

# Package ‘PRONE’

April 25, 2025

**Type** Package

**Title** The PROteomics Normalization Evaluator

**Version** 1.3.0

**Description** High-throughput omics data are often affected by systematic biases introduced throughout all the steps of a clinical study, from sample collection to quantification. Normalization methods aim to adjust for these biases to make the actual biological signal more prominent. However, selecting an appropriate normalization method is challenging due to the wide range of available approaches. Therefore, a comparative evaluation of unnormalized and normalized data is essential in identifying an appropriate normalization strategy for a specific data set. This R package provides different functions for preprocessing, normalizing, and evaluating different normalization approaches. Furthermore, normalization methods can be evaluated on downstream steps, such as differential expression analysis and statistical enrichment analysis. Spike-in data sets with known ground truth and real-world data sets of biological experiments acquired by either tandem mass tag (TMT) or label-free quantification (LFQ) can be analyzed.

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**Imports** dplyr, magrittr, data.table, RColorBrewer, ggplot2, S4Vectors, ComplexHeatmap, stringr, NormalyzerDE, tibble, limma, MASS, edgeR, matrixStats, preprocessCore, stats, gtools, methods, ROTS, ComplexUpset, tidyr, purrr, circlize, gprofiler2, plotROC, MSnbase, UpSetR, dendsort, vsn, Biobase, reshape2, POMA, ggtext, scales, DEqMS, vegan

**Suggests** testthat (>= 3.0.0), knitr, rmarkdown, BiocStyle, DT

**BugReports** <https://github.com/daisybio/PRONE/issues>

**URL** <https://github.com/daisybio/PRONE>

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PRONE-package

*PRONE: The PROteomics Normalization Evaluator*

---

## Description

High-throughput omics data are often affected by systematic biases introduced throughout all the steps of a clinical study, from sample collection to quantification. Normalization methods aim to adjust for these biases to make the actual biological signal more prominent. However, selecting an appropriate normalization method is challenging due to the wide range of available approaches. Therefore, a comparative evaluation of unnormalized and normalized data is essential in identifying an appropriate normalization strategy for a specific data set. This R package provides different functions for preprocessing, normalizing, and evaluating different normalization approaches. Furthermore, normalization methods can be evaluated on downstream steps, such as differential expression analysis and statistical enrichment analysis. Spike-in data sets with known ground truth and real-world data sets of biological experiments acquired by either tandem mass tag (TMT) or label-free quantification (LFQ) can be analyzed.

## Author(s)

**Maintainer:** Lis Arend <lis.arend@tum.de> ([ORCID](#))

## See Also

Useful links:

- <https://github.com/daisybio/PRONE>
- Report bugs at <https://github.com/daisybio/PRONE/issues>

---

|                  |                                             |
|------------------|---------------------------------------------|
| apply_thresholds | <i>Apply other thresholds to DE results</i> |
|------------------|---------------------------------------------|

---

### Description

Apply other thresholds to DE results

### Usage

```
apply_thresholds(
  de_res,
  logFC = TRUE,
  logFC_up = 1,
  logFC_down = -1,
  p_adj = TRUE,
  alpha = 0.05
)
```

### Arguments

|            |                                                                                                        |
|------------|--------------------------------------------------------------------------------------------------------|
| de_res     | data table resulting of run_DE                                                                         |
| logFC      | Boolean specifying whether to apply a logFC threshold (TRUE) or not (FALSE)                            |
| logFC_up   | Upper log2 fold change threshold (dividing into up regulated)                                          |
| logFC_down | Lower log2 fold change threshold (dividing into down regulated)                                        |
| p_adj      | Boolean specifying whether to apply a threshold on adjusted p-values (TRUE) or on raw p-values (FALSE) |
| alpha      | Threshold for adjusted p-values or p-values                                                            |

### Value

data table updating the Change column with the newly applied thresholds

### Examples

```
data(tuberculosis_TMT_de_res)
de_res <- apply_thresholds(tuberculosis_TMT_de_res, logFC = FALSE,
  p_adj = TRUE, alpha = 0.01)
```

---

|                       |                                                                                                     |
|-----------------------|-----------------------------------------------------------------------------------------------------|
| check_DEqMS_parameter | <i>Helper function to check whether the DEqMS_PSMs_column is in the SummarizedExperiment object</i> |
|-----------------------|-----------------------------------------------------------------------------------------------------|

---

### Description

Helper function to check whether the DEqMS\_PSMs\_column is in the SummarizedExperiment object

**Usage**

```
check_DEqMS_parameter(se, DEqMS_PSMs_column)
```

**Arguments**

|                   |                                                                                      |
|-------------------|--------------------------------------------------------------------------------------|
| se                | SummarizedExperiment containing all necessary information of the proteomics data set |
| DEqMS_PSMs_column | String specifying which column name to use for DEqMS (default NULL)                  |

**Value**

None

---

|                     |                                         |
|---------------------|-----------------------------------------|
| check_DE_parameters | <i>Check parameters for DE analysis</i> |
|---------------------|-----------------------------------------|

---

**Description**

Check parameters for DE analysis

**Usage**

```
check_DE_parameters(
  se,
  ain = NULL,
  condition = NULL,
  comparisons = NULL,
  DE_method = "limma",
  covariate = NULL,
  logFC = TRUE,
  logFC_up = 1,
  logFC_down = -1,
  p_adj = TRUE,
  p_adj_method = "BH",
  alpha = 0.05,
  B = 100,
  K = 500,
  DEqMS_PSMs_column = NULL
)
```

**Arguments**

|           |                                                                                                                                                       |
|-----------|-------------------------------------------------------------------------------------------------------------------------------------------------------|
| se        | SummarizedExperiment containing all necessary information of the proteomics data set                                                                  |
| ain       | Vector of strings which assay should be used as input (default NULL). If NULL then all normalization of the se object are plotted next to each other. |
| condition | column name of condition (if NULL, condition saved in SummarizedExperiment will be taken)                                                             |

|                   |                                                                                                        |
|-------------------|--------------------------------------------------------------------------------------------------------|
| comparisons       | Vector of comparisons that are performed in the DE analysis (from specify_comparisons method)          |
| DE_method         | String specifying which DE method should be applied (limma, ROTS, DEqMS)                               |
| covariate         | String specifying which column to include as covariate into limma                                      |
| logFC             | Boolean specifying whether to apply a logFC threshold (TRUE) or not (FALSE)                            |
| logFC_up          | Upper log2 fold change threshold (dividing into up regulated)                                          |
| logFC_down        | Lower log2 fold change threshold (dividing into down regulated)                                        |
| p_adj             | Boolean specifying whether to apply a threshold on adjusted p-values (TRUE) or on raw p-values (FALSE) |
| p_adj_method      | String specifying the method for adjusted p-values                                                     |
| alpha             | Threshold for adjusted p-values or p-values                                                            |
| B                 | Number of bootstrapping for ROTS                                                                       |
| K                 | Number of top-ranked features for reproducibility optimization                                         |
| DEqMS_PSMs_column | String specifying which column name to use for DEqMS (default NULL)                                    |

**Value**

list of checked assays and condition column name

---

|                    |                                                                                             |
|--------------------|---------------------------------------------------------------------------------------------|
| check_input_assays | <i>Helper function to check whether all given assays are in SummarizedExperiment object</i> |
|--------------------|---------------------------------------------------------------------------------------------|

---

**Description**

Helper function to check whether all given assays are in SummarizedExperiment object

**Usage**

```
check_input_assays(se, ain)
```

**Arguments**

|     |                                                                                                                                                       |
|-----|-------------------------------------------------------------------------------------------------------------------------------------------------------|
| se  | SummarizedExperiment containing all necessary information of the proteomics data set                                                                  |
| ain | Vector of strings which assay should be used as input (default NULL). If NULL then all normalization of the se object are plotted next to each other. |

**Value**

NULL if no methods in SummarizedExperiment object, else all available methods ready for visualization

---

`check_plot_DE_parameters`*Helper function to check the parameters for plotting the DE results*

---

**Description**

Helper function to check the parameters for plotting the DE results

**Usage**

```
check_plot_DE_parameters(de_res, ain, comparisons)
```

**Arguments**

|                          |                                                                                                                  |
|--------------------------|------------------------------------------------------------------------------------------------------------------|
| <code>de_res</code>      | data table resulting of <code>run_DE</code>                                                                      |
| <code>ain</code>         | String of normalization methods to visualize (must be valid normalization methods saved in <code>de_res</code> ) |
| <code>comparisons</code> | Vector of comparisons (must be valid comparisons saved in <code>de_res</code> )                                  |

**Value**

list of valid inputs for plotting functions

---

`check_stats_spiked_DE_parameters`*Helper function to check the parameters for plotting the DE stats of spike-in data sets*

---

**Description**

Helper function to check the parameters for plotting the DE stats of spike-in data sets

**Usage**

```
check_stats_spiked_DE_parameters(stats, ain, comparisons)
```

**Arguments**

|                          |                                                                                                                  |
|--------------------------|------------------------------------------------------------------------------------------------------------------|
| <code>stats</code>       | data table resulting of <code>get_spiked_stats_DE</code>                                                         |
| <code>ain</code>         | String of normalization methods to visualize (must be valid normalization methods saved in <code>de_res</code> ) |
| <code>comparisons</code> | Vector of comparisons (must be valid comparisons saved in <code>de_res</code> )                                  |

**Value**

list of valid inputs for plotting functions

**Description**

Outlier detection via POMA R Package

**Usage**

```
detect_outliers_POMA(  
  se,  
  ain = "log2",  
  condition = NULL,  
  method = "euclidean",  
  type = "median",  
  group = TRUE,  
  coeff = 1.5  
)
```

**Arguments**

|           |                                                                                                                                                                          |
|-----------|--------------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| se        | SummarizedExperiment containing all necessary information of the proteomics data set                                                                                     |
| ain       | String which data type should be used (default raw)                                                                                                                      |
| condition | Column name of condition (if NULL, condition saved in SummarizedExperiment will be taken)                                                                                |
| method    | String specifying the method that should be used to calculate the distance matrix                                                                                        |
| type      | String specifying the type of distance calculation to centroid or spatial median                                                                                         |
| group     | String specifying if the outlier detection should be performed multi-variate (with conditions) or on the complete data set                                               |
| coeff     | This value corresponds to the classical 1.5 in $Q3 + 1.5 * IQR$ formula to detect outliers. By changing this value, the permissiveness in outlier detection will change. |

**Value**

list of two ggplot objects and a data.table with outlier samples

**Examples**

```
data(tuberculosis_TMT_se)  
poma_res <- detect_outliers_POMA(tuberculosis_TMT_se, ain="raw",  
                                condition = NULL, method="euclidean",  
                                type="median", group=TRUE, coeff = 1.5)
```

---

|             |                              |
|-------------|------------------------------|
| eigenMSNorm | <i>EigenMS Normalization</i> |
|-------------|------------------------------|

---

### Description

EigenMS fits an analysis of variance model to estimate the effects of the experimental factors on the data using the knowledge about the experimental design, and then applies singular value decomposition to identify systematic trends contributing to significant variation not explained by the experimental factors. Log2-scaled data should be used as input (`on_raw = FALSE`).

### Usage

```
eigenMSNorm(se, ain = "log2", aout = "EigenMS", on_raw = FALSE)
```

### Arguments

|                     |                                                                                         |
|---------------------|-----------------------------------------------------------------------------------------|
| <code>se</code>     | SummarizedExperiment containing all necessary information of the proteomic dataset      |
| <code>ain</code>    | String which assay should be used as input                                              |
| <code>aout</code>   | String which assay should be used to save normalized data                               |
| <code>on_raw</code> | Boolean specifying whether normalization should be performed on raw or log2-scaled data |

### Value

SummarizedExperiment containing the EigenMS normalized data as assay (on log2 scale)

### Examples

```
data(tuberculosis_TMT_se)
tuberculosis_TMT_se <- eigenMSNorm(tuberculosis_TMT_se, ain = "log2",
                                   aout = "EigenMS", on_raw = FALSE)
```

---

|             |                                                                                        |
|-------------|----------------------------------------------------------------------------------------|
| export_data | <i>Export the SummarizedExperiment object, the meta data, and the normalized data.</i> |
|-------------|----------------------------------------------------------------------------------------|

---

### Description

Export the SummarizedExperiment object, the meta data, and the normalized data.

### Usage

```
export_data(se, out_dir, ain = NULL)
```

**Arguments**

|         |                                                                                                                        |
|---------|------------------------------------------------------------------------------------------------------------------------|
| se      | SummarizedExperiment containing all necessary information of the proteomics data set                                   |
| out_dir | Path of output directory                                                                                               |
| ain     | Vector of strings which assay should be downloaded (default NULL). If NULL then all assays of the se object are saved. |

**Value**

Nothing

**Examples**

```
data(tuberculosis_TMT_se)
## Not run: export_data(tuberculosis_TMT_se, out_dir = "data/",
  ain = c("IRS_on_RobNorm", "IRS_on_Median"))
## End(Not run)
```

---

|               |                                                                              |
|---------------|------------------------------------------------------------------------------|
| express_to_DT | <i>Helper function to transform an expression data frame to a data table</i> |
|---------------|------------------------------------------------------------------------------|

---

**Description**

Helper function to transform an expression data frame to a data table

**Usage**

```
express_to_DT(expr_data, column_names, row_names)
```

**Arguments**

|              |                                                      |
|--------------|------------------------------------------------------|
| expr_data    | Expression data frame containing the expression data |
| column_names | Column names of the expression data                  |
| row_names    | Row names of the expression data                     |

**Value**

Data table containing the expression data

---

extract\_consensus\_DE\_candidates

*Extract consensus DE candidates*

---

### Description

Extract consensus DE candidates

### Usage

```
extract_consensus_DE_candidates(
  de_res,
  ain = NULL,
  comparisons = NULL,
  norm_thr = 0.8,
  per_comparison = FALSE
)
```

### Arguments

|                |                                                                                                               |
|----------------|---------------------------------------------------------------------------------------------------------------|
| de_res         | data table resulting of run_DE                                                                                |
| ain            | Vector of strings of normalization methods to visualize (must be valid normalization methods saved in de_res) |
| comparisons    | Vector of comparisons (must be valid comparisons saved in de_res)                                             |
| norm_thr       | Threshold for the number of normalization methods that must agree on a DE candidate                           |
| per_comparison | Logical indicating if the consensus should be calculated per comparison                                       |

### Value

data table with consensus DE candidates

### Examples

```
data(tuberculosis_TMT_de_res)
extract_consensus_DE_candidates(tuberculosis_TMT_de_res, ain = NULL,
  comparisons = NULL, norm_thr = 0.8, per_comparison = TRUE)
```

---

extract\_limma\_DE

*Extract the DE results from eBayes fit of perform\_limma function.*

---

### Description

Extract the DE results from eBayes fit of perform\_limma function.

