

# Package ‘DEP’

April 11, 2018

**Title** Differential Enrichment analysis of Proteomics data

**Version** 1.0.1

**Description** This package provides an integrated analysis workflow for robust and reproducible analysis of mass spectrometry proteomics data for differential protein expression or differential enrichment.

It requires tabular input (e.g. txt files) as generated by quantitative analysis softwares of raw mass spectrometry data, such as MaxQuant or IsobarQuant. Functions are provided for data preparation, filtering, variance normalization and imputation of missing values, as well as statistical testing of differentially enriched / expressed proteins. It also includes tools to check intermediate steps in the workflow, such as normalization and missing values imputation. Finally, visualization tools are provided to explore the results, including heatmap, volcano plot and barplot representations. For scientists with limited experience in R, the package also contains wrapper functions that entail the complete analysis workflow and generate a report. Even easier to use are the interactive Shiny apps that are provided by the package.

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**R topics documented:**

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add_rejections	<i>Mark significant proteins</i>
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## Description

add\_rejections marks significant proteins based on defined cutoffs.

## Usage

```
add_rejections(diff, alpha = 0.05, lfc = 1)
```

## Arguments

diff	SummarizedExperiment, Proteomics dataset on which differential enrichment analysis has been performed (output from <a href="#">test_diff()</a> ).
alpha	Numeric(1), Sets the threshold for the adjusted P value.
lfc	Numeric(1), Sets the threshold for the log2 fold change.

## Value

A SummarizedExperiment object annotated with logical columns indicating significant proteins.

## Examples

```
# Load example
data <- UbiLength
data <- data[data$Reverse != "+" & data$Potential.contaminant != "+",]
data_unique <- make_unique(data, "Gene.names", "Protein.IDs", delim = ";")

# Make SummarizedExperiment
columns <- grep("LFQ.", colnames(data_unique))
exp_design <- UbiLength_ExpDesign
se <- make_se(data_unique, columns, exp_design)

# Filter, normalize and impute missing values
filt <- filter_missval(se, thr = 0)
norm <- normalize_vsn(filt)
imputed <- impute(norm, fun = "MinProb", q = 0.01)

# Test for differentially expressed proteins
diff <- test_diff(imputed, "control", "Ctrl")
dep <- add_rejections(diff, alpha = 0.05, lfc = 1)
```

## analyze\_dep

*Differential expression analysis***Description**

`analyze_dep` tests for differential expression of proteins based on protein-wise linear models and empirical Bayes statistics using [limma](#).

**Usage**

```
analyze_dep(se, type = c("all", "control", "manual"), control = NULL,
            alpha = 0.05, lfc = 1, test = NULL, design_formula = formula(~0 +
              condition))
```

**Arguments**

<code>se</code>	SummarizedExperiment, Proteomics data with unique names and identifiers annotated in 'name' and 'ID' columns. Additionally, the colData should contain sample annotation including 'label', 'condition' and 'replicate' columns. The appropriate columns and objects can be generated using <a href="#">make_se</a> or <a href="#">make_se_parse</a> .
<code>type</code>	"all", "control" or "manual", The type of contrasts that will be tested. This can be all possible pairwise comparisons ("all"), limited to the comparisons versus the control ("control"), or manually defined contrasts ("manual").
<code>control</code>	Character(1), The condition to which contrasts are generated (a control condition would be most appropriate).
<code>alpha</code>	Numeric(1), Sets the threshold for the adjusted P value.
<code>lfc</code>	Numeric(1), Sets the threshold for the log2 fold change.
<code>test</code>	Character, The contrasts that will be tested if type = "manual". These should be formatted as "SampleA_vs_SampleB" or c("SampleA_vs_SampleC", "SampleB_vs_SampleC").
<code>design_formula</code>	Formula, Used to create the design matrix.

**Value**

A SummarizedExperiment object containing FDR estimates of differential expression and logical columns indicating significant proteins.

