE statistics object.            |
| jobName       | Name of processing run.                    |
| currentLayout | Layout used for document.                  |
| pageno        | Current page number.                       |
| type          | Specify whether to plot 'Volcano' or 'MA'. |
| sigThres      | FDR threshold for DE coloring.             |

**Value**

None

---

|                |                                                                      |
|----------------|----------------------------------------------------------------------|
| preprocessData | <i>Replace empty values (0 or empty field) with NA in input data</i> |
|----------------|----------------------------------------------------------------------|

---

**Description**

Replace empty values (0 or empty field) with NA in input data

**Usage**

```
preprocessData(dataMatrix, quiet = FALSE)
```

**Arguments**

|            |                                |
|------------|--------------------------------|
| dataMatrix | Matrix with raw data.          |
| quiet      | Don't show diagnostic messages |

**Value**

Parsed rawdata where 0 values are replaced with NA

---

|           |                                                                                                                                               |
|-----------|-----------------------------------------------------------------------------------------------------------------------------------------------|
| printMeta | <i>Print meta information for Normalyzer plot page ! Needs refactoring to reduce redundancy in code ! Needs double check of functionality</i> |
|-----------|-----------------------------------------------------------------------------------------------------------------------------------------------|

---

**Description**

Print meta information for Normalyzer plot page ! Needs refactoring to reduce redundancy in code ! Needs double check of functionality

**Usage**

```
printMeta(plotname, pageno, jobname, currentLayout)
```

**Arguments**

|               |                         |
|---------------|-------------------------|
| plotname      | Name of current plot.   |
| pageno        | Current page number.    |
| jobname       | Name of ongoing job.    |
| currentLayout | Custom viewport layout. |

**Value**

None

---

|            |                                                                      |
|------------|----------------------------------------------------------------------|
| printPlots | <i>Generate PDF grid page filling it with provided list of plots</i> |
|------------|----------------------------------------------------------------------|

---

**Description**

Generate PDF grid page filling it with provided list of plots

**Usage**

```
printPlots(plotlist, plotname, pageno, jobname, currentLayout)
```

**Arguments**

|               |                                                        |
|---------------|--------------------------------------------------------|
| plotlist      | List of target plots to display.                       |
| plotname      | List of names corresponding to the provided plot list. |
| pageno        | Current page number.                                   |
| jobname       | Name of ongoing job.                                   |
| currentLayout | Custom viewport layout.                                |

**Value**

None

---

|                           |                                                         |
|---------------------------|---------------------------------------------------------|
| reduceTechnicalReplicates | <i>Remove technical replicates from data and design</i> |
|---------------------------|---------------------------------------------------------|

---

**Description**

Collapses sample values into their average. If only one value is present due to NA-values in other technical replicates, then that value is used.

**Usage**

```
reduceTechnicalReplicates(se, techRepColName, sampleColName)
```

**Arguments**

|                |                                                                                                       |
|----------------|-------------------------------------------------------------------------------------------------------|
| se             | Summarized experiment where the assay contains the data to be reduced, and the colData the data frame |
| techRepColName | Technical replicates column name in colData                                                           |
| sampleColName  | Sample names column name in colData                                                                   |

**Details**

Takes a SummarizedExperiment where the data is present as the assay and the colData contains the design conditions. In the design conditions there should be one column with the technical replicate groups and one column containing the sample names

**Value**

reducedSe Summarized experiment with reduced data

**Examples**

```
testData <- as.matrix(data.frame(
  c(1,1,1),
  c(1,2,1),
  c(7,7,7),
  c(7,9,7)))
colnames(testData) <- c("a1", "a2", "b1", "b2")
designDf <- data.frame(
  sample=c("a1", "a2", "b1", "b2"),
  techrep=c("a", "a", "b", "b"))
se <- SummarizedExperiment::SummarizedExperiment(
  assay=testData,
  colData=designDf
)
statObj <- reduceTechnicalReplicates(se, "techrep", "sample")
```

---

|             |                                       |
|-------------|---------------------------------------|
| setupJobDir | <i>Create empty directory for run</i> |
|-------------|---------------------------------------|

---

**Description**

Creates a directory at provided path named to the jobname.

**Usage**

```
setupJobDir(jobName, outputDir)
```

**Arguments**

|           |                                                         |
|-----------|---------------------------------------------------------|
| jobName   | Name of the run.                                        |
| outputDir | Path to directory where to create the output directory. |

**Value**

Path to newly created directory.