**Usage**

```
extract_limma_DE(
  fit,
  comparisons,
  logFC = TRUE,
  logFC_up = 1,
  logFC_down = -1,
  p_adj = TRUE,
  alpha = 0.05
)
```

**Arguments**

|             |                                                                                                        |
|-------------|--------------------------------------------------------------------------------------------------------|
| fit         | eBayes object resulting from perform_limma method                                                      |
| comparisons | Vector of comparisons that are performed in the DE analysis (from specify_comparisons method)          |
| logFC       | Boolean specifying whether to apply a logFC threshold (TRUE) or not (FALSE)                            |
| logFC_up    | Upper log2 fold change threshold (dividing into up regulated)                                          |
| logFC_down  | Lower log2 fold change threshold (dividing into down regulated)                                        |
| p_adj       | Boolean specifying whether to apply a threshold on adjusted p-values (TRUE) or on raw p-values (FALSE) |
| alpha       | Threshold for adjusted p-values or p-values                                                            |

**Value**

Data table with limma DE results

---

filter\_out\_complete\_NA\_proteins

*Remove proteins with NAs in all samples*

---

**Description**

Remove proteins with NAs in all samples

**Usage**

```
filter_out_complete_NA_proteins(se)
```

**Arguments**

|    |                                                                                      |
|----|--------------------------------------------------------------------------------------|
| se | SummarizedExperiment containing all necessary information of the proteomics data set |
|----|--------------------------------------------------------------------------------------|

**Value**

filtered SummarizedExperiment object

**Examples**

```
data(tuberculosis_TMT_se)
tuberculosis_TMT_se <- filter_out_complete_NA_proteins(tuberculosis_TMT_se)
```

---

```
filter_out_NA_proteins_by_threshold
```

*Filter proteins based on their NA pattern using a specific threshold*

---

**Description**

Filter proteins based on their NA pattern using a specific threshold

**Usage**

```
filter_out_NA_proteins_by_threshold(se, thr = 0.8)
```

**Arguments**

|     |                                                                                      |
|-----|--------------------------------------------------------------------------------------|
| se  | SummarizedExperiment containing all necessary information of the proteomics data set |
| thr | Threshold for the minimum fraction of valid values allowed for any protein           |

**Value**

filtered SummarizedExperiment object

**Examples**

```
data(tuberculosis_TMT_se)
tuberculosis_TMT_se <- filter_out_NA_proteins_by_threshold(tuberculosis_TMT_se,
  thr = 0.8)
```

---

```
filter_out_proteins_by_ID
```

*Remove proteins by their ID*

---

**Description**

Remove proteins by their ID

**Usage**

```
filter_out_proteins_by_ID(se, protein_ids)
```

**Arguments**

|             |                                                                                      |
|-------------|--------------------------------------------------------------------------------------|
| se          | SummarizedExperiment containing all necessary information of the proteomics data set |
| protein_ids | Vector of protein IDs that should be kept                                            |

**Value**

filtered SummarizedExperiment object

**Examples**

```
data(tuberculosis_TMT_se)
tuberculosis_TMT_se <- filter_out_proteins_by_ID(tuberculosis_TMT_se,
  protein_ids = c("P0A8V2", "P0A8V2"))
```

---

filter\_out\_proteins\_by\_value

*Remove proteins by value in specific column*

---

**Description**

Remove proteins by value in specific column

**Usage**

```
filter_out_proteins_by_value(se, column_name = "Reverse", values = c("+"))
```

**Arguments**

|             |                                                                                      |
|-------------|--------------------------------------------------------------------------------------|
| se          | SummarizedExperiment containing all necessary information of the proteomics data set |
| column_name | name of column of which proteins with a specific value should be removed             |
| values      | value of the column defining the proteins that should be removed                     |

**Value**

filtered SummarizedExperiment object

**Examples**

```
data(tuberculosis_TMT_se)
tuberculosis_TMT_se <- filter_out_proteins_by_value(tuberculosis_TMT_se,
  column_name = "Reverse", values = c("+"))
```

---

|                 |                                                                                          |
|-----------------|------------------------------------------------------------------------------------------|
| get_color_value | <i>Helper function to get correct value for coloration of plots (color_by parameter)</i> |
|-----------------|------------------------------------------------------------------------------------------|

---

**Description**

Helper function to get correct value for coloration of plots (color\_by parameter)

**Usage**

```
get_color_value(se, color_by)
```

**Arguments**

|          |                                                                                                                                                            |
|----------|------------------------------------------------------------------------------------------------------------------------------------------------------------|
| se       | SummarizedExperiment containing all necessary information of the proteomics data set                                                                       |
| color_by | String specifying the column to color the samples (If NULL, the condition column of the SummarizedExperiment object is used. If "No", no color bar added.) |

**Value**

String of column to color or NULL if no color should be applied

---

|                 |                                                                                          |
|-----------------|------------------------------------------------------------------------------------------|
| get_complete_dt | <i>Function to get a long data table of all intensities of all kind of normalization</i> |
|-----------------|------------------------------------------------------------------------------------------|

---

**Description**

Function to get a long data table of all intensities of all kind of normalization

**Usage**

```
get_complete_dt(se, ain = NULL)
```

**Arguments**

|     |                                                                                                                                                                          |
|-----|--------------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| se  | SummarizedExperiment containing all necessary information of the proteomics data set                                                                                     |
| ain | String which assay should be used as input (default NULL) If NULL then all normalization of the SummarizedExperiment object are plotted next to each other (except raw). |

**Value**

data table

---

|                     |                                                                                                   |
|---------------------|---------------------------------------------------------------------------------------------------|
| get_complete_pca_dt | <i>Function to get a long data table of all PCA1 and PCA2 values of all kind of normalization</i> |
|---------------------|---------------------------------------------------------------------------------------------------|

---

**Description**

Function to get a long data table of all PCA1 and PCA2 values of all kind of normalization

**Usage**

```
get_complete_pca_dt(se, ain = NULL)
```

**Arguments**

|     |                                                                                                                                                                          |
|-----|--------------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| se  | SummarizedExperiment containing all necessary information of the proteomics data set                                                                                     |
| ain | String which assay should be used as input (default NULL) If NULL then all normalization of the SummarizedExperiment object are plotted next to each other (except raw). |

**Value**

data table

---

|                     |                                                     |
|---------------------|-----------------------------------------------------|
| get_condition_value | <i>Helper function to check the condition value</i> |
|---------------------|-----------------------------------------------------|

---

**Description**

Helper function to check the condition value

**Usage**

```
get_condition_value(se, condition)
```

**Arguments**

|           |                                                                                           |
|-----------|-------------------------------------------------------------------------------------------|
| se        | SummarizedExperiment containing all necessary information of the proteomics data set      |
| condition | column name of condition (if NULL, condition saved in SummarizedExperiment will be taken) |

**Value**

String of column for condition

---

|                 |                                                                                        |
|-----------------|----------------------------------------------------------------------------------------|
| get_facet_value | <i>Helper function to get correct value for faceting of plots (facet_by parameter)</i> |
|-----------------|----------------------------------------------------------------------------------------|

---

**Description**

Helper function to get correct value for faceting of plots (facet\_by parameter)

**Usage**

```
get_facet_value(se, facet_by)
```

**Arguments**

|          |                                                                                           |
|----------|-------------------------------------------------------------------------------------------|
| se       | SummarizedExperiment containing all necessary information of the proteomics data set      |
| facet_by | String specifying the column to facet the samples (If NULL or "No", no faceting is done.) |

**Value**

String of column to facet or NULL if no faceting should be done

---

|                 |                                                                                               |
|-----------------|-----------------------------------------------------------------------------------------------|
| get_label_value | <i>Helper function to get correct value for sample labeling of plots (label_by parameter)</i> |
|-----------------|-----------------------------------------------------------------------------------------------|

---

**Description**

Helper function to get correct value for sample labeling of plots (label\_by parameter)

**Usage**

```
get_label_value(se, label_by)
```

**Arguments**

|          |                                                                                                                                                                  |
|----------|------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| se       | SummarizedExperiment containing all necessary information of the proteomics data set                                                                             |
| label_by | String specifying the column to label the samples (If NULL, the labels column of the SummarizedExperiment object is used. If "No", no labeling of samples done.) |

**Value**

String of column to label or NULL if no label should be applied

---

|                 |                                                                        |
|-----------------|------------------------------------------------------------------------|
| get_NA_overview | <i>Function returning some values on the numbers of NA in the data</i> |
|-----------------|------------------------------------------------------------------------|

---

**Description**

Function returning some values on the numbers of NA in the data

**Usage**

```
get_NA_overview(se, ain = "log2")
```

**Arguments**

|     |                                                                                      |
|-----|--------------------------------------------------------------------------------------|
| se  | SummarizedExperiment containing all necessary information of the proteomics data set |
| ain | String which data type should be used (default raw)                                  |

**Value**

list with total amount of values in the data, amount of NA values, and the percentage of NAs

**Examples**

```
data(tuberculosis_TMT_se)
get_NA_overview(tuberculosis_TMT_se, ain="log2")
```

---

|                           |                                                                             |
|---------------------------|-----------------------------------------------------------------------------|
| get_normalization_methods | <i>Function to return available normalization methods' identifier names</i> |
|---------------------------|-----------------------------------------------------------------------------|

---

**Description**

Function to return available normalization methods' identifier names

**Usage**

```
get_normalization_methods()
```

**Value**

Vector of normalization methods

**Examples**

```
get_normalization_methods()
```

---

|                 |                                         |
|-----------------|-----------------------------------------|
| get_overview_DE | <i>Get overview table of DE results</i> |
|-----------------|-----------------------------------------|

---

**Description**

Get overview table of DE results

**Usage**

```
get_overview_DE(de_res)
```

**Arguments**

|        |                                |
|--------|--------------------------------|
| de_res | data table resulting of run_DE |
|--------|--------------------------------|

**Value**

data table of numbers of DE proteins per comparison and per normalization method

**Examples**

```
data(tuberculosis_TMT_de_res)
get_overview_DE(tuberculosis_TMT_de_res)
```

---

|                       |                                                 |
|-----------------------|-------------------------------------------------|
| get_proteins_by_value | <i>Get proteins by value in specific column</i> |
|-----------------------|-------------------------------------------------|

---

**Description**

Get proteins by value in specific column

**Usage**

```
get_proteins_by_value(se, column_name = "Reverse", values = c("+"))
```

**Arguments**

|             |                                                                                      |
|-------------|--------------------------------------------------------------------------------------|
| se          | SummarizedExperiment containing all necessary information of the proteomics data set |
| column_name | name of column of which proteins with a specific value should be identified          |
| values      | value of the column defining the proteins that should be identified                  |

**Value**

vector of protein IDs

**Examples**

```
data(tuberculosis_TMT_se)
proteins <- get_proteins_by_value(tuberculosis_TMT_se,
                                column_name = "Potential.contaminant", values = c("+"))
```

---

|                 |                                                                                       |
|-----------------|---------------------------------------------------------------------------------------|
| get_shape_value | <i>Helper function to get correct value for shaping of plots (shape_by parameter)</i> |
|-----------------|---------------------------------------------------------------------------------------|

---

**Description**

Helper function to get correct value for shaping of plots (shape\_by parameter)

**Usage**

```
get_shape_value(se, shape_by)
```

**Arguments**

|          |                                                                                          |
|----------|------------------------------------------------------------------------------------------|
| se       | SummarizedExperiment containing all necessary information of the proteomics data set     |
| shape_by | String specifying the column to shape the samples (If NULL or "No", no shaping is done.) |

**Value**

String of column to shape or NULL if no shaping should be done

---

|                     |                                                                    |
|---------------------|--------------------------------------------------------------------|
| get_spiked_stats_DE | <i>Get performance metrics of DE results of spike-in data set.</i> |
|---------------------|--------------------------------------------------------------------|

---

**Description**

Get performance metrics of DE results of spike-in data set.

**Usage**

```
get_spiked_stats_DE(se, de_res)
```

**Arguments**

|        |                                                                                      |
|--------|--------------------------------------------------------------------------------------|
| se     | SummarizedExperiment containing all necessary information of the proteomics data set |
| de_res | data table resulting of run_DE                                                       |

**Value**

data table with multiple performance metrics of the DE results

**Examples**

```
data(spike_in_se)
data(spike_in_de_res)
stats <- get_spiked_stats_DE(spike_in_se, spike_in_de_res)
```

---

|               |                                      |
|---------------|--------------------------------------|
| globalIntNorm | <i>Total Intensity Normalization</i> |
|---------------|--------------------------------------|

---

### Description

Intensities of each variable in a sample are divided with the sum of intensities of all variables in the sample and multiplied with the median or mean of sum of intensities of all variables in all samples. Raw data should be taken as input (on\_raw = TRUE).

### Usage

```
globalIntNorm(  
  se,  
  ain = "raw",  
  aout = "GlobalMedian",  
  type = "median",  
  on_raw = TRUE  
)
```

### Arguments

|        |                                                                                         |
|--------|-----------------------------------------------------------------------------------------|
| se     | SummarizedExperiment containing all necessary information of the proteomics data set    |
| ain    | String which assay should be used as input                                              |
| aout   | String which assay should be used to save normalized data                               |
| type   | String whether to use median or mean to calculate the scaling factor                    |
| on_raw | Boolean specifying whether normalization should be performed on raw or log2-scaled data |

### Value

SummarizedExperiment containing the total intensity normalized data as assay (on log2 scale)

### Examples

```
data(tuberculosis_TMT_se)  
tuberculosis_TMT_se <- globalIntNorm(tuberculosis_TMT_se, ain = "raw",  
                                     aout = "GlobalMedian",  
                                     type = "median",  
                                     on_raw = TRUE)
```

---

|                |                                                                                            |
|----------------|--------------------------------------------------------------------------------------------|
| globalMeanNorm | <i>Total Intensity Normalization Using the Mean for the Calculation of Scaling Factors</i> |
|----------------|--------------------------------------------------------------------------------------------|

---

**Description**

Intensities of each variable in a sample are divided with the sum of intensities of all variables in the sample and multiplied with the mean of sum of intensities of all variables in all samples. Raw data should be taken as input (on\_raw = TRUE).

**Usage**

```
globalMeanNorm(se, ain = "raw", aout = "GlobalMean", on_raw = TRUE)
```

**Arguments**

|        |                                                                                         |
|--------|-----------------------------------------------------------------------------------------|
| se     | SummarizedExperiment containing all necessary information of the proteomics data set    |
| ain    | String which assay should be used as input                                              |
| aout   | String which assay should be used to save normalized data                               |
| on_raw | Boolean specifying whether normalization should be performed on raw or log2-scaled data |

**Value**

SummarizedExperiment containing the total intensity normalized data as assay (on log2 scale)

**Examples**

```
data(tuberculosis_TMT_se)
tuberculosis_TMT_se <- globalMeanNorm(tuberculosis_TMT_se, ain = "raw",
                                     aout = "GlobalMean", on_raw = TRUE)
```

---

|                  |                                                                                              |
|------------------|----------------------------------------------------------------------------------------------|
| globalMedianNorm | <i>Total Intensity Normalization Using the Median for the Calculation of Scaling Factors</i> |
|------------------|----------------------------------------------------------------------------------------------|

---

**Description**

Intensities of each variable in a sample are divided with the sum of intensities of all variables in the sample and multiplied with the median of sum of intensities of all variables in all samples. Raw data should be taken as input (on\_raw = TRUE).

**Usage**

```
globalMedianNorm(se, ain = "raw", aout = "GlobalMedian", on_raw = TRUE)
```

**Arguments**

|        |                                                                                         |
|--------|-----------------------------------------------------------------------------------------|
| se     | SummarizedExperiment containing all necessary information of the proteomics data set    |
| ain    | String which assay should be used as input                                              |
| aout   | String which assay should be used to save normalized data                               |
| on_raw | Boolean specifying whether normalization should be performed on raw or log2-scaled data |

**Value**

SummarizedExperiment containing the total intensity normalized data as assay (on log2 scale)

**Examples**

```
data(tuberculosis_TMT_se)
tuberculosis_TMT_se <- globalMedianNorm(tuberculosis_TMT_se, ain = "raw",
                                         aout = "GlobalMedian", on_raw = TRUE)
```

---

|           |                                                                                                                                                                                                                                                                                                                                             |
|-----------|---------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| impute_se | <i>Method to impute SummarizedExperiment. This method performs a mixed imputation on the proteins. It uses a k-nearest neighbor imputation for proteins with missing values at random (MAR) and imputes missing values by random draws from a left-shifted Gaussian distribution for proteins with missing values not at random (MNAR).</i> |
|-----------|---------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|

---

**Description**

Method to impute SummarizedExperiment. This method performs a mixed imputation on the proteins. It uses a k-nearest neighbor imputation for proteins with missing values at random (MAR) and imputes missing values by random draws from a left-shifted Gaussian distribution for proteins with missing values not at random (MNAR).