**Examples**

```
# Load datasets
data <- UbiLength
exp_design <- UbiLength_ExpDesign

# Import and process data
se <- import_MaxQuant(data, exp_design)
processed <- process(se)

# Differential protein expression analysis
dep <- analyze_dep(processed, "control", "Ctrl")
dep <- analyze_dep(processed, "control", "Ctrl",
                   alpha = 0.01, lfc = log2(1.5))
dep <- analyze_dep(processed, "manual", test = c("Ubi6_vs_Ubi4"))
```

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DEP	<i>DEP: A package for Differential Enrichment analysis of Proteomics data.</i>
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## Description

This package provides an integrated analysis workflow for robust and reproducible analysis of mass spectrometry proteomics data for differential protein expression or differential enrichment. It requires tabular input (e.g. txt files) as generated by quantitative analysis softwares of raw mass spectrometry data, such as [MaxQuant](#) or [IsobarQuant](#). Functions are provided for data preparation, filtering, variance normalization and imputation of missing values, as well as statistical testing of differentially enriched / expressed proteins. It also includes tools to check intermediate steps in the workflow, such as normalization and missing values imputation. Finally, visualization tools are provided to explore the results, including heatmap, volcano plot and barplot representations. For scientists with limited experience in R, the package also entails wrapper functions that entail the complete analysis workflow and generate a report. Even easier to use are the interactive Shiny apps that are provided by the package.

## Shiny apps

- [run\\_app](#): Shiny apps for interactive analysis.

## Workflow functions

- [LFQ](#): Label-free quantification (LFQ) workflow wrapper.
- [TMT](#): Tandem-mass-tags (TMT) workflow wrapper.
- [report](#): Create a rmarkdown report wrapper.

## Wrapper functions

- [import\\_MaxQuant](#): Import data from MaxQuant into a SummarizedExperiment object.
- [import\\_IsobarQuant](#): Import data from IsobarQuant into a SummarizedExperiment object.
- [process](#): Perform filtering, normalization and imputation on protein data.
- [analyze\\_dep](#): Differential protein expression analysis.
- [plot\\_all](#): Visualize the results in different types of plots.

## Main functions

- [make\\_unique](#): Generate unique names.
- [make\\_se\\_parse](#): Turn data.frame into SummarizedExperiment by parsing column names.
- [make\\_se](#): Turn data.frame into SummarizedExperiment using an experimental design.
- [filter\\_missval](#): Filter on missing values.
- [normalize\\_vsn](#): Normalize data using vsn.
- [impute](#): Impute missing values.
- [test\\_diff](#): Differential enrichment analysis.
- [add\\_rejections](#): Mark significant proteins.
- [get\\_results](#): Generate a results table.

### Visualization functions

- `plot_single`: Barplot for a protein of interest.
- `plot_volcano`: Volcano plot for a specified contrast.
- `plot_heatmap`: Heatmap of all significant proteins.
- `plot_normalization`: Boxplots to inspect normalization.
- `plot_detect`: Density and CumSum plots of proteins with and without missing values.
- `plot_imputation`: Density plots to inspect imputation.
- `plot_missval`: Heatmap to inspect missing values.
- `plot_numbers`: Barplot of proteins identified.
- `plot_frequency`: Barplot of protein identification overlap between conditions.
- `plot_coverage`: Barplot of the protein coverage in conditions.
- `plot_pca`: PCA plot of top variable proteins.
- `plot_cor`: Correlation matrix.
- `plot_p_hist`: P value histogram.
- `plot_cond_freq`: Barplot of the number of significant conditions per protein.
- `plot_cond_overlap`: Barplot of the number of proteins for overlapping conditions.
- `plot_cond`: Barplot of the frequency of significant conditions per protein and the overlap in proteins between conditions.

### Gene Set Enrichment Analysis functions

- `test_gsea`: Gene Set Enrichment Analysis using enrichR.
- `plot_gsea`: Barplot of enriched gene sets.

### Additional functions

- `get_df_wide`: Generate a wide data.frame from a SummarizedExperiment.
- `get_df_long`: Generate a long data.frame from a SummarizedExperiment.
- `se2msn`: SummarizedExperiment object to MSnSet object conversion.
- `manual_impute`: Imputation by random draws from a manually defined distribution.
- `get_prefix`: Obtain the longest common prefix.