**Examples**

```
setupJobDir("job_name", "path/to/outdir")
```

---

|               |                                                                                                                        |
|---------------|------------------------------------------------------------------------------------------------------------------------|
| setupPlotting | <i>Setup PDF report settings by initializing the color palette, format for the PDF report and the graphical device</i> |
|---------------|------------------------------------------------------------------------------------------------------------------------|

---

**Description**

Setup PDF report settings by initializing the color palette, format for the PDF report and the graphical device

**Usage**

```
setupPlotting(currentJob, jobDir, suffix)
```

**Arguments**

|            |                                   |
|------------|-----------------------------------|
| currentJob | Name of current run.              |
| jobDir     | Path to output directory for run. |
| suffix     | Text to add to output filename.   |

**Value**

None

---

|                        |                                                                |
|------------------------|----------------------------------------------------------------|
| setupRawContrastObject | <i>Prepare SummarizedExperiment object for statistics data</i> |
|------------------------|----------------------------------------------------------------|

---

**Description**

Prepare SummarizedExperiment object for statistics data

**Usage**

```
setupRawContrastObject(dataPath, designPath, sampleColName)
```

**Arguments**

|               |                                                          |
|---------------|----------------------------------------------------------|
| dataPath      | Path to raw data matrix                                  |
| designPath    | Path to design matrix                                    |
| sampleColName | Name for column in design matrix containing sample names |

**Value**

experimentObj Prepared instance of SummarizedExperiment



**Examples**

```
data_path <- system.file(package="NormalyzerDE", "extdata", "tiny_data.tsv")
design_path <- system.file(package="NormalyzerDE", "extdata", "tiny_design.tsv")
sumExpObj <- setupRawContrastObject(data_path, design_path, "sample")
```

---

|                    |                                                                                                                             |
|--------------------|-----------------------------------------------------------------------------------------------------------------------------|
| setupRawDataObject | <i>Prepare SummarizedExperiment object for raw data to be normalized containing data, design and annotation information</i> |
|--------------------|-----------------------------------------------------------------------------------------------------------------------------|

---

**Description**

Prepare SummarizedExperiment object for raw data to be normalized containing data, design and annotation information

**Usage**

```
setupRawDataObject(
  dataPath,
  designPath,
  inputFormat = "default",
  zeroToNA = FALSE,
  sampleColName = "sample",
  groupColName = "group"
)
```

**Arguments**

|               |                                                                                               |
|---------------|-----------------------------------------------------------------------------------------------|
| dataPath      | File path to data matrix.                                                                     |
| designPath    | File path to design matrix.                                                                   |
| inputFormat   | Type of matrix for data, can be either 'default', 'proteios', 'maxquantprot' or 'maxquantpep' |
| zeroToNA      | If TRUE zeroes in the data is automatically converted to NA values                            |
| sampleColName | Column name for column containing sample names                                                |
| groupColName  | Column name for column containing condition levels                                            |

**Value**

experimentObj SummarizedExperiment object loaded with the data

**Examples**

```
data_path <- system.file(package="NormalyzerDE", "extdata", "tiny_data.tsv")
design_path <- system.file(package="NormalyzerDE", "extdata", "tiny_design.tsv")
df <- setupRawDataObject(data_path, design_path)
```

---

|               |                                                                                        |
|---------------|----------------------------------------------------------------------------------------|
| setupTestData | <i>Generate a random test dataset with features, sample values and retention times</i> |
|---------------|----------------------------------------------------------------------------------------|

---

**Description**

Generate a random test dataset with features, sample values and retention times

**Usage**

```
setupTestData(nSamples, nFeatures, rtMin = 40, rtMax = 80, mean = 20, sd = 4)
```

**Arguments**

|           |                                           |
|-----------|-------------------------------------------|
| nSamples  | Number of samples                         |
| nFeatures | Number of features                        |
| rtMin     | Minimum retention time                    |
| rtMax     | Maximum retention time                    |
| mean      | Mean value for sample intensities         |
| sd        | Standard deviation for sample intensities |

**Value**

Test dataset

**Examples**

```
df <- setupTestData(6, 20)
df <- setupTestData(6, 20, mean=15, sd=1)
```

---

|                           |                                                  |
|---------------------------|--------------------------------------------------|
| validateSampleReplication | <i>Check whether all samples have replicates</i> |
|---------------------------|--------------------------------------------------|

---

**Description**

Check whether all samples have replicates

**Usage**

```
validateSampleReplication(
  dataMatrix,
  groups,
  requireReplicates = TRUE,
  quiet = FALSE
)
```

**Arguments**

|                   |                                                                |
|-------------------|----------------------------------------------------------------|
| dataMatrix        | Prepared matrix containing expression data.                    |
| groups            | Vector containing condition levels                             |
| requireReplicates | By default stops processing if not all samples have replicates |

**Value**

None

---

|                 |                                                                                                                                                                                             |
|-----------------|---------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| verifyContrasts | <i>Check that a given contrast string is valid given a particular design matrix. Each level tested for in the contrast should be present in the condition column for the design matrix.</i> |
|-----------------|---------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|

---

**Description**

Mainly meant to verify strings received during server usage.