**Usage**

```
impute_se(se, ain = NULL, condition = NULL)
```

**Arguments**

|           |                                                                                                                                                       |
|-----------|-------------------------------------------------------------------------------------------------------------------------------------------------------|
| se        | SummarizedExperiment containing all necessary information of the proteomics dataset                                                                   |
| ain       | Vector of strings which assay should be used as input (default NULL). If NULL then all normalization of the se object are plotted next to each other. |
| condition | name of column of colData(se) representing the conditions of the data                                                                                 |

**Value**

SummarizedExperiment with imputed intensities

## Examples

```
data(tuberculosis_TMT_se)
tuberculosis_TMT_se <- remove_samples_manually(tuberculosis_TMT_se,
  column = "Label", values = c("1.HC.Pool1"))
tuberculosis_TMT_se <- impute_se(tuberculosis_TMT_se, ain = NULL,
  condition = NULL)
```

---

irsNorm

*Internal Reference Scaling Normalization*

---

## Description

IRS makes different measurements of the same thing all exactly the same and puts all of the intensities on the same scale. Raw data should be taken as input (`on_raw = TRUE`)

## Usage

```
irsNorm(se, ain = "raw", aout = "IRS", on_raw = TRUE)
```

## Arguments

|        |                                                                                         |
|--------|-----------------------------------------------------------------------------------------|
| se     | SummarizedExperiment containing all necessary information of the proteomic dataset      |
| ain    | String which assay should be used as input                                              |
| aout   | String which assay should be used to save normalized data                               |
| on_raw | Boolean specifying whether normalization should be performed on raw or log2-scaled data |

## Value

SummarizedExperiment containing the IRS normalized data as assay (on log2 scale)

## Examples

```
data(tuberculosis_TMT_se)
tuberculosis_TMT_se <- irsNorm(tuberculosis_TMT_se, ain = "raw",
  aout = "IRS", on_raw = TRUE)
```

---

|           |                                          |
|-----------|------------------------------------------|
| limmaNorm | <i>limma::removeBatchEffects (limBE)</i> |
|-----------|------------------------------------------|

---

### Description

Log2-scaled data should be used as input (on\_raw = FALSE).

### Usage

```
limmaNorm(se, ain = "log2", aout = "limBE", on_raw = FALSE)
```

### Arguments

|        |                                                                                         |
|--------|-----------------------------------------------------------------------------------------|
| se     | SummarizedExperiment containing all necessary information of the proteomic dataset      |
| ain    | String which assay should be used as input                                              |
| aout   | String which assay should be used to save normalized data                               |
| on_raw | Boolean specifying whether normalization should be performed on raw or log2-scaled data |

### Value

SummarizedExperiment containing the limBE normalized data as assay (on log2 scale)

### See Also

[removeBatchEffect\(\)](#)

### Examples

```
data(tuberculosis_TMT_se)
tuberculosis_TMT_se <- limmaNorm(tuberculosis_TMT_se, ain = "log2",
                                aout = "limBE", on_raw = FALSE)
```

---

|           |                                                                    |
|-----------|--------------------------------------------------------------------|
| load_data | <i>Load real-world proteomics data into a SummarizedExperiment</i> |
|-----------|--------------------------------------------------------------------|

---

### Description

Load real-world proteomics data into a SummarizedExperiment

**Usage**

```
load_data(
  data,
  md,
  protein_column = "Protein.IDs",
  gene_column = "Gene.Names",
  ref_samples = NULL,
  batch_column = NULL,
  condition_column = NULL,
  label_column = NULL
)
```

**Arguments**

|                               |                                                                                                                     |
|-------------------------------|---------------------------------------------------------------------------------------------------------------------|
| <code>data</code>             | tabular data table with rows = proteins and columns = samples (such as protein-Groups.txt of MaxQuant)              |
| <code>md</code>               | experimental design table (requires a column named "Column" for the column names of the sample intensities in data) |
| <code>protein_column</code>   | name of the column in data containing the protein IDs                                                               |
| <code>gene_column</code>      | name of the column in data containing the gene names                                                                |
| <code>ref_samples</code>      | reference samples if TMT experiment provided (names of samples)                                                     |
| <code>batch_column</code>     | name of the column in md defining the batches                                                                       |
| <code>condition_column</code> | name of the column in md defining the condition (can still be changed afterwards)                                   |
| <code>label_column</code>     | name of the column in md containing simple sample names (for visualization)                                         |

**Value**

SummarizedExperiment object

**Examples**

```
data_path <- readPRONE_example("tuberculosis_protein_intensities.csv")
md_path <- readPRONE_example("tuberculosis_metadata.csv")
data <- read.csv(data_path)
md <- read.csv(md_path)
md$Column <- stringr::str_replace_all(md$Column, " ", ".")
ref_samples <- md[md$Group == "ref",]$Column
se <- load_data(data, md, protein_column = "Protein.IDs",
  gene_column = "Gene.names", ref_samples = ref_samples,
  batch_column = "Pool", condition_column = "Group",
  label_column = "Label")
```

---

|                 |                                                                  |
|-----------------|------------------------------------------------------------------|
| load_spike_data | <i>Load spike-in proteomics data into a SummarizedExperiment</i> |
|-----------------|------------------------------------------------------------------|

---

## Description

Load spike-in proteomics data into a SummarizedExperiment

## Usage

```
load_spike_data(
  data,
  md,
  spike_column,
  spike_value,
  spike_concentration,
  protein_column = "Protein.IDs",
  gene_column = "Gene.Names",
  ref_samples = NULL,
  batch_column = NULL,
  condition_column = NULL,
  label_column = NULL
)
```

## Arguments

|                     |                                                                                                                     |
|---------------------|---------------------------------------------------------------------------------------------------------------------|
| data                | tabular data table with rows = proteins and columns = samples (such as protein-Groups.txt of MaxQuant)              |
| md                  | experimental design table (requires a column named "Column" for the column names of the sample intensities in data) |
| spike_column        | name of the column specifying which proteins are the spike-ins                                                      |
| spike_value         | String value specifying the spike-in proteins in the spike-in column                                                |
| spike_concentration | name of the column in md defining the spike-in concentration per sample                                             |
| protein_column      | name of the column in data containing the protein IDs                                                               |
| gene_column         | name of the column in data containing the gene names                                                                |
| ref_samples         | reference samples if TMT experiment provided (names of samples)                                                     |
| batch_column        | name of the column in md defining the batches                                                                       |
| condition_column    | name of the column in md defining the condition (can still be changed afterwards)                                   |
| label_column        | name of the column in md containing simple sample names (for visualization)                                         |

## Value

SummarizedExperiment object

**Examples**

```

data_path <- readPRONE_example("Ecoli_human_MaxLFQ_protein_intensities.csv")
md_path <- readPRONE_example("Ecoli_human_MaxLFQ_metadata.csv")
data <- read.csv(data_path)
md <- read.csv(md_path)
mixed <- grepl("Homo sapiens.*Escherichia|Escherichia.*Homo sapiens", data$Fasta.headers)
data <- data[!mixed,]
data$Spiked <- rep("HUMAN", nrow(data))
data$Spiked[grepl("ECOLI", data$Fasta.headers)] <- "ECOLI"
se <- load_spike_data(data, md, spike_column = "Spiked", spike_value = "ECOLI",
  spike_concentration = "Concentration", protein_column = "Protein.IDs",
  gene_column = "Gene.names", ref_samples = NULL, batch_column = NULL,
  condition_column = "Condition", label_column = "Label")

```

loessCycNorm

*Cyclic Loess Normalization of limma***Description**

Two samples of the data are MA transformed and normalized at a time, and all pairs of samples are iterated through. Log2-scaled data should be taken as input (on\_raw = FALSE).

**Usage**

```
loessCycNorm(se, ain = "log2", aout = "LoessCyc", on_raw = FALSE)
```

**Arguments**

|        |                                                                                         |
|--------|-----------------------------------------------------------------------------------------|
| se     | SummarizedExperiment containing all necessary information of the proteomic dataset      |
| ain    | String which assay should be used as input                                              |
| aout   | String which assay should be used to save normalized data                               |
| on_raw | Boolean specifying whether normalization should be performed on raw or log2-scaled data |

**Value**

SummarizedExperiment containing the loessCyc normalized data as assay (on log2 scale)

**See Also**

[normalizeCyclicLoess\(\)](#)

**Examples**

```

data(tuberculosis_TMT_se)
tuberculosis_TMT_se <- loessCycNorm(tuberculosis_TMT_se, ain = "log2",
  aout = "LoessCyc", on_raw = FALSE)

```

---

|            |                                          |
|------------|------------------------------------------|
| loessFNorm | <i>Fast Loess Normalization of limma</i> |
|------------|------------------------------------------|

---

**Description**

Using mean intensities over all the samples as its reference A sample. Log2-scaled data should be used as input (on\_raw = FALSE).

**Usage**

```
loessFNorm(se, ain = "log2", aout = "LoessF", on_raw = FALSE)
```

**Arguments**

|        |                                                                                         |
|--------|-----------------------------------------------------------------------------------------|
| se     | SummarizedExperiment containing all necessary information of the proteomic dataset      |
| ain    | String which assay should be used as input                                              |
| aout   | String which assay should be used to save normalized data                               |
| on_raw | Boolean specifying whether normalization should be performed on raw or log2-scaled data |

**Value**

SummarizedExperiment containing the LoessF normalized data as assay (on log2 scale)

**See Also**

[normalizeCyclicLoess\(\)](#)

**Examples**

```
data(tuberculosis_TMT_se)
tuberculosis_TMT_se <- loessFNorm(tuberculosis_TMT_se, ain = "log2",
                                aout = "LoessCyc", on_raw = FALSE)
```

---

|          |                           |
|----------|---------------------------|
| meanNorm | <i>Mean Normalization</i> |
|----------|---------------------------|

---

**Description**

The intensity of each protein group in a given sample is divided by the mean of the intensities of all protein groups in that sample and then multiplied by the mean of mean of sum of intensities of all protein groups in all samples.

**Usage**

```
meanNorm(se, ain = "raw", aout = "Mean", on_raw = TRUE)
```



---

|            |                             |
|------------|-----------------------------|
| medianNorm | <i>Median Normalization</i> |
|------------|-----------------------------|

---

### Description

The intensity of each protein group in a given sample is divided by the median of the intensities of all protein groups in that sample and then multiplied by the mean of median of sum of intensities of all protein groups in all samples.

### Usage

```
medianNorm(se, ain = "raw", aout = "Median", on_raw = TRUE)
```

### Arguments

|        |                                                                                         |
|--------|-----------------------------------------------------------------------------------------|
| se     | SummarizedExperiment containing all necessary information of the proteomic dataset      |
| ain    | String which assay should be used as input                                              |
| aout   | String which assay should be used to save normalized data                               |
| on_raw | Boolean specifying whether normalization should be performed on raw or log2-scaled data |

### Value

SummarizedExperiment containing the median normalized data as assay (on log2 scale)

### Examples

```
data(tuberculosis_TMT_se)
tuberculosis_TMT_se <- medianNorm(tuberculosis_TMT_se, ain = "raw",
                                  aout = "Median", on_raw = TRUE)
```

---

|              |                                                                                                                                    |
|--------------|------------------------------------------------------------------------------------------------------------------------------------|
| normalize_se | <i>Normalize SummarizedExperiment object using single normalization methods or specified combinations of normalization methods</i> |
|--------------|------------------------------------------------------------------------------------------------------------------------------------|

---

### Description

Normalize SummarizedExperiment object using single normalization methods or specified combinations of normalization methods

**Usage**

```
normalize_se(
  se,
  methods,
  combination_pattern = "_on_",
  on_raw = NULL,
  gamma.0 = 0.1,
  reduce_correlation_by = 1,
  NormicsVSN_quantile = 0.8,
  top_x = 50,
  VSN_quantile = 0.9
)
```

**Arguments**

|                       |                                                                                                                                                                                                                                                                                                                                                                              |
|-----------------------|------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| se                    | SummarizedExperiment containing all necessary information of the proteomics data set                                                                                                                                                                                                                                                                                         |
| methods               | Vector of normalization methods to apply for normalizing the proteomics data of the SummarizedExperiment object (identifier of normalization methods can be retrieved using <code>get_normalization_methods()</code> )                                                                                                                                                       |
| combination_pattern   | String specifying how normalization methods are combined. For instance, <code>methods = c("IRS", "Median_on_IRS")</code> , <code>combination_pattern = "_on_"</code> .                                                                                                                                                                                                       |
| on_raw                | Logical indicating if the normalization should be performed on the raw data or on log2-transformed data. If <code>on_raw = NULL</code> (default), the normalization is performed on the default method specific <code>on_raw</code> setting (suggestion based on publications).                                                                                              |
| gamma.0               | Numeric representing the exponent of the weighted density of RobNorm normalization. When the sample size is small, the fitted population of some proteins could be locally trapped such that the variance of those proteins was very small under a large gamma. To avoid this, a small gamma is recommended. When sample size smaller than 40, then set gamma to 0.5 or 0.1. |
| reduce_correlation_by | If the data is too big for the computation of the params, increase this parameter by 2,3,4.... The whole data will still be normalized, but the params are calculated on every second row etc.                                                                                                                                                                               |
| NormicsVSN_quantile   | The quantile that is used for the resistant least trimmed sum of squares regression. A value of 0.8 means focusing on the central 80% of the data, reducing the influence of outliers.                                                                                                                                                                                       |
| top_x                 | Number of reference proteins extracted for the calculation of parameters                                                                                                                                                                                                                                                                                                     |
| VSN_quantile          | Numeric of length 1. The quantile that is used for the resistant least trimmed sum of squares regression (see <code>vsn2 lts.quantile</code> )                                                                                                                                                                                                                               |

**Value**

SummarizedExperiment object with normalized data saved as assays

## Examples

```
data(tuberculosis_TMT_se)
tuberculosis_TMT_se <- normalize_se(tuberculosis_TMT_se,
  methods = c("IRS_on_GlobalMedian", "IRS_on_Median",
    "limBE_on_NormicsVSN"), on_raw = NULL,
  combination_pattern = "_on_", gamma.0 = 0.1,
  reduce_correlation_by = 1, NormicsVSN_quantile = 0.8, top_x = 50,
  VSN_quantile = 0.9)
```

---

normalize\_se\_combination

*Normalize SummarizedExperiment object using combinations of normalization methods*

---

## Description

Normalize SummarizedExperiment object using combinations of normalization methods

## Usage

```
normalize_se_combination(
  se,
  methods,
  ains,
  on_raw = NULL,
  combination_pattern = "_on_",
  gamma.0 = 0.1,
  reduce_correlation_by = 1,
  NormicsVSN_quantile = 0.8,
  top_x = 50,
  VSN_quantile = 0.9
)
```