### Example data

- `UbiLength`: Ubiquitin interactors of different linear ubiquitin lengths (UbIA-MS dataset) (Zhang, Smits, van Tilburg et al. Mol. Cell 2017).
- `UbiLength_ExpDesign`: Experimental design of the UbiLength dataset.
- `DiUbi`: Ubiquitin interactors for different diubiquitin-linkages (UbIA-MS dataset) (Zhang, Smits, van Tilburg et al. Mol. Cell 2017).
- `DiUbi_ExpDesign`: Experimental design of the DiUbi dataset.

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DiUbi	<i>DiUbi - Ubiquitin interactors for different diubiquitin-linkages (UbIA-MS dataset)</i>
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## Description

The DiUbi dataset contains label free quantification (LFQ) and intensity-based absolute quantification (iBAQ) data for ubiquitin interactors of different diubiquitin-linkages, generated by Zhang et al 2017. The dataset contains the proteingroups output file from [MaxQuant](#).

## Usage

DiUbi

## Format

A data.frame with 4071 observations and 102 variables:

**Protein.IDs** Uniprot IDs

**Majority.protein.IDs** Uniprot IDs of major protein(s) in the protein group

**Protein.names** Full protein names

**Gene.names** Gene name

**Fasta.headers** Header as present in the Uniprot fasta file

**Peptides** Number of peptides identified for this protein group

**Razor...unique.peptides** Number of peptides used for the quantification of this protein group

**Unique.peptides** Number of peptides identified which are unique for this protein group

**Intensity columns (30)** Raw mass spectrometry intensity, A.U.

**iBAQ columns (30)** iBAQ normalized mass spectrometry intensity, A.U.

**LFQ.intensity columns (30)** LFQ normalized mass spectrometry intensity, A.U.

**Only.identified.by.site** The protein is only identified by a modification site if marked ('+')

**Reverse** The protein is identified in the decoy database if marked ('+')

**Potential.contaminant** The protein is a known contaminant if marked ('+')

**id** The protein group ID

## Value

A data.frame.

## Source

Zhang, Smits, van Tilburg, et al (2017). An interaction landscape of ubiquitin signaling. Molecular Cell 65(5): 941-955. doi: [10.1016/j.molcel.2017.01.004](https://doi.org/10.1016/j.molcel.2017.01.004).

**DiUbi\_ExpDesign**      *Experimental design of the DiUbi dataset*

### Description

The DiUbi\_ExpDesign object annotates 30 different samples of the DiUbi dataset in 10 conditions and 3 replicates.

### Usage

```
DiUbi_ExpDesign
```

### Format

A data.frame with 30 observations and 3 variables:

**label** Label names

**condition** Experimental conditions

**replicate** Replicate number

### Value

A data.frame.

### Source

Zhang, Smits, van Tilburg, et al (2017). An interaction landscape of ubiquitin signaling. Molecular Cell 65(5): 941-955. doi: [10.1016/j.molcel.2017.01.004](https://doi.org/10.1016/j.molcel.2017.01.004).

**filter\_missval**      *Filter on missing values*

### Description

filter\_missval filters a proteomics dataset based on missing values. The dataset is filtered for proteins that have a maximum of 'thr' missing values in at least one condition.

### Usage

```
filter_missval(se, thr = 0)
```

### Arguments

<b>se</b>	SummarizedExperiment, Proteomics data (output from <a href="#">make_se()</a> or <a href="#">make_se_parse()</a> ).
<b>thr</b>	Integer(1), Sets the threshold for the allowed number of missing values in at least one condition.

### Value

A filtered SummarizedExperiment object.

## Examples

```
# Load example
data <- UbiLength
data <- data[data$Reverse != "+" & data$Potential.contaminant != "+",]
data_unique <- make_unique(data, "Gene.names", "Protein.IDs", delim = ";")

# Make SummarizedExperiment
columns <- grep("LFQ.", colnames(data_unique))
exp_design <- UbiLength_ExpDesign
se <- make_se(data_unique, columns, exp_design)

# Filter
stringent_filter <- filter_missval(se, thr = 0)
less_stringent_filter <- filter_missval(se, thr = 1)
```

get\_df\_long

*Generate a long data.frame from a SummarizedExperiment*

## Description

`get_df_long` generate a wide data.frame from a SummarizedExperiment.