**Usage**

```
verifyContrasts(designLevels, contrasts)
```

**Arguments**

|              |                                                                                                      |
|--------------|------------------------------------------------------------------------------------------------------|
| designLevels | Vector containing condition levels present in design                                                 |
| contrasts    | A string containing one or several (comma delimited) strings for which contrasts should be performed |

**Value**

None

---

|                    |                                                                |
|--------------------|----------------------------------------------------------------|
| verifyDesignMatrix | <i>Verify that design matrix setup matches the data matrix</i> |
|--------------------|----------------------------------------------------------------|

---

**Description**

Verify that design matrix setup matches the data matrix

**Usage**

```
verifyDesignMatrix(fullMatrix, designMatrix, sampleCol)
```

**Arguments**

|              |                                                |
|--------------|------------------------------------------------|
| fullMatrix   | Dataframe with input data.                     |
| designMatrix | Dataframe with design setup.                   |
| sampleCol    | Column in design matrix containing sample IDs. |

**Value**

None

---

 verifyMultipleSamplesPresent

*Check whether more than one sample is present*


---

**Description**

Check whether more than one sample is present

**Usage**

```
verifyMultipleSamplesPresent(
  dataMatrix,
  groups,
  requireReplicates = TRUE,
  quiet = FALSE
)
```

**Arguments**

dataMatrix      Prepared dataframe.

groups          Vector containing condition levels

requireReplicates      By default stops processing if not all samples have replicates

**Value**

None

---

 verifySummarizedExperiment

*Verify that design matrix setup matches the data matrix*


---

**Description**

Verify that design matrix setup matches the data matrix

**Usage**

```
verifySummarizedExperiment(summarizedExp, sampleCol)
```

**Arguments**

sampleCol      Column in design matrix containing sample IDs.

fullMatrix      Dataframe with input data.

designMatrix      Dataframe with design setup.

**Value**

None

---

|                    |                                                                 |
|--------------------|-----------------------------------------------------------------|
| verifyValidNumbers | <i>Verify that input fields conform to the expected formats</i> |
|--------------------|-----------------------------------------------------------------|

---

**Description**

Verify that input fields conform to the expected formats

**Usage**

```
verifyValidNumbers(rawDataOnly, groups, noLogTransform = FALSE, quiet = FALSE)
```

**Arguments**

|             |                                   |
|-------------|-----------------------------------|
| rawDataOnly | Dataframe with input data.        |
| groups      | Condition levels for comparisons. |

**Value**

None

---

|                         |                                             |
|-------------------------|---------------------------------------------|
| writeNormalizedDatasets | <i>Write normalization matrices to file</i> |
|-------------------------|---------------------------------------------|

---

**Description**

Outputs each of the normalized datasets to the specified directory.

**Usage**

```
writeNormalizedDatasets(  
  nr,  
  jobdir,  
  includePairwiseComparisons = FALSE,  
  includeCvCol = FALSE,  
  includeAnovaP = FALSE,  
  normSuffix = "-normalized.txt",  
  rawDataName = "submitted_rawdata.txt"  
)
```

**Arguments**

|                                         |                                                               |
|-----------------------------------------|---------------------------------------------------------------|
| <code>nr</code>                         | Results object.                                               |
| <code>jobdir</code>                     | Path to output directory.                                     |
| <code>includePairwiseComparisons</code> | Include p-values for pairwise comparisons.                    |
| <code>includeCvCol</code>               | Include CV column in output.                                  |
| <code>includeAnovaP</code>              | Include ANOVA p-value in output.                              |
| <code>normSuffix</code>                 | String used to name output together with normalization names. |
| <code>rawdataName</code>                | Name of output raw data file.                                 |

**Value**

None

**Examples**

```
data(example_summarized_experiment)
normObj <- getVerifiedNormalizerObject("job_name", example_summarized_experiment)
normResults <- normMethods(normObj)
normResultsWithEval <- analyzeNormalizations(normResults)
outputDir <- tempdir()
writeNormalizedDatasets(normResultsWithEval, outputDir)
```

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