## Arguments

|                     |                                                                                                                                                                                                                                                                                 |
|---------------------|---------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| se                  | SummarizedExperiment containing all necessary information of the proteomics data set                                                                                                                                                                                            |
| methods             | Vector of normalization methods to apply for normalizing the proteomics data of the SummarizedExperiment object (identifier of normalization methods can be retrieved using <code>get_normalization_methods()</code> )                                                          |
| ains                | Vector of assays of SummarizedExperiment object to apply the normalization methods (e.g. if you want to perform Median normalization on IRS-normalized data)                                                                                                                    |
| on_raw              | Logical indicating if the normalization should be performed on the raw data or on log2-transformed data. If <code>on_raw = NULL</code> (default), the normalization is performed on the default method specific <code>on_raw</code> setting (suggestion based on publications). |
| combination_pattern | String to give name to combination of methods (e.g. <code>IRS_on_Median</code> → <code>"_on_"</code> )                                                                                                                                                                          |

|                       |                                                                                                                                                                                                                                                                                                                                                                              |
|-----------------------|------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| gamma.0               | Numeric representing the exponent of the weighted density of RobNorm normalization. When the sample size is small, the fitted population of some proteins could be locally trapped such that the variance of those proteins was very small under a large gamma. To avoid this, a small gamma is recommended. When sample size smaller than 40, then set gamma to 0.5 or 0.1. |
| reduce_correlation_by | If the data is too big for the computation of the params, increase this parameter by 2,3,4.... The whole data will still be normalized, but the params are calculated on every second row etc.                                                                                                                                                                               |
| NormicsVSN_quantile   | The quantile that is used for the resistant least trimmed sum of squares regression. A value of 0.8 means focusing on the central 80% of the data, reducing the influence of outliers.                                                                                                                                                                                       |
| top_x                 | Number of reference proteins extracted for the calculation of parameters                                                                                                                                                                                                                                                                                                     |
| VSN_quantile          | Numeric of length 1. The quantile that is used for the resistant least trimmed sum of squares regression. (see vsn2 lts.quantile)                                                                                                                                                                                                                                            |

**Value**

SummarizedExperiment object with normalized data saved as assays

**Examples**

```
data(tuberculosis_TMT_se)
tuberculosis_TMT_se <- normalize_se_combination(tuberculosis_TMT_se,
  methods = c("Median", "NormicsVSN"), ains = c("IRS"), on_raw = NULL,
  combination_pattern = "_on_", gamma.0 = 0.1,
  reduce_correlation_by = 1, NormicsVSN_quantile = 0.8, top_x = 50,
  VSN_quantile = 0.9)
```

---

|                     |                                                                                    |
|---------------------|------------------------------------------------------------------------------------|
| normalize_se_single | <i>Normalize SummarizedExperiment object using different normalization methods</i> |
|---------------------|------------------------------------------------------------------------------------|

---

**Description**

Normalize SummarizedExperiment object using different normalization methods

**Usage**

```
normalize_se_single(
  se,
  methods = NULL,
  on_raw = NULL,
  gamma.0 = 0.1,
  reduce_correlation_by = 1,
  NormicsVSN_quantile = 0.8,
  top_x = 50,
  VSN_quantile = 0.9
)
```

**Arguments**

|                       |                                                                                                                                                                                                                                                                                                                                                                              |
|-----------------------|------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| se                    | SummarizedExperiment containing all necessary information of the proteomics data set                                                                                                                                                                                                                                                                                         |
| methods               | Vector of normalization methods to apply for normalizing the proteomics data of the SummarizedExperiment object (identifier of normalization methods can be retrieved using <code>get_normalization_methods()</code> )                                                                                                                                                       |
| on_raw                | Logical indicating if the normalization should be performed on the raw data or on log2-transformed data. If <code>on_raw = NULL</code> (default), the normalization is performed on the default method specific <code>on_raw</code> setting (suggestion based on publications).                                                                                              |
| gamma.0               | Numeric representing the exponent of the weighted density of RobNorm normalization. When the sample size is small, the fitted population of some proteins could be locally trapped such that the variance of those proteins was very small under a large gamma. To avoid this, a small gamma is recommended. When sample size smaller than 40, then set gamma to 0.5 or 0.1. |
| reduce_correlation_by | If the data is too big for the computation of the params, increase this parameter by 2,3,4.... The whole data will still be normalized, but the params are calculated on every second row etc.                                                                                                                                                                               |
| NormicsVSN_quantile   | The quantile that is used for the resistant least trimmed sum of squares regression. A value of 0.8 means focusing on the central 80% of the data, reducing the influence of outliers.                                                                                                                                                                                       |
| top_x                 | Number of reference proteins extracted for the calculation of parameters                                                                                                                                                                                                                                                                                                     |
| VSN_quantile          | Numeric of length 1. The quantile that is used for the resistant least trimmed sum of squares regression. (see <code>vsn2 lts.quantile</code> )                                                                                                                                                                                                                              |

**Value**

SummarizedExperiment object with normalized data saved as assays

**Examples**

```
data(tuberculosis_TMT_se)
tuberculosis_TMT_se <- normalize_se_single(tuberculosis_TMT_se,
  methods = c("RobNorm", "Median", "NormicsVSN", "VSN"),
  on_raw = NULL, gamma.0 = 0.1, reduce_correlation_by = 1,
  NormicsVSN_quantile = 0.8, top_x = 50, VSN_quantile = 0.9)
```

---

normicsNorm

*Normics Normalization (Normics using VSN or using Median)*


---

**Description**

Log2-scaled data should be used as input (`on_raw = FALSE`).

**Usage**

```
normicsNorm(
  se,
  ain = "raw",
  aout = "NormicsVSN",
  method = "NormicsVSN",
  on_raw = TRUE,
  reduce_correlation_by = 1,
  NormicsVSN_quantile = 0.8,
  TMT_ratio = FALSE,
  top_x = 50
)
```

**Arguments**

|                       |                                                                                                                                                                                                |
|-----------------------|------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| se                    | SummarizedExperiment containing all necessary information of the proteomic dataset                                                                                                             |
| ain                   | String which assay should be used as input                                                                                                                                                     |
| aout                  | String which assay should be used to save normalized data                                                                                                                                      |
| method                | String specifying the method to use (NORMICS or NORMICSmedian)                                                                                                                                 |
| on_raw                | Boolean specifying whether normalization should be performed on raw or log2-scaled data                                                                                                        |
| reduce_correlation_by | If the data is too big for the computation of the params, increase this parameter by 2,3,4.... The whole data will still be normalized, but the params are calculated on every second row etc. |
| NormicsVSN_quantile   | The quantile that is used for the resistant least trimmed sum of squares regression. A value of 0.8 means focusing on the central 80% of the data, reducing the influence of outliers.         |
| TMT_ratio             | Indicates if the data involves Tandem Mass Tag (TMT) ratio-based measurements (common in proteomics). If TRUE, the method may handle the data differently.                                     |
| top_x                 | Number of reference proteins extracted for the calculation of parameters                                                                                                                       |

**Value**

SummarizedExperiment containing the NormicsVSN/NormicsMedian normalized data as assay (on log2 scale)

**Examples**

```
data(tuberculosis_TMT_se)
tuberculosis_TMT_se <- normicsNorm(tuberculosis_TMT_se, ain = "raw",
                                  aout = "NormicsVSN", method = "NormicsVSN",
                                  on_raw = TRUE)
```

---

|               |                      |
|---------------|----------------------|
| perform_DEqMS | <i>Perform DEqMS</i> |
|---------------|----------------------|

---

**Description**

Perform DEqMS

**Usage**

```
perform_DEqMS(  
  fit,  
  se,  
  DEqMS_PSMs_column = NULL,  
  logFC = TRUE,  
  logFC_up = 1,  
  logFC_down = -1,  
  p_adj = TRUE,  
  alpha = 0.05  
)
```

**Arguments**

|                   |                                                                                                                 |
|-------------------|-----------------------------------------------------------------------------------------------------------------|
| fit               | eBayes object resulting from perform_limma method                                                               |
| se                | SummarizedExperiment containing all necessary information of the proteomics data set                            |
| DEqMS_PSMs_column | String specifying which column name to use for DEqMS (default NULL). Any column of the rowData(se) is accepted. |
| logFC             | Boolean specifying whether to apply a logFC threshold (TRUE) or not (FALSE)                                     |
| logFC_up          | Upper log2 fold change threshold (dividing into up regulated)                                                   |
| logFC_down        | Lower log2 fold change threshold (dividing into down regulated)                                                 |
| p_adj             | Boolean specifying whether to apply a threshold on adjusted p-values (TRUE) or on raw p-values (FALSE)          |
| alpha             | Threshold for adjusted p-values or p-values                                                                     |

**Value**

data.table of DE results

---

|               |                                           |
|---------------|-------------------------------------------|
| perform_limma | <i>Fitting a linear model using limma</i> |
|---------------|-------------------------------------------|

---

**Description**

Fitting a linear model using limma

**Usage**

```
perform_limma(
  data,
  condition_vector,
  comparisons,
  covariate = NULL,
  trend = TRUE,
  robust = TRUE
)
```

**Arguments**

|                  |                                                                                                                                                                                                                                                  |
|------------------|--------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| data             | Data table of intensities (rows = proteins, cols = samples)                                                                                                                                                                                      |
| condition_vector | Vector of experimental design specifying the condition(s) to compare                                                                                                                                                                             |
| comparisons      | Vector of comparisons that are performed in the DE analysis (from specify_comparisons method)                                                                                                                                                    |
| covariate        | String specifying which column to include as covariate into limma                                                                                                                                                                                |
| trend            | logical, should an intensity-dependent trend be allowed for the prior variance? If FALSE then the prior variance is constant. Alternatively, trend can be a row-wise numeric vector, which will be used as the covariate for the prior variance. |
| robust           | logical, should the estimation of df.prior and var.prior be robustified against outlier sample variances?                                                                                                                                        |

**Value**

eBayes object

---

|               |                         |
|---------------|-------------------------|
| perform_ROTSt | <i>Performing ROTSt</i> |
|---------------|-------------------------|

---

**Description**

Performing ROTSt

**Usage**

```
perform_ROTSt(
  data,
  condition,
  comparisons,
  condition_name,
  coldata,
  logFC = TRUE,
  logFC_up = 1,
  logFC_down = -1,
  p_adj = TRUE,
  alpha = 0.05,
  B = 100,
  K = 500
)
```

**Arguments**

|                |                                                                                                        |
|----------------|--------------------------------------------------------------------------------------------------------|
| data           | Data table of intensities (rows = proteins, cols = samples)                                            |
| condition      | Vector of experimental design specifying the condition(s) to compare                                   |
| comparisons    | Vector of comparisons that are performed in the DE analysis (from specify_comparisons method)          |
| condition_name | String of name of condition in colData                                                                 |
| coldata        | colData of the SummarizedExperiment                                                                    |
| logFC          | Boolean specifying whether to apply a logFC threshold (TRUE) or not (FALSE)                            |
| logFC_up       | Upper log2 fold change threshold (dividing into up regulated)                                          |
| logFC_down     | Lower log2 fold change threshold (dividing into down regulated)                                        |
| p_adj          | Boolean specifying whether to apply a threshold on adjusted p-values (TRUE) or on raw p-values (FALSE) |
| alpha          | Threshold for adjusted p-values or p-values                                                            |
| B              | Number of bootstrapping for ROTS                                                                       |
| K              | Number of top-ranked features for reproducibility optimization                                         |

**Value**

Data table with DE results

---

|               |                                                                  |
|---------------|------------------------------------------------------------------|
| plot_boxplots | <i>Plot the distributions of the normalized data as boxplots</i> |
|---------------|------------------------------------------------------------------|

---

**Description**

Plot the distributions of the normalized data as boxplots

**Usage**

```
plot_boxplots(
  se,
  ain = NULL,
  color_by = NULL,
  label_by = NULL,
  facet_norm = TRUE,
  ncol = 3
)
```

**Arguments**

|          |                                                                                                                                                            |
|----------|------------------------------------------------------------------------------------------------------------------------------------------------------------|
| se       | SummarizedExperiment containing all necessary information of the proteomics data set                                                                       |
| ain      | Vector of strings which assay should be used as input (default NULL). If NULL then all normalization of the se object are plotted next to each other.      |
| color_by | String specifying the column to color the samples (If NULL, the condition column of the SummarizedExperiment object is used. If "No", no color bar added.) |

|            |                                                                                                                                                                  |
|------------|------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| label_by   | String specifying the column to label the samples (If NULL, the labels column of the SummarizedExperiment object is used. If "No", no labeling of samples done.) |
| facet_norm | Boolean specifying whether to facet by normalization methods (default TRUE). If FALSE, list of plots of the different normalized data is returned.               |
| ncol       | Number of columns in plot (for faceting)                                                                                                                         |

**Value**

if facet\_norm = TRUE, ggplot object, else list of ggplot objects

**Examples**

```
data(tuberculosis_TMT_se)
plot_boxplots(tuberculosis_TMT_se, ain = NULL, color_by = NULL, label_by = NULL,
              facet_norm = TRUE, ncol = 3)
plot_boxplots(tuberculosis_TMT_se, ain = c("log2", "IRS_on_RobNorm"), color_by = "Pool",
              label_by = "Label", facet_norm = FALSE)
```

---

plot\_condition\_overview

*Barplot showing the number of samples per condition*

---

**Description**

Barplot showing the number of samples per condition

**Usage**

```
plot_condition_overview(se, condition = NULL)
```

**Arguments**

|           |                                                                                           |
|-----------|-------------------------------------------------------------------------------------------|
| se        | SummarizedExperiment containing all necessary information of the proteomics data set      |
| condition | column name of condition (if NULL, condition saved in SummarizedExperiment will be taken) |

**Value**

ggplot object

**Examples**

```
data(tuberculosis_TMT_se)
plot_condition_overview(tuberculosis_TMT_se, condition = NULL)
```

---

|                |                                                  |
|----------------|--------------------------------------------------|
| plot_densities | <i>Plot the densities of the normalized data</i> |
|----------------|--------------------------------------------------|

---

### Description

Plot the densities of the normalized data

### Usage

```
plot_densities(se, ain = NULL, color_by = NULL, facet_norm = TRUE, ncol = 3)
```

### Arguments

|            |                                                                                                                                                            |
|------------|------------------------------------------------------------------------------------------------------------------------------------------------------------|
| se         | SummarizedExperiment containing all necessary information of the proteomics data set                                                                       |
| ain        | Vector of strings which assay should be used as input (default NULL). If NULL then all normalization of the se object are plotted next to each other.      |
| color_by   | String specifying the column to color the samples (If NULL, the condition column of the SummarizedExperiment object is used. If "No", no color bar added.) |
| facet_norm | Boolean specifying whether to facet by normalization methods (default TRUE). If FALSE, list of plots of the different normalized data is returned.         |
| ncol       | Number of columns in plot (for faceting)                                                                                                                   |

### Value

if facet\_norm = TRUE, ggplot object, else list of ggplot objects

### Examples

```
data(tuberculosis_TMT_se)
plot_densities(tuberculosis_TMT_se, ain = NULL, color_by = NULL,
               facet_norm = TRUE, ncol = 3)
plot_densities(tuberculosis_TMT_se, ain = c("log2", "IRS_on_RobNorm"),
               color_by = "Label",
               facet_norm = FALSE)
```

---

|                          |                                                                                                                                                                                                                                      |
|--------------------------|--------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| plot_fold_changes_spiked | <i>Boxplot of log fold changes of spike-in and background proteins for specific normalization methods and comparisons. The ground truth (calculated based on the concentrations of the spike-ins) is shown as a horizontal line.</i> |
|--------------------------|--------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|

---

### Description

Boxplot of log fold changes of spike-in and background proteins for specific normalization methods and comparisons. The ground truth (calculated based on the concentrations of the spike-ins) is shown as a horizontal line.