## Usage

```
get_df_long(se)
```

## Arguments

se	SummarizedExperiment, Proteomics data (output from <code>make_se()</code> or <code>make_se_parse()</code> ).
----	--

## Value

A data.frame object containing all data in a wide format, where each row represents a single measurement.

## Examples

```
# Load example
data <- UbiLength
data <- data[data$Reverse != "+" & data$Potential.contaminant != "+",]
data_unique <- make_unique(data, "Gene.names", "Protein.IDs", delim = ";")

# Make SummarizedExperiment
columns <- grep("LFQ.", colnames(data_unique))
exp_design <- UbiLength_ExpDesign
se <- make_se(data_unique, columns, exp_design)

# Filter, normalize and impute missing values
filt <- filter_missval(se, thr = 0)
norm <- normalize_vsn(filt)
imputed <- impute(norm, fun = "MinProb", q = 0.01)

# Test for differentially expressed proteins
```

```
diff <- test_diff(imputed, "control", "Ctrl")
dep <- add_rejections(diff, alpha = 0.05, lfc = 1)

# Get a long data.frame
long <- get_df_long(dep)
colnames(long)
```

**get\_df\_wide***Generate a wide data.frame from a SummarizedExperiment***Description**

`get_df_wide` generate a wide data.frame from a `SummarizedExperiment`.

**Usage**

```
get_df_wide(se)
```

**Arguments**

se	SummarizedExperiment, Proteomics data (output from <code>make_se()</code> or <code>make_se_parse()</code> ).
----	--

**Value**

A data.frame object containing all data in a wide format, where each row represents a protein.

**Examples**

```
# Load example
data <- UbiLength
data <- data[data$Reverse != "+" & data$Potential.contaminant != "+",]
data_unique <- make_unique(data, "Gene.names", "Protein.IDs", delim = ";")

# Make SummarizedExperiment
columns <- grep("LFQ.", colnames(data_unique))
exp_design <- UbiLength_ExpDesign
se <- make_se(data_unique, columns, exp_design)

# Filter, normalize and impute missing values
filt <- filter_missval(se, thr = 0)
norm <- normalize_vsn(filt)
imputed <- impute(norm, fun = "MinProb", q = 0.01)

# Test for differentially expressed proteins
diff <- test_diff(imputed, "control", "Ctrl")
dep <- add_rejections(diff, alpha = 0.05, lfc = 1)

# Get a wide data.frame
wide <- get_df_wide(dep)
colnames(wide)
```

---

get_prefix	<i>Obtain the longest common prefix</i>
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---

### Description

`get_prefix` returns the longest common prefix of the supplied words.

### Usage

```
get_prefix(words)
```

### Arguments

`words` Character vector, A list of words.

### Value

A character vector containing the prefix.

### Examples

```
# Load example
data <- UbiLength
columns <- grep("LFQ.", colnames(data))

# Get prefix
names <- colnames(data[, columns])
get_prefix(names)
```

---

get_results	<i>Generate a results table</i>
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---

### Description

`get_results` generates a results table from a proteomics dataset on which differential enrichment analysis was performed.

### Usage

```
get_results(dep)
```

### Arguments

`dep` SummarizedExperiment, Data object for which differentially enriched proteins are annotated (output from `test_diff()` and `add_rejections()`).

### Value

A data.frame object containing all results variables from the performed analysis.

## Examples

```
# Load example
data <- UbiLength
data <- data[data$Reverse != "+" & data$Potential.contaminant != "+",]
data_unique <- make_unique(data, "Gene.names", "Protein.IDs", delim = ";")

# Make SummarizedExperiment
columns <- grep("LFQ.", colnames(data_unique))
exp_design <- UbiLength_ExpDesign
se <- make_se(data_unique, columns, exp_design)

# Filter, normalize and impute missing values
filt <- filter_missval(se, thr = 0)
norm <- normalize_vsn(filt)
imputed <- impute(norm, fun = "MinProb", q = 0.01)

# Test for differentially expressed proteins
diff <- test_diff(imputed, "control", "Ctrl")
dep <- add_rejections(diff, alpha = 0.05, lfc = 1)

# Get results
results <- get_results(dep)
colnames(results)

significant_proteins <- results[results$significant,]
nrow(significant_proteins)
head(significant_proteins)
```

*import\_IsobarQuant      Import from IsobarQuant*

## Description

`import_IsobarQuant` imports a protein table from IsobarQuant and converts it into a SummarizedExperiment object.