**Usage**

```
plot_fold_changes_spiked(se, de_res, condition, ain = NULL, comparisons = NULL)
```

**Arguments**

|             |                                                                                                              |
|-------------|--------------------------------------------------------------------------------------------------------------|
| se          | SummarizedExperiment containing all necessary information of the proteomics data set                         |
| de_res      | data table resulting of run_DE                                                                               |
| condition   | column name of condition (if NULL, condition saved in SummarizedExperiment will be taken)                    |
| ain         | Vector of strings of normalization methods to visualize (must be valid normalization methods saved in stats) |
| comparisons | Vector of comparisons (must be valid comparisons saved in stats)                                             |

**Value**

ggplot object

**Examples**

```
data(spike_in_se)
data(spike_in_de_res)
plot_fold_changes_spiked(spike_in_se, spike_in_de_res,
                        condition = "Condition", ain = NULL,
                        comparisons = NULL)
```

---

|              |                                                                                                                           |
|--------------|---------------------------------------------------------------------------------------------------------------------------|
| plot_heatmap | <i>Plot a heatmap of the sample intensities with optional column annotations for a selection of normalization methods</i> |
|--------------|---------------------------------------------------------------------------------------------------------------------------|

---

**Description**

Plot a heatmap of the sample intensities with optional column annotations for a selection of normalization methods

**Usage**

```
plot_heatmap(
  se,
  ain = NULL,
  color_by = c("Group", "Pool"),
  label_by = NULL,
  only_refs = FALSE
)
```

**Arguments**

|           |                                                                                                                                                                         |
|-----------|-------------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| se        | SummarizedExperiment containing all necessary information of the proteomics data set                                                                                    |
| ain       | Vector of strings which assay should be used as input (default NULL). If NULL then all normalization of the se object are plotted next to each other.                   |
| color_by  | Vector of strings specifying the columns to color the samples (If NULL, the condition column of the SummarizedExperiment object is used. If "No", no color bars added.) |
| label_by  | String specifying the column in the metadata used to label the samples for the UpSet plot                                                                               |
| only_refs | Logical, if TRUE, only reference samples (ComRef) are included in the plot                                                                                              |

**Value**

list of ggplot objects

**Examples**

```
data(tuberculosis_TMT_se)
plot_heatmap(tuberculosis_TMT_se, ain = c("log2"), color_by = NULL,
             label_by = NULL, only_refs = FALSE)
```

---

|                 |                              |
|-----------------|------------------------------|
| plot_heatmap_DE | <i>Heatmap of DE results</i> |
|-----------------|------------------------------|

---

**Description**

Heatmap of DE results

**Usage**

```
plot_heatmap_DE(
  se,
  de_res,
  ain,
  comparison,
  condition = NULL,
  label_by = NULL,
  pvalue_column = "adj.P.Val",
  col_vector = NULL
)
```

**Arguments**

|        |                                                                                                                             |
|--------|-----------------------------------------------------------------------------------------------------------------------------|
| se     | SummarizedExperiment containing all necessary information of the proteomics data set (including the normalized intensities) |
| de_res | data table resulting of run_DE                                                                                              |
| ain    | Vector of strings of normalization methods to visualize (must be valid normalization methods saved in de_res)               |

|               |                                                                                                                                                                  |
|---------------|------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| comparison    | String of comparison (must be a valid comparison saved in de_res)                                                                                                |
| condition     | column name of condition (if NULL, condition saved in SummarizedExperiment will be taken)                                                                        |
| label_by      | String specifying the column to label the samples (If NULL, the labels column of the SummarizedExperiment object is used. If "No", no labeling of samples done.) |
| pvalue_column | column name of p-values in de_res                                                                                                                                |
| col_vector    | Vector of colors to use for the heatmap. If NULL, default colors are used.                                                                                       |

**Value**

list of ComplexHeatmaps for each method

**Examples**

```
data(tuberculosis_TMT_se)
data(tuberculosis_TMT_de_res)
plot_heatmap_DE(tuberculosis_TMT_se, tuberculosis_TMT_de_res, ain = NULL,
                comparison = "PTB-HC",
                condition = NULL, label_by = NULL,
                pvalue_column = "adj.P.Val", col_vector = NULL)
```

---

plot\_histogram\_spiked *Plot histogram of the spike-in and background protein intensities per condition.*

---

**Description**

Plot histogram of the spike-in and background protein intensities per condition.

**Usage**

```
plot_histogram_spiked(se, condition = NULL)
```

**Arguments**

|           |                                                                                           |
|-----------|-------------------------------------------------------------------------------------------|
| se        | SummarizedExperiment containing all necessary information of the proteomics data set      |
| condition | column name of condition (if NULL, condition saved in SummarizedExperiment will be taken) |

**Value**

ggplot object

**Examples**

```
data(spike_in_se)
plot_histogram_spiked(spike_in_se, condition = NULL)
```

---

plot\_identified\_spiked\_proteins

*Plot number of identified spike-in proteins per sample.*


---

### Description

Plot number of identified spike-in proteins per sample.

### Usage

```
plot_identified_spiked_proteins(se, color_by = NULL, label_by = NULL)
```

### Arguments

|          |                                                                                                                                                                    |
|----------|--------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| se       | SummarizedExperiment containing all necessary information of the proteomics data set                                                                               |
| color_by | String specifying the column to color the samples (If NULL, the condition column of the SummarizedExperiment object is used. If "No", no color bar added.)         |
| label_by | String specifying the column to label the samples (If NULL, the labels column of the SummarizedExperiment object is used. If "No", no labeling of samples done.)#' |

### Value

ggplot object

### Examples

```
data(spike_in_se)
plot_identified_spiked_proteins(spike_in_se, color_by = NULL,
                                label_by = NULL)
```

---

plot\_intersection\_enrichment

*Functional enrichment analysis for analyzing the DE results of different normalization methods and biologically interpreting the results*


---

### Description

Functional enrichment analysis for analyzing the DE results of different normalization methods and biologically interpreting the results

**Usage**

```
plot_intersection_enrichment(
  se,
  de_res,
  comparison,
  ain = NULL,
  id_column = "Gene.Names",
  organism = "hsapiens",
  source = "KEGG",
  signif_thr = 0.05
)
```

**Arguments**

|            |                                                                                                               |
|------------|---------------------------------------------------------------------------------------------------------------|
| se         | SummarizedExperiment containing all necessary information of the proteomics data set                          |
| de_res     | data table resulting of run_DE                                                                                |
| comparison | String of comparison (must be a valid comparison saved in de_res)                                             |
| ain        | Vector of strings of normalization methods to visualize (must be valid normalization methods saved in de_res) |
| id_column  | String specifying the column of the rowData of the SummarizedExperiment object which includes the gene names  |
| organism   | Organism name (gprofiler parameter)                                                                           |
| source     | Data source to use (gprofiler parameter, example: KEGG)                                                       |
| signif_thr | Significance threshold                                                                                        |

**Value**

list of ggplot objects

**Examples**

```
data(tuberculosis_TMT_se)
data(tuberculosis_TMT_de_res)
plot_intersection_enrichment(tuberculosis_TMT_se, tuberculosis_TMT_de_res,
  ain = c("IRS_on_Median", "IRS_on_RobNorm", "RobNorm"),
  comparison = "Rx-TBL", id_column = "Gene.Names",
  organism = "hsapiens", source = "GO:BP", signif_thr = 0.05)
```

---

plot\_intragroup\_correlation

*Plot intragroup correlation of the normalized data*

---

**Description**

Plot intragroup correlation of the normalized data

**Usage**

```
plot_intragroup_correlation(
  se,
  ain = NULL,
  condition = NULL,
  method = "pearson"
)
```

**Arguments**

|           |                                                                                                                                                       |
|-----------|-------------------------------------------------------------------------------------------------------------------------------------------------------|
| se        | SummarizedExperiment containing all necessary information of the proteomics data set                                                                  |
| ain       | Vector of strings which assay should be used as input (default NULL). If NULL then all normalization of the se object are plotted next to each other. |
| condition | column name of condition (if NULL, condition saved in SummarizedExperiment will be taken)                                                             |
| method    | String specifying the method for correlation calculation (pearson, spearman or kendall)                                                               |

**Value**

ggplot object (boxplot)

**Examples**

```
data(tuberculosis_TMT_se)
plot_intragroup_correlation(tuberculosis_TMT_se, ain = NULL,
                           condition = NULL, method = "pearson")
```

---

|                     |                                                                                     |
|---------------------|-------------------------------------------------------------------------------------|
| plot_intragroup_PCV | <i>Plot intragroup pooled coefficient of variation (PCV) of the normalized data</i> |
|---------------------|-------------------------------------------------------------------------------------|

---

**Description**

Plot intragroup pooled coefficient of variation (PCV) of the normalized data

**Usage**

```
plot_intragroup_PCV(se, ain = NULL, condition = NULL, diff = FALSE)
```

**Arguments**

|           |                                                                                                                                                       |
|-----------|-------------------------------------------------------------------------------------------------------------------------------------------------------|
| se        | SummarizedExperiment containing all necessary information of the proteomics data set                                                                  |
| ain       | Vector of strings which assay should be used as input (default NULL). If NULL then all normalization of the se object are plotted next to each other. |
| condition | Column name of condition (if NULL, condition saved in SummarizedExperiment will be taken)                                                             |

**diff** Boolean indicating whether to visualize the reduction of intragroup variation (PCV) compared to "log" (TRUE) or a normal boxplot of intragroup variation (PCV) for each normalization method (FALSE).

### Value

ggplot object (boxplot)

### Examples

```
data(tuberculosis_TMT_se)
plot_intragroup_PCV(tuberculosis_TMT_se, ain = NULL,
                    condition = NULL, diff = FALSE)
```

---

|                     |                                                                                 |
|---------------------|---------------------------------------------------------------------------------|
| plot_intragroup_PEV | <i>Plot intragroup pooled estimate of variance (PEV) of the normalized data</i> |
|---------------------|---------------------------------------------------------------------------------|

---

### Description

Plot intragroup pooled estimate of variance (PEV) of the normalized data

### Usage

```
plot_intragroup_PEV(se, ain = NULL, condition = NULL, diff = FALSE)
```

### Arguments

|                  |                                                                                                                                                                                                       |
|------------------|-------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| <b>se</b>        | SummarizedExperiment containing all necessary information of the proteomics data set                                                                                                                  |
| <b>ain</b>       | Vector of strings which assay should be used as input (default NULL). If NULL then all normalization of the se object are plotted next to each other.                                                 |
| <b>condition</b> | column name of condition (if NULL, condition saved in SummarizedExperiment will be taken)                                                                                                             |
| <b>diff</b>      | Boolean indicating whether to visualize the reduction of intragroup variation (PEV) compared to "log" (TRUE) or a normal boxplot of intragroup variation (PEV) for each normalization method (FALSE). |

### Value

ggplot object (boxplot)

### Examples

```
data(tuberculosis_TMT_se)
plot_intragroup_PEV(tuberculosis_TMT_se, ain = NULL,
                    condition = NULL, diff = FALSE)
```

---

|                     |                                                                                       |
|---------------------|---------------------------------------------------------------------------------------|
| plot_intragroup_PMA | <i>Plot intragroup pooled median absolute deviation (PMAD) of the normalized data</i> |
|---------------------|---------------------------------------------------------------------------------------|

---

### Description

Plot intragroup pooled median absolute deviation (PMAD) of the normalized data

### Usage

```
plot_intragroup_PMA(se, ain = NULL, condition = NULL, diff = FALSE)
```

### Arguments

|           |                                                                                                                                                                                                         |
|-----------|---------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| se        | SummarizedExperiment containing all necessary information of the proteomics data set                                                                                                                    |
| ain       | Vector of strings which assay should be used as input (default NULL). If NULL then all normalization of the se object are plotted next to each other.                                                   |
| condition | column name of condition (if NULL, condition saved in SummarizedExperiment will be taken)                                                                                                               |
| diff      | Boolean indicating whether to visualize the reduction of intragroup variation (PMAD) compared to "log" (TRUE) or a normal boxplot of intragroup variation (PMAD) for each normalization method (FALSE). |

### Value

ggplot object (boxplot)

### Examples

```
data(tuberculosis_TMT_se)
plot_intragroup_PMA(tuberculosis_TMT_se, ain = NULL,
                    condition = NULL, diff = FALSE)
```

---

|                      |                                                                                         |
|----------------------|-----------------------------------------------------------------------------------------|
| plot_jaccard_heatmap | <i>Jaccard similarity heatmap of DE proteins of the different normalization methods</i> |
|----------------------|-----------------------------------------------------------------------------------------|

---

### Description

Jaccard similarity heatmap of DE proteins of the different normalization methods

### Usage

```
plot_jaccard_heatmap(
  de_res,
  ain = NULL,
  comparisons = NULL,
  plot_type = "single"
)
```

**Arguments**

|             |                                                                                                                                                                    |
|-------------|--------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| de_res      | data table resulting of run_DE                                                                                                                                     |
| ain         | Vector of strings of normalization methods to visualize (must be valid normalization methods saved in de_res)                                                      |
| comparisons | Vector of comparisons (must be valid comparisons saved in de_res)                                                                                                  |
| plot_type   | String indicating whether to plot a single plot per comparison ("single"), facet by comparison ("facet_comp"), or include all comparisons in a single plot ("all") |

**Value**

ggplot object (list of objects if plot\_type == "single")

**Examples**

```
data(tuberculosis_TMT_de_res)
plot_jaccard_heatmap(tuberculosis_TMT_de_res, ain = NULL,
                     comparisons = NULL, plot_type = "all")
```

---

plot\_logFC\_thresholds\_spiked

*Line plot of number of true and false positives when applying different logFC thresholds*

---

**Description**

Line plot of number of true and false positives when applying different logFC thresholds

**Usage**

```
plot_logFC_thresholds_spiked(
  se,
  de_res,
  condition,
  ain = NULL,
  comparisons = NULL,
  nrow = 2,
  alpha = 0.05
)
```

**Arguments**

|           |                                                                                                              |
|-----------|--------------------------------------------------------------------------------------------------------------|
| se        | SummarizedExperiment containing all necessary information of the proteomics data set                         |
| de_res    | data table resulting of run_DE                                                                               |
| condition | column name of condition (if NULL, condition saved in SummarizedExperiment will be taken)                    |
| ain       | Vector of strings of normalization methods to visualize (must be valid normalization methods saved in stats) |

|             |                                                                  |
|-------------|------------------------------------------------------------------|
| comparisons | Vector of comparisons (must be valid comparisons saved in stats) |
| nrow        | number of rows for facet wrap                                    |
| alpha       | threshold for adjusted p-values                                  |

**Value**

list of ggplot objects

**Examples**

```
data(spike_in_se)
data(spike_in_de_res)
plot_logFC_thresholds_spiked(spike_in_se, spike_in_de_res,
                             condition = "Condition", ain = NULL,
                             comparisons = NULL, nrow = 2, alpha = 0.05)
```

---

plot\_markers\_boxplots *Boxplots of intensities of specific markers*

---

**Description**

Boxplots of intensities of specific markers

**Usage**

```
plot_markers_boxplots(
  se,
  markers,
  ain = NULL,
  id_column = "Protein.IDs",
  color_by = NULL,
  shape_by = NULL,
  facet_norm = TRUE,
  facet_marker = FALSE
)
```

**Arguments**

|           |                                                                                                                                                            |
|-----------|------------------------------------------------------------------------------------------------------------------------------------------------------------|
| se        | SummarizedExperiment containing all necessary information of the proteomics data set                                                                       |
| markers   | Vector of the IDs of the markers to plot                                                                                                                   |
| ain       | Vector of strings of normalization methods to visualize (must be valid normalization methods saved in de_res)                                              |
| id_column | String specifying the column of the rowData of the SummarizedExperiment object which includes the IDs of the markers                                       |
| color_by  | String specifying the column to color the samples (If NULL, the condition column of the SummarizedExperiment object is used. If "No", no color bar added.) |
| shape_by  | String specifying the column to shape the samples (If NULL or "No", no shaping of samples is done.)                                                        |

|              |                                                                                                            |
|--------------|------------------------------------------------------------------------------------------------------------|
| facet_norm   | Boolean indicating whether to facet by normalization method (TRUE) or not (FALSE)                          |
| facet_marker | Boolean indicating whether to facet by comparison (TRUE) or not (FALSE). Only valid if facet_norm = FALSE. |

**Value**

ggplot object

**Examples**

```
data(tuberculosis_TMT_se)
plot_markers_boxplots(tuberculosis_TMT_se, markers = c("Q7Z7F0", "Q13790"),
  ain = c("log2"), id_column = "Protein.IDs",
  color_by = NULL,
  shape_by = "Pool",
  facet_norm = FALSE,
  facet_marker = TRUE)
```

---

plot\_NA\_density

*Plot the intensity distribution of proteins with and without NAs*

---

**Description**

Plot the intensity distribution of proteins with and without NAs

**Usage**

```
plot_NA_density(se)
```

**Arguments**

|    |                                                                                      |
|----|--------------------------------------------------------------------------------------|
| se | SummarizedExperiment containing all necessary information of the proteomics data set |
|----|--------------------------------------------------------------------------------------|

**Value**

ggplot object

**Examples**

```
data(tuberculosis_TMT_se)
plot_NA_density(tuberculosis_TMT_se)
```

---

|                   |                                                                                                        |
|-------------------|--------------------------------------------------------------------------------------------------------|
| plot_NA_frequency | <i>Plot protein identification overlap (x = identified in number of Samples, y=number of proteins)</i> |
|-------------------|--------------------------------------------------------------------------------------------------------|

---

### Description

Plot protein identification overlap (x = identified in number of Samples, y=number of proteins)

### Usage

```
plot_NA_frequency(se)
```

### Arguments

|    |                                                                                      |
|----|--------------------------------------------------------------------------------------|
| se | SummarizedExperiment containing all necessary information of the proteomics data set |
|----|--------------------------------------------------------------------------------------|

### Value

ggplot object

### Examples

```
data(tuberculosis_TMT_se)
plot_NA_frequency(tuberculosis_TMT_se)
```

---

|                 |                                       |
|-----------------|---------------------------------------|
| plot_NA_heatmap | <i>Plot heatmap of the NA pattern</i> |
|-----------------|---------------------------------------|

---

### Description

Plot heatmap of the NA pattern

### Usage

```
plot_NA_heatmap(
  se,
  color_by = NULL,
  label_by = NULL,
  cluster_samples = TRUE,
  cluster_proteins = TRUE,
  show_row_dend = TRUE,
  show_column_dend = FALSE,
  col_vector = NULL
)
```

**Arguments**

|                  |                                                                                                                                                                  |
|------------------|------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| se               | SummarizedExperiment containing all necessary information of the proteomics data set                                                                             |
| color_by         | String specifying the column to color the samples (If NULL, the condition column of the SummarizedExperiment object is used. If "No", no color bar added.)       |
| label_by         | String specifying the column to label the samples (If NULL, the labels column of the SummarizedExperiment object is used. If "No", no labeling of samples done.) |
| cluster_samples  | Boolean. TRUE if samples should be clustered, else FALSE.                                                                                                        |
| cluster_proteins | Boolean. TRUE if proteins should be clustered, else FALSE.                                                                                                       |
| show_row_dend    | Boolean. TRUE if row dendrogram should be shown.                                                                                                                 |
| show_column_dend | Boolean. TRUE if column dendrogram should be shown.                                                                                                              |
| col_vector       | Vector of colors for the color bar. If NULL, default colors are used.                                                                                            |

**Value**

ComplexHeatmap plot (only showing proteins with at least one missing value)

**Examples**

```
data(tuberculosis_TMT_se)
plot_NA_heatmap(tuberculosis_TMT_se, color_by = NULL,
                label_by = NULL, cluster_samples = TRUE,
                cluster_proteins = TRUE, show_row_dend = TRUE,
                show_column_dend = FALSE,
                col_vector = NULL)
```

---

plot\_nr\_prot\_samples    *Plot number of non-zero proteins per sample*

---

**Description**

Plot number of non-zero proteins per sample

**Usage**

```
plot_nr_prot_samples(se, ain = "raw", color_by = NULL, label_by = NULL)
```

**Arguments**

|          |                                                                                                                                                                  |
|----------|------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| se       | SummarizedExperiment containing all necessary information of the proteomics data set                                                                             |
| ain      | String which data type should be used (default raw)                                                                                                              |
| color_by | String specifying the column to color the samples (If NULL, the condition column of the SummarizedExperiment object is used. If "No", no color bar added.)       |
| label_by | String specifying the column to label the samples (If NULL, the labels column of the SummarizedExperiment object is used. If "No", no labeling of samples done.) |

**Value**

ggplot object

**Examples**

```
data(tuberculosis_TMT_se)
plot_nr_prot_samples(tuberculosis_TMT_se, ain="raw", color_by = "Group",
                     label_by = "Label")
```

---

plot\_overview\_DE\_bar    *Overview plots of DE results*

---

**Description**

Overview plots of DE results

**Usage**

```
plot_overview_DE_bar(
  de_res,
  ain = NULL,
  comparisons = NULL,
  plot_type = "single"
)
```

**Arguments**

|             |                                                                                                                                                                                                                                                                     |
|-------------|---------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| de_res      | data table resulting of run_DE                                                                                                                                                                                                                                      |
| ain         | Vector of strings of normalization methods to visualize (must be valid normalization methods saved in de_res)                                                                                                                                                       |
| comparisons | Vector of comparisons (must be valid comparisons saved in de_res)                                                                                                                                                                                                   |
| plot_type   | String indicating whether to plot a single plot per comparison ("single"), facet by comparison ("facet_comp"), stack the number of DE per comparison ("stacked"), or stack the number of DE per comparison but facet by up- and down-regulated ("facet_regulation") |

**Value**

list of ggplot objects or single object if plot\_type = facet or stacked

**Examples**

```
data(tuberculosis_TMT_de_res)
plot_overview_DE_bar(tuberculosis_TMT_de_res, ain = NULL, comparisons = NULL,
                     plot_type = "facet_regulation")
```

---

plot\_overview\_DE\_tile *Overview heatmap plot of DE results*

---

### Description

Overview heatmap plot of DE results

### Usage

```
plot_overview_DE_tile(de_res, ain = NULL, comparisons = NULL)
```

### Arguments

|             |                                                                                                               |
|-------------|---------------------------------------------------------------------------------------------------------------|
| de_res      | data table resulting of run_DE                                                                                |
| ain         | Vector of strings of normalization methods to visualize (must be valid normalization methods saved in de_res) |
| comparisons | Vector of comparisons (must be valid comparisons saved in de_res)                                             |

### Value

ggplot object

### Examples

```
data(tuberculosis_TMT_de_res)
plot_overview_DE_tile(tuberculosis_TMT_de_res, ain = NULL,
                      comparisons = NULL)
```

---

plot\_PCA *PCA plot of the normalized data*

---

### Description

PCA plot of the normalized data

### Usage

```
plot_PCA(
  se,
  ain = NULL,
  color_by = NULL,
  label_by = NULL,
  shape_by = NULL,
  facet_norm = TRUE,
  facet_by = NULL,
  ellipse = FALSE,
  ncol = 3
)
```

**Arguments**

|            |                                                                                                                                                                                                                                                                                                                                                         |
|------------|---------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| se         | SummarizedExperiment containing all necessary information of the proteomics data set                                                                                                                                                                                                                                                                    |
| ain        | Vector of strings which assay should be used as input (default NULL). If NULL then all normalization of the se object are plotted next to each other.                                                                                                                                                                                                   |
| color_by   | String specifying the column to color the samples (If NULL, the condition column of the SummarizedExperiment object is used. If "No", no color bar added.)                                                                                                                                                                                              |
| label_by   | String specifying the column to label the samples (If NULL, the labels column of the SummarizedExperiment object is used. If "No", no labeling of samples done.)                                                                                                                                                                                        |
| shape_by   | String specifying the column to shape the samples (If NULL or "No", no shaping of samples is done.)                                                                                                                                                                                                                                                     |
| facet_norm | Boolean specifying whether to facet by normalization methods (default TRUE). If FALSE, list of plots of the different normalized data is returned. However, then you can also facet by any column of the metadata.                                                                                                                                      |
| facet_by   | String specifying the column to facet the samples (If facet = FALSE, the plot will not be faceted by the normalization methods, but instead a list of plots of each normalization method is returned. Then, the PCA plot can be faceted by any column of the metadata, for instance by "Batch". If facet_by is NULL or "No", no faceting is performed.) |
| ellipse    | Boolean to indicate if ellipses should be drawn                                                                                                                                                                                                                                                                                                         |
| ncol       | Number of columns in plot (for faceting)                                                                                                                                                                                                                                                                                                                |

**Value**

if facet\_norm = TRUE, ggplot object, else list of ggplot objects

**Examples**

```
data(tuberculosis_TMT_se)
plot_PCA(tuberculosis_TMT_se, ain = NULL, color_by = NULL, label_by = NULL,
        shape_by = "Pool",
        facet_norm = TRUE, ncol = 3)
plot_PCA(tuberculosis_TMT_se, ain = c("IRS_on_RobNorm"), color_by = "Group",
        label_by = "Label", facet_norm = FALSE, facet_by = "Pool")
```

---

|                      |                                                                                                                                                        |
|----------------------|--------------------------------------------------------------------------------------------------------------------------------------------------------|
| plot_profiles_spiked | <i>Plot profiles of the spike-in and background proteins using the log2 average protein intensities as a function of the different concentrations.</i> |
|----------------------|--------------------------------------------------------------------------------------------------------------------------------------------------------|

---

**Description**

Plot profiles of the spike-in and background proteins using the log2 average protein intensities as a function of the different concentrations.

**Usage**

```
plot_profiles_spiked(se, xlab = "Concentration")
```

**Arguments**

|      |                                                                                      |
|------|--------------------------------------------------------------------------------------|
| se   | SummarizedExperiment containing all necessary information of the proteomics data set |
| xlab | String for the x-label of the plot                                                   |

**Value**

ggplot object

**Examples**

```
data(spike_in_se)
plot_profiles_spiked(spike_in_se, xlab = "Concentration")
```

---

|                     |                                                                                                                                                                                                                              |
|---------------------|------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| plot_pvalues_spiked | <i>Boxplot of p-values of spike-in and background proteins for specific normalization methods and comparisons. The ground truth (calculated based on the concentrations of the spike-ins) is shown as a horizontal line.</i> |
|---------------------|------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|

---

**Description**

Boxplot of p-values of spike-in and background proteins for specific normalization methods and comparisons. The ground truth (calculated based on the concentrations of the spike-ins) is shown as a horizontal line.

**Usage**

```
plot_pvalues_spiked(se, de_res, ain = NULL, comparisons = NULL)
```

**Arguments**

|             |                                                                                                              |
|-------------|--------------------------------------------------------------------------------------------------------------|
| se          | SummarizedExperiment containing all necessary information of the proteomics data set                         |
| de_res      | data table resulting of run_DE                                                                               |
| ain         | Vector of strings of normalization methods to visualize (must be valid normalization methods saved in stats) |
| comparisons | Vector of comparisons (must be valid comparisons saved in stats)                                             |

**Value**

ggplot object

**Examples**

```
data(spike_in_se)
data(spike_in_de_res)
plot_pvalues_spiked(spike_in_se, spike_in_de_res, ain = NULL,
                    comparisons = NULL)
```

---

|                     |                                                                                                                  |
|---------------------|------------------------------------------------------------------------------------------------------------------|
| plot_ROC_AUC_spiked | <i>Plot ROC curve and barplot of AUC values for each method for a specific comparison or for all comparisons</i> |
|---------------------|------------------------------------------------------------------------------------------------------------------|

---

**Description**

Plot ROC curve and barplot of AUC values for each method for a specific comparison or for all comparisons

**Usage**

```
plot_ROC_AUC_spiked(se, de_res, ain = NULL, comparisons = NULL)
```

**Arguments**

|             |                                                                                                              |
|-------------|--------------------------------------------------------------------------------------------------------------|
| se          | SummarizedExperiment containing all necessary information of the proteomics data set                         |
| de_res      | data table resulting of run_DE                                                                               |
| ain         | Vector of strings of normalization methods to visualize (must be valid normalization methods saved in stats) |
| comparisons | Vector of comparisons (must be valid comparisons saved in stats)                                             |

**Value**

list of ggplot objects

**Examples**

```
data(spike_in_se)
data(spike_in_de_res)
plot_ROC_AUC_spiked(spike_in_se, spike_in_de_res)
```

---

|                           |                                                              |
|---------------------------|--------------------------------------------------------------|
| plot_stats_spiked_heatmap | <i>Heatmap of performance metrics for spike-in data sets</i> |
|---------------------------|--------------------------------------------------------------|

---

**Description**

Heatmap of performance metrics for spike-in data sets

**Usage**

```
plot_stats_spiked_heatmap(
  stats,
  ain = NULL,
  comparisons = NULL,
  metrics = c("Accuracy", "Precision", "F1Score")
)
```

**Arguments**

|             |                                                                                                              |
|-------------|--------------------------------------------------------------------------------------------------------------|
| stats       | data table with multiple metrics of the DE results (resulting of get_spiked_stats_DE)                        |
| ain         | Vector of strings of normalization methods to visualize (must be valid normalization methods saved in stats) |
| comparisons | Vector of comparisons (must be valid comparisons saved in stats)                                             |
| metrics     | vector of Strings specifying the metrics (must be colnames of stats)                                         |

**Value**

ggplot object

**Examples**

```
data(spike_in_se)
data(spike_in_de_res)
stats <- get_spiked_stats_DE(spike_in_se, spike_in_de_res)
plot_stats_spiked_heatmap(stats, ain = NULL, comparisons = NULL,
                           metrics = c("F1Score", "Accuracy"))
```

---

plot\_tot\_int\_samples    *Plot total protein intensity per sample*

---

**Description**

Plot total protein intensity per sample

**Usage**

```
plot_tot_int_samples(se, ain = "raw", color_by = NULL, label_by = NULL)
```

**Arguments**

|          |                                                                                                                                                                  |
|----------|------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| se       | SummarizedExperiment containing all necessary information of the proteomics data set                                                                             |
| ain      | String which data type should be used (default raw)                                                                                                              |
| color_by | String specifying the column to color the samples (If NULL, the condition column of the SummarizedExperiment object is used. If "No", no color bar added.)       |
| label_by | String specifying the column to label the samples (If NULL, the labels column of the SummarizedExperiment object is used. If "No", no labeling of samples done.) |

**Value**

list of a ggplot object and the dataframe of outliers

**Examples**

```
data(tuberculosis_TMT_se)
plot_tot_int_samples(tuberculosis_TMT_se, ain="raw", color_by = NULL,
                      label_by = NULL)
```

---

|                       |                                                                                               |
|-----------------------|-----------------------------------------------------------------------------------------------|
| plot_TP_FP_spiked_bar | <i>Barplot of true and false positives for specific comparisons and normalization methods</i> |
|-----------------------|-----------------------------------------------------------------------------------------------|

---

### Description

Barplot of true and false positives for specific comparisons and normalization methods

### Usage

```
plot_TP_FP_spiked_bar(stats, ain = NULL, comparisons = NULL)
```

### Arguments

|             |                                                                                                              |
|-------------|--------------------------------------------------------------------------------------------------------------|
| stats       | data table with multiple metrics of the DE results (resulting of get_spiked_stats_DE)                        |
| ain         | Vector of strings of normalization methods to visualize (must be valid normalization methods saved in stats) |
| comparisons | Vector of comparisons (must be valid comparisons saved in stats)                                             |