## Usage

```
import_IsobarQuant(proteins, expdesign, intensities = "signal_sum",
                   names = "gene_name", ids = "protein_id", delim = "[|]")
```

## Arguments

<code>proteins</code>	Data.frame, Protein table for which unique names will be created.
<code>expdesign</code>	Data.frame, Experimental design with 'label', 'condition' and 'replicate' information. See <a href="#">UbiLength_ExpDesign</a> for an example experimental design.
<code>intensities</code>	Character(1), Prefix of the columns containing sample intensities.
<code>names</code>	Character(1), Name of the column containing feature names.
<code>ids</code>	Character(1), Name of the column containing feature IDs.
<code>delim</code>	Character(1), Sets the delimiter separating the feature names within on protein group.

`import_MaxQuant`

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### Value

A SummarizedExperiment object with log2-transformed values and "name" and "ID" columns containing unique names and identifiers.

### Examples

```
## Not run:  
# Load data  
isobarquant_table <- read.csv("testfile.txt", header = TRUE,  
                               stringsAsFactors = FALSE, sep = "\t")  
exp_design <- read.csv("test_experimental_design.txt", header = TRUE,  
                       stringsAsFactors = FALSE, sep = "\t")  
# Import data  
se <- import_IsobarQuant(isobarquant_table, exp_design)  
  
## End(Not run)
```

---

`import_MaxQuant`      *Import from MaxQuant*

---

### Description

`import_MaxQuant` imports a protein table from MaxQuant and converts it into a SummarizedExperiment object.

### Usage

```
import_MaxQuant(proteins, expdesign, filter = c("Reverse",  
                                              "Potential.contaminant"), intensities = "LFQ", names = "Gene.names",  
                                              ids = "Protein.IDs", delim = ";")
```

### Arguments

<code>proteins</code>	Data.frame, Protein table originating from MaxQuant.
<code>expdesign</code>	Data.frame, Experimental design with 'label', 'condition' and 'replicate' information. See <a href="#">UbiLength_ExpDesign</a> for an example experimental design.
<code>filter</code>	Character, Name of the column(s) containing features to be filtered on.
<code>intensities</code>	Character(1), Prefix of the columns containing sample intensities.
<code>names</code>	Character(1), Name of the column containing feature names.
<code>ids</code>	Character(1), Name of the column containing feature IDs.
<code>delim</code>	Character(1), Sets the delimiter separating the feature names within one protein group.

### Value

A SummarizedExperiment object with log2-transformed values and "name" and "ID" columns containing unique names and identifiers.

## Examples

```
# Load example data and experimental design
data <- UbiLength
exp_design <- UbiLength_ExpDesign

# Import data
se <- import_MaxQuant(data, exp_design)
```

impute	<i>Impute missing values</i>
--------	------------------------------

## Description

impute imputes missing values in a proteomics dataset.

## Usage

```
impute(se, fun = c("bpca", "knn", "QRILC", "MLE", "MinDet", "MinProb", "man",
"min", "zero", "mixed", "nbavg"), ...)
```

## Arguments

se	SummarizedExperiment, Proteomics data (output from <a href="#">make_se()</a> or <a href="#">make_se_parse()</a> ). It is advised to first remove proteins with too many missing values using <a href="#">filter_missval()</a> and normalize the data using <a href="#">normalize_vsn()</a> .
fun	"bpca", "knn", "QRILC", "MLE", "MinDet", "MinProb", "man", "min", "zero", "mixed" or "nbavg", Function used for data imputation based on <a href="#">manual_impute</a> and <a href="#">impute</a> .
...	Additional arguments for imputation functions as depicted in <a href="#">manual_impute</a> and <a href="#">impute</a> .

## Value

An imputed SummarizedExperiment object.

## Examples

```
# Load example
data <- UbiLength
data <- data[data$Reverse != "+" & data$Potential.contaminant != "+",]
data_unique <- make_unique(data, "Gene.names", "Protein.IDs", delim = ";")

# Make SummarizedExperiment
columns <- grep("LFQ.", colnames(data_unique))
exp_design <- UbiLength_ExpDesign
se <- make_se(data_unique, columns, exp_design)

# Filter and normalize
filt <- filter_missval(se, thr = 0)
norm <- normalize_vsn(filt)

# Impute missing values using different functions
```

```
imputed_MinProb <- impute(norm, fun = "MinProb", q = 0.05)
imputed_QRILC <- impute(norm, fun = "QRILC")

imputed_knn <- impute(norm, fun = "knn", k = 10, rowmax = 0.9)
imputed_MLE <- impute(norm, fun = "MLE")

imputed_manual <- impute(norm, fun = "man", shift = 1.8, scale = 0.3)
```

LFQ

*LFQ workflow*

## Description

LFQ is a wrapper function running the entire differential enrichment/expression analysis workflow for label free quantification (LFQ)-based proteomics data. The protein table from **MaxQuant** is used as direct input.

## Usage

```
LFQ(proteins, expdesign, fun = c("man", "bpca", "knn", "QRILC", "MLE",
"MinDet", "MinProb", "min", "zero", "mixed", "nbavg"), type = c("all",
"control", "manual"), control = NULL, test = NULL, filter = c("Reverse",
"Potential.contaminant"), name = "Gene.names", ids = "Protein.IDs",
alpha = 0.05, lfc = 1)
```

## Arguments

<code>proteins</code>	Data.frame, The data object.
<code>expdesign</code>	Data.frame, The experimental design object.
<code>fun</code>	"man", "bpca", "knn", "QRILC", "MLE", "MinDet", "MinProb", "min", "zero", "mixed" or "nbavg", Function used for data imputation based on <code>manual_impute</code> and <code>impute</code> .
<code>type</code>	'all', 'control' or 'manual', The type of contrasts that will be generated.
<code>control</code>	Character(1), The sample name to which the contrasts are generated (the control sample would be most appropriate).
<code>test</code>	Character, The contrasts that will be tested if type = "manual". These should be formatted as "SampleA_vs_SampleB" or c("SampleA_vs_SampleC", "SampleB_vs_SampleC").
<code>filter</code>	Character, Name(s) of the column(s) to be filtered on.
<code>name</code>	Character(1), Name of the column representing gene names.
<code>ids</code>	'Character(1), Name of the column representing protein IDs.
<code>alpha</code>	Numeric(1), sets the false discovery rate threshold.
<code>lfc</code>	Numeric(1), sets the log fold change threshold.

**Value**

A list of 9 objects:

<code>data</code>	data.frame containing the original data
<code>se</code>	SummarizedExperiment object containing the original data
<code>filt</code>	SummarizedExperiment object containing the filtered data
<code>norm</code>	SummarizedExperiment object containing the normalized data
<code>imputed</code>	SummarizedExperiment object containing the imputed data
<code>diff</code>	SummarizedExperiment object containing FDR estimates of differential expression
<code>dep</code>	SummarizedExperiment object annotated with logical columns indicating significant proteins
<code>results</code>	data.frame containing all results variables from the performed analysis
<code>param</code>	data.frame containing the test parameters

**Examples**

```
data <- UbiLength
expdesign <- UbiLength_ExpDesign
results <- LFQ(data, expdesign, 'MinProb', 'control', 'Ctrl')
```

**make\_se**

*Data.frame to SummarizedExperiment object conversion using an experimental design*

**Description**

`make_se` creates a `SummarizedExperiment` object based on two `data.frames`: the protein table and experimental design.

**Usage**

```
make_se(proteins_unique, columns, expdesign)
```

**Arguments**

<code>proteins_unique</code>	Data.frame, Protein table with unique names annotated in the 'name' column (output from <a href="#">make_unique()</a> ).
<code>columns</code>	Integer vector, Column numbers indicating the columns containing the assay data.
<code>expdesign</code>	Data.frame, Experimental design with 'label', 'condition' and 'replicate' information. See <a href="#">UbiLength_ExpDesign</a> for an example experimental design.

**Value**

A `SummarizedExperiment` object with log2-transformed values.