### Value

ggplot object (barplot)

### Examples

```
data(spike_in_se)
data(spike_in_de_res)
stats <- get_spiked_stats_DE(spike_in_se, spike_in_de_res)
plot_TP_FP_spiked_bar(stats, ain = NULL, comparisons = NULL)
```

---

|                       |                                                                                               |
|-----------------------|-----------------------------------------------------------------------------------------------|
| plot_TP_FP_spiked_box | <i>Boxplot of true and false positives for specific comparisons and normalization methods</i> |
|-----------------------|-----------------------------------------------------------------------------------------------|

---

### Description

Boxplot of true and false positives for specific comparisons and normalization methods

### Usage

```
plot_TP_FP_spiked_box(stats, ain = NULL, comparisons = NULL)
```

### Arguments

|             |                                                                                                              |
|-------------|--------------------------------------------------------------------------------------------------------------|
| stats       | data table with multiple metrics of the DE results (resulting of get_spiked_stats_DE)                        |
| ain         | Vector of strings of normalization methods to visualize (must be valid normalization methods saved in stats) |
| comparisons | Vector of comparisons (must be valid comparisons saved in stats)                                             |

**Value**

ggplot object (barplot)

**Examples**

```
data(spike_in_se)
data(spike_in_de_res)
stats <- get_spiked_stats_DE(spike_in_se, spike_in_de_res)
plot_TP_FP_spiked_box(stats, ain = NULL, comparisons = NULL)
```

---

plot\_TP\_FP\_spiked\_scatter

*Scatterplot of true positives and false positives (median with errorbars as Q1, and Q3) for all comparisons*

---

**Description**

Scatterplot of true positives and false positives (median with errorbars as Q1, and Q3) for all comparisons

**Usage**

```
plot_TP_FP_spiked_scatter(stats, ain = NULL, comparisons = NULL)
```

**Arguments**

|             |                                                                                                              |
|-------------|--------------------------------------------------------------------------------------------------------------|
| stats       | data table with multiple metrics of the DE results (resulting of get_spiked_stats_DE)                        |
| ain         | Vector of strings of normalization methods to visualize (must be valid normalization methods saved in stats) |
| comparisons | Vector of comparisons (must be valid comparisons saved in stats)                                             |

**Value**

ggplot object

**Examples**

```
data(spike_in_se)
data(spike_in_de_res)
stats <- get_spiked_stats_DE(spike_in_se, spike_in_de_res)
plot_TP_FP_spiked_scatter(stats, ain = NULL, comparisons = NULL)
```

plot\_upset

*Create an UpSet Plot from SummarizedExperiment Data***Description**

This function generates an UpSet plot from a given SummarizedExperiment object. It allows for the visualization of overlaps between sets defined by a specific column in the metadata. The function supports subsetting to reference samples and customizable color mapping.

**Usage**

```
plot_upset(
  se,
  color_by = NULL,
  label_by = NULL,
  mb.ratio = c(0.7, 0.3),
  only_refs = FALSE
)
```

**Arguments**

|           |                                                                                                                               |
|-----------|-------------------------------------------------------------------------------------------------------------------------------|
| se        | SummarizedExperiment containing all necessary information of the proteomics data set                                          |
| color_by  | String specifying the column to color the samples (If NULL, the condition column of the SummarizedExperiment object is used.) |
| label_by  | String specifying the column in the metadata used to label the samples for the UpSet plot                                     |
| mb.ratio  | A numeric vector of length 2, specifying the barplot and matrix area ratios                                                   |
| only_refs | Logical, if TRUE, only reference samples (ComRef) are included in the plot                                                    |

**Value**

ggplot object

**Examples**

```
data(tuberculosis_TMT_se)
plot_upset(tuberculosis_TMT_se, color_by = NULL, label_by = NULL,
  mb.ratio = c(0.7, 0.3), only_refs = FALSE)
```

---

|               |                                                                         |
|---------------|-------------------------------------------------------------------------|
| plot_upset_DE | <i>Upset plots of DE results of the different normalization methods</i> |
|---------------|-------------------------------------------------------------------------|

---

**Description**

Upset plots of DE results of the different normalization methods

**Usage**

```
plot_upset_DE(
  de_res,
  ain = NULL,
  comparisons = NULL,
  min_degree = 2,
  plot_type = "single"
)
```

**Arguments**

|             |                                                                                                                                |
|-------------|--------------------------------------------------------------------------------------------------------------------------------|
| de_res      | data table resulting of run_DE                                                                                                 |
| ain         | Vector of strings of normalization methods to visualize (must be valid normalization methods saved in de_res)                  |
| comparisons | Vector of comparisons (must be valid comparisons saved in de_res)                                                              |
| min_degree  | Minimal degree of an intersection for it to be included                                                                        |
| plot_type   | String indicating whether to plot a single plot per comparison ("single") or stack the number of DE per comparison ("stacked") |

**Value**

list of plots and intersection tables (split by comparison if plot\_type == "single")

**Examples**

```
data(tuberculosis_TMT_de_res)
plot_upset_DE(tuberculosis_TMT_de_res,
  ain = c("IRS_on_RobNorm", "IRS_on_Median"),
  comparisons = NULL, min_degree = 2,
  plot_type = "stacked")
```

---

|                 |                                    |
|-----------------|------------------------------------|
| plot_volcano_DE | <i>Volcano plots of DE results</i> |
|-----------------|------------------------------------|

---

**Description**

Volcano plots of DE results

**Usage**

```
plot_volcano_DE(
  de_res,
  ain = NULL,
  comparisons = NULL,
  facet_norm = TRUE,
  facet_comparison = FALSE
)
```

**Arguments**

|                  |                                                                                                               |
|------------------|---------------------------------------------------------------------------------------------------------------|
| de_res           | data table resulting of run_DE                                                                                |
| ain              | Vector of strings of normalization methods to visualize (must be valid normalization methods saved in de_res) |
| comparisons      | Vector of comparisons (must be valid comparisons saved in de_res)                                             |
| facet_norm       | Boolean indicating whether to facet by normalization method (TRUE) or not (FALSE)                             |
| facet_comparison | Boolean indicating whether to facet by comparison (TRUE) or not (FALSE). Only valid if facet_norm = FALSE.    |

**Value**

list of ggplot objects

**Examples**

```
data(tuberculosis_TMT_de_res)
plot_volcano_DE(tuberculosis_TMT_de_res, ain = NULL,
  comparisons = NULL, facet_norm = TRUE,
  facet_comparison = FALSE)
```

---

quantileNorm

*Quantile Normalization of preprocessCore package.*


---

**Description**

Forces distributions of the samples to be the same on the basis of the quantiles of the samples by replacing each protein of a sample with the mean of the corresponding quantile. Log2-scaled data should be taken as input (on\_raw = FALSE)

**Usage**

```
quantileNorm(se, ain = "log2", aout = "Quantile", on_raw = FALSE)
```

**Arguments**

|        |                                                                                         |
|--------|-----------------------------------------------------------------------------------------|
| se     | SummarizedExperiment containing all necessary information of the proteomic dataset      |
| ain    | String which assay should be used as input                                              |
| aout   | String which assay should be used to save normalized data                               |
| on_raw | Boolean specifying whether normalization should be performed on raw or log2-scaled data |

**Value**

SummarizedExperiment containing the quantile normalized data as assay (on log2 scale)

**See Also**

`normalize.quantiles()`

**Examples**

```
data(tuberculosis_TMT_se)
tuberculosis_TMT_se <- quantileNorm(tuberculosis_TMT_se, ain = "log2",
                                   aout = "Quantile", on_raw = FALSE)
```

---

|                   |                                             |
|-------------------|---------------------------------------------|
| readPRONE_example | <i>Helper function to read example data</i> |
|-------------------|---------------------------------------------|

---

**Description**

Helper function to read example data

**Usage**

```
readPRONE_example(path = NULL)
```

**Arguments**

|      |                                                                         |
|------|-------------------------------------------------------------------------|
| path | NULL to get all example data set files, otherwise specify the file name |
|------|-------------------------------------------------------------------------|

**Value**

If path=NULL a character vector with the file names, otherwise the path to the specific file

**Examples**

```
readPRONE_example()
```

---

`remove_assays_from_SE` *Remove normalization assays from a SummarizedExperiment object*

---

### Description

Remove normalization assays from a SummarizedExperiment object

### Usage

```
remove_assays_from_SE(se, assays_to_remove)
```

### Arguments

`se` SummarizedExperiment containing all necessary information of the proteomics data set

`assays_to_remove` Character vector of assay names to remove from the SummarizedExperiment object

### Value

SummarizedExperiment object with the normalization assays removed

### Examples

```
data(tuberculosis_TMT_se)
tuberculosis_TMT_se <- remove_assays_from_SE(tuberculosis_TMT_se,
                                             assays_to_remove = c("IRS_on_RobNorm"))
```

---

`remove_POMA_outliers` *Remove outliers samples detected by the detect\_outliers\_POMA function*

---

### Description

Remove outliers samples detected by the detect\_outliers\_POMA function

### Usage

```
remove_POMA_outliers(se, poma_res_outliers)
```

### Arguments

`se` SummarizedExperiment containing all necessary information of the proteomics data set

`poma_res_outliers` Outliers data.table returned by the detect\_outliers\_POMA function

**Value**

filtered SummarizedExperiment object

**Examples**

```
data(tuberculosis_TMT_se)
poma_res <- detect_outliers_POMA(tuberculosis_TMT_se)
tuberculosis_TMT_se <- remove_POMA_outliers(tuberculosis_TMT_se, poma_res$outliers)
```

---

remove\_reference\_samples

*Remove reference samples of SummarizedExperiment object (reference samples specified during loading)*

---

**Description**

Remove reference samples of SummarizedExperiment object (reference samples specified during loading)

**Usage**

```
remove_reference_samples(se)
```

**Arguments**

|    |                                                                                      |
|----|--------------------------------------------------------------------------------------|
| se | SummarizedExperiment containing all necessary information of the proteomics data set |
|----|--------------------------------------------------------------------------------------|

**Value**

filtered SummarizedExperiment object

**Examples**

```
data(tuberculosis_TMT_se)
tuberculosis_TMT_se <- remove_reference_samples(tuberculosis_TMT_se)
```

---

remove\_samples\_manually

*Remove samples with specific value in column manually*

---

**Description**

Remove samples with specific value in column manually

**Usage**

```
remove_samples_manually(se, column, values)
```

**Arguments**

|        |                                                                                                                                     |
|--------|-------------------------------------------------------------------------------------------------------------------------------------|
| se     | SummarizedExperiment containing all necessary information of the proteomics data set                                                |
| column | String specifying the column of the meta data (samples with the specified value in this column will be removed)                     |
| values | Vector of Strings specifying the value for the removal of samples (samples with this value in the specified column will be removed) |

**Value**

filtered SummarizedExperiment object

**Examples**

```
data(tuberculosis_TMT_se)
tuberculosis_TMT_se <- remove_samples_manually(tuberculosis_TMT_se,
                                              column = "Label", values = c("1.HC.Pool1"))
```

---

r1rMACycNorm

*Cyclic Linear Regression Normalization on MA Transformed Data*


---

**Description**

No reference, but MA transformation and normalization of samples is done pairwise between two samples with A = average of two samples and M = difference. The process is iterated through all samples pairs. Log2 data should be taken as input (on\_raw = FALSE).

**Usage**

```
r1rMACycNorm(
  se,
  ain = "log2",
  aout = "R1rMACyc",
  on_raw = FALSE,
  iterations = 3
)
```

**Arguments**

|            |                                                                                         |
|------------|-----------------------------------------------------------------------------------------|
| se         | SummarizedExperiment containing all necessary information of the proteomic dataset      |
| ain        | String which assay should be used as input                                              |
| aout       | String which assay should be used to save normalized data                               |
| on_raw     | Boolean specifying whether normalization should be performed on raw or log2-scaled data |
| iterations | Number of cyclic iterations to be performed                                             |

## Value

SummarizedExperiment containing the RlrMACyc normalized data as assay (on log2 scale)

## Examples

```
data(tuberculosis_TMT_se)
tuberculosis_TMT_se <- rlrMACycNorm(tuberculosis_TMT_se, ain = "log2",
                                   aout = "RlrMACyc", on_row = FALSE, iterations=3)
```

---

rlrMANorm

*Linear Regression Normalization on MA Transformed Data*

## Description

Similar to Rlr, but data are MA transformed before normalization, (A = median sample, M = difference of that sample to A). Log2 data should be taken as input (on\_raw = FALSE).

## Usage

```
rlrMANorm(se, ain = "log2", aout = "RlrMA", on_raw = FALSE)
```

## Arguments

|        |                                                                                         |
|--------|-----------------------------------------------------------------------------------------|
| se     | SummarizedExperiment containing all necessary information of the proteomic dataset      |
| ain    | String which assay should be used as input                                              |
| aout   | String which assay should be used to save normalized data                               |
| on_raw | Boolean specifying whether normalization should be performed on raw or log2-scaled data |

## Value

SummarizedExperiment containing the RlrMA normalized data as assay (on log2 scale)

## Examples

[illegible]

rlrNorm

*Robust Linear Regression Normalization of NormalyzerDE.***Description**

Uses median values over all samples as reference sample to which all the other samples in the data are normalized to. Log2 data should be taken as input (on\_raw = FALSE).

**Usage**

```
rlrNorm(se, ain = "log2", aout = "Rlr", on_raw = FALSE)
```

**Arguments**

|        |                                                                                         |
|--------|-----------------------------------------------------------------------------------------|
| se     | SummarizedExperiment containing all necessary information of the proteomic dataset      |
| ain    | String which assay should be used as input                                              |
| aout   | String which assay should be used to save normalized data                               |
| on_raw | Boolean specifying whether normalization should be performed on raw or log2-scaled data |

**Value**

SummarizedExperiment containing the rlr normalized data as assay (on log2 scale)

**Examples**

```
data(tuberculosis_TMT_se)
tuberculosis_TMT_se <- rlrNorm(tuberculosis_TMT_se, ain = "log2",
                              aout = "Rlr", on_raw = FALSE)
```

RobNorm

*Original RobNorm Function***Description**

To robustly normalize expression data (Author: Meng Wang, <https://github.com/mwgrassgreen/RobNorm>).

**Usage**

```
RobNorm(X.0, gamma.0 = 0.5, tol = 10^(-4), step = 200)
```

**Arguments**

|         |                                                                               |
|---------|-------------------------------------------------------------------------------|
| X.0     | The expression matrix in log scale.                                           |
| gamma.0 | The density exponent parameter gamma, in practice, taking gamma.0 = 0.5 or 1. |
| tol     | The tolerance for iterations (default: 10 <sup>-4</sup> ).                    |
| step    | The step limit (default: 50).                                                 |

**Value**

Normalized expression data

---

|             |                              |
|-------------|------------------------------|
| robnormNorm | <i>RobNorm Normalization</i> |
|-------------|------------------------------|

---

**Description**

Log2-scaled data should be used as input (on\_raw = FALSE).

**Usage**

```
robnormNorm(se, ain = "log2", aout = "RobNorm", on_raw = FALSE, gamma.0 = 0.1)
```

**Arguments**

|         |                                                                                                                                                                                                                                                                                                                                                     |
|---------|-----------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| se      | SummarizedExperiment containing all necessary information of the proteomic dataset                                                                                                                                                                                                                                                                  |
| ain     | String which assay should be used as input                                                                                                                                                                                                                                                                                                          |
| aout    | String which assay should be used to save normalized data                                                                                                                                                                                                                                                                                           |
| on_raw  | Boolean specifying whether normalization should be performed on raw or log2-scaled data                                                                                                                                                                                                                                                             |
| gamma.0 | Numeric representing the exponent of the weighted density. When the sample size is small, the fitted population of some proteins could be locally trapped such that the variance of those proteins was very small under a large gamma. To avoid this, a small gamma is recommended. When sample size smaller than 40, then set gamma to 0.5 or 0.1. |

**Value**

SummarizedExperiment containing the RobNorm normalized data as assay (on log2 scale)

**Examples**

```
data(tuberculosis_TMT_se)
tuberculosis_TMT_se <- robnormNorm(tuberculosis_TMT_se, ain = "log2",
                                   aout = "RobNorm", on_raw = FALSE, gamma.0 = 0.1)
```

run\_DE

*Run DE analysis of a selection of normalized data sets***Description**

Run DE analysis of a selection of normalized data sets

**Usage**

```
run_DE(
  se,
  comparisons,
  ain = NULL,
  condition = NULL,
  DE_method = "limma",
  covariate = NULL,
  logFC = TRUE,
  logFC_up = 1,
  logFC_down = -1,
  p_adj = TRUE,
  alpha = 0.05,
  B = 100,
  K = 500,
  trend = TRUE,
  robust = TRUE,
  DEqMS_PSMs_column = NULL
)
```

**Arguments**

|             |                                                                                                                                                       |
|-------------|-------------------------------------------------------------------------------------------------------------------------------------------------------|
| se          | SummarizedExperiment containing all necessary information of the proteomics data set                                                                  |
| comparisons | Vector of comparisons that are performed in the DE analysis (from specify_comparisons method)                                                         |
| ain         | Vector of strings which assay should be used as input (default NULL). If NULL then all normalization of the se object are plotted next to each other. |
| condition   | column name of condition (if NULL, condition saved in SummarizedExperiment will be taken)                                                             |
| DE_method   | String specifying which DE method should be applied (limma, ROTS, DEqMS)                                                                              |
| covariate   | String specifying which column to include as covariate into limma                                                                                     |
| logFC       | Boolean specifying whether to apply a logFC threshold (TRUE) or not (FALSE)                                                                           |
| logFC_up    | Upper log2 fold change threshold (dividing into up regulated)                                                                                         |
| logFC_down  | Lower log2 fold change threshold (dividing into down regulated)                                                                                       |
| p_adj       | Boolean specifying whether to apply a threshold on adjusted p-values (TRUE) or on raw p-values (FALSE)                                                |
| alpha       | Threshold for adjusted p-values or p-values                                                                                                           |
| B           | Number of bootstrapping for ROTS                                                                                                                      |

|                   |                                                                                                                                                                                                                                                  |
|-------------------|--------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| K                 | Number of top-ranked features for reproducibility optimization                                                                                                                                                                                   |
| trend             | logical, should an intensity-dependent trend be allowed for the prior variance? If FALSE then the prior variance is constant. Alternatively, trend can be a row-wise numeric vector, which will be used as the covariate for the prior variance. |
| robust            | logical, should the estimation of df.prior and var.prior be robustified against outlier sample variances?                                                                                                                                        |
| DEqMS_PSMs_column | String specifying which column name to use for DEqMS (default NULL). Any column of the rowData(se) is accepted.                                                                                                                                  |

**Value**

Data table of DE results of selected normalized data sets

**Examples**

```
data(tuberculosis_TMT_se)
comparisons <- specify_comparisons(tuberculosis_TMT_se, condition = NULL,
                                   sep = NULL, control = NULL)
de_res <- run_DE(tuberculosis_TMT_se, comparisons,
                 ain = NULL, condition = NULL, DE_method = "limma",
                 logFC = TRUE, logFC_up = 1, logFC_down = -1, p_adj = TRUE,
                 alpha = 0.05, B = 100, K = 500, trend = TRUE, robust = TRUE)
```

---

|               |                                                        |
|---------------|--------------------------------------------------------|
| run_DE_single | <i>Run DE analysis on a single normalized data set</i> |
|---------------|--------------------------------------------------------|

---

**Description**

Run DE analysis on a single normalized data set

**Usage**

```
run_DE_single(
  se,
  method,
  comparisons,
  condition = NULL,
  DE_method = "limma",
  covariate = NULL,
  logFC = TRUE,
  logFC_up = 1,
  logFC_down = -1,
  p_adj = TRUE,
  alpha = 0.05,
  B = 100,
  K = 500,
  trend = TRUE,
  robust = TRUE,
  DEqMS_PSMs_column = NULL
)
```

**Arguments**

|                   |                                                                                                                                                                                                                                                  |
|-------------------|--------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| se                | SummarizedExperiment containing all necessary information of the proteomics data set                                                                                                                                                             |
| method            | String specifying which assay should be used as input                                                                                                                                                                                            |
| comparisons       | Vector of comparisons that are performed in the DE analysis (from specify_comparisons method)                                                                                                                                                    |
| condition         | column name of condition (if NULL, condition saved in SummarizedExperiment will be taken)                                                                                                                                                        |
| DE_method         | String specifying which DE method should be applied (limma, ROTS, DEqMS)                                                                                                                                                                         |
| covariate         | String specifying which column to include as covariate into limma                                                                                                                                                                                |
| logFC             | Boolean specifying whether to apply a logFC threshold (TRUE) or not (FALSE)                                                                                                                                                                      |
| logFC_up          | Upper log2 fold change threshold (dividing into up regulated)                                                                                                                                                                                    |
| logFC_down        | Lower log2 fold change threshold (dividing into down regulated)                                                                                                                                                                                  |
| p_adj             | Boolean specifying whether to apply a threshold on adjusted p-values (TRUE) or on raw p-values (FALSE)                                                                                                                                           |
| alpha             | Threshold for adjusted p-values or p-values                                                                                                                                                                                                      |
| B                 | Number of bootstrapping for ROTS                                                                                                                                                                                                                 |
| K                 | Number of top-ranked features for reproducibility optimization                                                                                                                                                                                   |
| trend             | logical, should an intensity-dependent trend be allowed for the prior variance? If FALSE then the prior variance is constant. Alternatively, trend can be a row-wise numeric vector, which will be used as the covariate for the prior variance. |
| robust            | logical, should the estimation of df.prior and var.prior be robustified against outlier sample variances?                                                                                                                                        |
| DEqMS_PSMs_column | String specifying which column name to use for DEqMS (default NULL). Any column of the rowData(se) is accepted.                                                                                                                                  |

**Value**

Data table of DE results

---

|                     |                                                                                                                        |
|---------------------|------------------------------------------------------------------------------------------------------------------------|
| specify_comparisons | <i>Create vector of comparisons for DE analysis (either by single condition (sep = NULL) or by combined condition)</i> |
|---------------------|------------------------------------------------------------------------------------------------------------------------|

---

**Description**

Create vector of comparisons for DE analysis (either by single condition (sep = NULL) or by combined condition)

**Usage**

```
specify_comparisons(se, condition = NULL, sep = NULL, control = NULL)
```

**Arguments**

|           |                                                                                                                |
|-----------|----------------------------------------------------------------------------------------------------------------|
| se        | SummarizedExperiment containing all necessary information of the proteomics data set                           |
| condition | Column name of condition (if NULL, condition saved in SummarizedExperiment will be taken)                      |
| sep       | Separator that separates both groups in the condition vector (NULL if condition composed only of single group) |
| control   | String of control samples (how the control condition is named) (NULL if no control sample)                     |

**Value**

Vector of comparisons for DE analysis

**Examples**

```
data(tuberculosis_TMT_se)
comparisons <- specify_comparisons(tuberculosis_TMT_se, condition = NULL,
                                   sep = NULL, control = NULL)
```

---

spectraCounteBayes\_DEqMS

*Additional function of the DEqMS package*

---

**Description**

Additional function of the DEqMS package

**Usage**

```
spectraCounteBayes_DEqMS(fit, coef_col)
```

**Arguments**

|          |                                                                                                                                                         |
|----------|---------------------------------------------------------------------------------------------------------------------------------------------------------|
| fit      | linear model from function perform_limma                                                                                                                |
| coef_col | an integer vector indicating the column(s) of fit\$coefficients for which the function is to be performed. if not specified, all coefficients are used. |

**Value**

list object

---

`spike_in_de_res`*Example data.table of DE results of a spike-in proteomics data set*

---

**Description**

A `data.table` containing the DE results of the `spike_in_se` data set (limma,  $\logFC > 1$ ,  $\logFC < -1$ ,  $p_{\text{adj}} < 0.05$ )

**Usage**

```
data(spike_in_de_res)
```

**Format**

An object of class `data.table` (inherits from `data.frame`) with 7500 rows and 10 columns.

**Source**

Jürgen Cox, Marco Y. Hein, Christian A. Luber, Igor Paron, Nagarjuna Nagaraj, and Matthias Mann. Accurate Proteome-wide Label-free Quantification by Delayed Normalization and Maximal Peptide Ratio Extraction, Termed MaxLFQ. *Molecular & Cellular Proteomics* 13.9 (Sept. 2014), pp. 2513–2526. <<https://doi.org/10.1074/mcp.M113.031591>>.

---

`spike_in_se`*Example SummarizedExperiment of a spike-in proteomics data set*

---

**Description**

A `SummarizedExperiment` containing the raw and log2-scaled data of 301 proteins measured in 20 samples. Due to size restriction, we only included the relevant columns of the original protein-Groups.txt of MaxQuant.

**Usage**

```
data(spike_in_se)
```

**Format**

An object of class `SummarizedExperiment` with 1500 rows and 6 columns.

**Source**

Jürgen Cox, Marco Y. Hein, Christian A. Luber, Igor Paron, Nagarjuna Nagaraj, and Matthias Mann. Accurate Proteome-wide Label-free Quantification by Delayed Normalization and Maximal Peptide Ratio Extraction, Termed MaxLFQ. *Molecular & Cellular Proteomics* 13.9 (Sept. 2014), pp. 2513–2526. <<https://doi.org/10.1074/mcp.M113.031591>>.

---

|                   |                                                                   |
|-------------------|-------------------------------------------------------------------|
| subset_SE_by_norm | <i>Subset SummarizedExperiment object by normalization assays</i> |
|-------------------|-------------------------------------------------------------------|

---

**Description**

Subset SummarizedExperiment object by normalization assays

**Usage**

```
subset_SE_by_norm(se, ain)
```

**Arguments**

|     |                                                                                      |
|-----|--------------------------------------------------------------------------------------|
| se  | SummarizedExperiment containing all necessary information of the proteomics data set |
| ain | Character vector of assay names to keep in the SummarizedExperiment object           |

**Value**

SummarizedExperiment object with only the selected normalization assays

**Examples**

```
data(tuberculosis_TMT_se)
tuberculosis_TMT_se <- subset_SE_by_norm(tuberculosis_TMT_se,
  ain = c("raw", "log2", "IRS_on_RobNorm"))
```

---

|           |                                                              |
|-----------|--------------------------------------------------------------|
| tib_to_DF | <i>Helper function to transform a tibble to a data table</i> |
|-----------|--------------------------------------------------------------|

---

**Description**

Helper function to transform a tibble to a data table

**Usage**

```
tib_to_DF(expr_data, column_names, row_names)
```

**Arguments**

|              |                                                  |
|--------------|--------------------------------------------------|
| expr_data    | Tibble data frame containing the expression data |
| column_names | Column names of the expression data              |
| row_names    | Row names of the expression data                 |

**Value**

Data table containing the expression data

---

|         |                                                                                |
|---------|--------------------------------------------------------------------------------|
| tmmNorm | <i>Weighted Trimmed Mean of M Values (TMM) Normalization of edgeR package.</i> |
|---------|--------------------------------------------------------------------------------|

---

### Description

Raw data should be taken as input (on\_raw = TRUE).

### Usage

```
tmmNorm(se, ain = "raw", aout = "TMM", on_raw = TRUE)
```

### Arguments

|        |                                                                                         |
|--------|-----------------------------------------------------------------------------------------|
| se     | SummarizedExperiment containing all necessary information of the proteomic dataset      |
| ain    | String which assay should be used as input                                              |
| aout   | String which assay should be used to save normalized data                               |
| on_raw | Boolean specifying whether normalization should be performed on raw or log2-scaled data |

### Value

SummarizedExperiment containing the TMM normalized data as assay (on log2 scale)

### See Also

[calcNormFactors\(\)](#)

### Examples

```
data(tuberculosis_TMT_se)
tuberculosis_TMT_se <- tmmNorm(tuberculosis_TMT_se, ain = "raw",
                              aout = "TMM", on_raw = TRUE)
```

---

|                         |                                                                             |
|-------------------------|-----------------------------------------------------------------------------|
| tuberculosis_TMT_de_res | <i>Example data.table of DE results of a real-world proteomics data set</i> |
|-------------------------|-----------------------------------------------------------------------------|

---

### Description

A data.table containing the DE results of the tuberculosis\_TMT\_se data set (limma, logFC > 1, logFC < -1, p.adj < 0.05)

### Usage

```
data(tuberculosis_TMT_de_res)
```

**Format**

An object of class `data.table` (inherits from `data.frame`) with 9030 rows and 9 columns.

**Source**

Biadlegne et al. Mycobacterium tuberculosis Affects Protein and Lipid Content of Circulating Exosomes in Infected Patients Depending on Tuberculosis Disease State. *Biomedicines* 10.4 (Mar. 2022), p. 783. doi: 10.3390/biomedicines10040783.

---

|                     |                                                                         |
|---------------------|-------------------------------------------------------------------------|
| tuberculosis_TMT_se | <i>Example SummarizedExperiment of a real-world proteomics data set</i> |
|---------------------|-------------------------------------------------------------------------|

---

**Description**

A `SummarizedExperiment` containing the raw and log2-scaled data of 301 proteins measured in 20 samples

**Usage**

```
data(tuberculosis_TMT_se)
```

**Format**

An object of class `SummarizedExperiment` with 301 rows and 20 columns.

**Source**

Biadlegne et al. Mycobacterium tuberculosis Affects Protein and Lipid Content of Circulating Exosomes in Infected Patients Depending on Tuberculosis Disease State. *Biomedicines* 10.4 (Mar. 2022), p. 783. doi: 10.3390/biomedicines10040783.

---

|         |                                                               |
|---------|---------------------------------------------------------------|
| vsnNorm | <i>Variance Stabilization Normalization of limma package.</i> |
|---------|---------------------------------------------------------------|

---

**Description**

Raw data should be taken as input (`on_raw = TRUE`).

**Usage**

```
vsnNorm(se, ain = "raw", aout = "VSN", on_raw = TRUE, VSN_quantile = 0.9)
```

**Arguments**

|                           |                                                                                                                                                             |
|---------------------------|-------------------------------------------------------------------------------------------------------------------------------------------------------------|
| <code>se</code>           | <code>SummarizedExperiment</code> containing all necessary information of the proteomic dataset                                                             |
| <code>ain</code>          | String which assay should be used as input                                                                                                                  |
| <code>aout</code>         | String which assay should be used to save normalized data                                                                                                   |
| <code>on_raw</code>       | Boolean specifying whether normalization should be performed on raw or log2-scaled data                                                                     |
| <code>VSN_quantile</code> | Numeric of length 1. The quantile that is used for the resistant least trimmed sum of squares regression (see <code>vsn2</code> <code>lts.quantile</code> ) |

**Value**

SummarizedExperiment containing the vsn normalized data as assay (on log2-scale)

**See Also**

[normalizeVSN\(\)](#)

**Examples**

```
data(tuberculosis_TMT_se)
tuberculosis_TMT_se <- vsnNorm(tuberculosis_TMT_se, ain = "raw",
                              aout = "VSN", on_raw = TRUE, VSN_quantile = 0.9)
```

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