## Examples

```
# Load example
data <- UbiLength
data <- data[data$Reverse != "+" & data$Potential.contaminant != "+",]
data_unique <- make_unique(data, "Gene.names", "Protein.IDs", delim = ";")

# Make SummarizedExperiment
columns <- grep("LFQ.", colnames(data_unique))
exp_design <- UbiLength_ExpDesign
se <- make_se(data_unique, columns, exp_design)
```

make\_se\_parse

*Data.frame to SummarizedExperiment object conversion using parsing from column names*

## Description

make\_se\_parse creates a SummarizedExperiment object based on a single data.frame.

## Usage

```
make_se_parse(proteins_unique, columns, mode = c("char", "delim"),
             chars = 1, sep = "_")
```

## Arguments

proteins_unique	Data.frame, Protein table with unique names annotated in the 'name' column (output from <a href="#">make_unique()</a> ).
columns	Integer vector, Column numbers indicating the columns containing the assay data.
mode	"char" or "delim", The mode of parsing the column headers. "char" will parse the last number of characters as replicate number and requires the 'chars' parameter. "delim" will parse on the separator and requires the 'sep' parameter.
chars	Numeric(1), The number of characters to take at the end of the column headers as replicate number (only for mode == "char").
sep	Character(1), The separator used to parse the column header (only for mode == "delim").

## Value

A SummarizedExperiment object with log2-transformed values.

## Examples

```
# Load example
data <- UbiLength
data <- data[data$Reverse != "+" & data$Potential.contaminant != "+",]
data_unique <- make_unique(data, "Gene.names", "Protein.IDs", delim = ";")

# Make SummarizedExperiment
```

---

```
columns <- grep("LFQ.", colnames(data_unique))
se <- make_se_parse(data_unique, columns, mode = "char", chars = 1)
se <- make_se_parse(data_unique, columns, mode = "delim", sep = "_")
```

---

**make\_unique** *Make unique names*

---

### Description

`make_unique` generates unique identifiers for a proteomics dataset based on "name" and "id" columns.

### Usage

```
make_unique(proteins, names, ids, delim = ";")
```

### Arguments

<code>proteins</code>	Data.frame, Protein table for which unique names will be created.
<code>names</code>	Character(1), Name of the column containing feature names.
<code>ids</code>	Character(1), Name of the column containing feature IDs.
<code>delim</code>	Character(1), Sets the delimiter separating the feature names within one protein group.

### Value

A data.frame with the additional variables "name" and "ID" containing unique names and identifiers, respectively.

### Examples

```
# Load example
data <- UbiLength

# Check colnames and pick the appropriate columns
colnames(data)
data_unique <- make_unique(data, "Gene.names", "Protein.IDs", delim = ";")
```

---

**manual\_impute** *Imputation by random draws from a manually defined distribution*

---

### Description

`manual_impute` imputes missing values in a proteomics dataset by random draws from a manually defined distribution.

### Usage

```
manual_impute(se, scale = 0.3, shift = 1.8)
```

**Arguments**

- se** SummarizedExperiment, Proteomics data (output from [make\\_se\(\)](#) or [make\\_se\\_parse\(\)](#)). It is advised to first remove proteins with too many missing values using [filter\\_missval\(\)](#) and normalize the data using [normalize\\_vsn\(\)](#).
- scale** Numeric(1), Sets the width of the distribution relative to the standard deviation of the original distribution.
- shift** Numeric(1), Sets the left-shift of the distribution (in standard deviations) from the median of the original distribution.

**Value**

An imputed SummarizedExperiment object.

**Examples**

```
# Load example
data <- UbiLength
data <- data[data$Reverse != "+" & data$Potential.contaminant != "+",]
data_unique <- make_unique(data, "Gene.names", "Protein.IDs", delim = ";")

# Make SummarizedExperiment
columns <- grep("LFQ.", colnames(data_unique))
exp_design <- UbiLength_ExpDesign
se <- make_se(data_unique, columns, exp_design)

# Filter and normalize
filt <- filter_missval(se, thr = 0)
norm <- normalize_vsn(filt)

# Impute missing values manually
imputed_manual <- impute(norm, fun = "man", shift = 1.8, scale = 0.3)
```

meanSdPlot

*Plot row standard deviations versus row means***Description**

meanSdPlot generates a hexagonal heatmap of the row standard deviations versus row means from SummarizedExperiment objects. See [meanSdPlot](#).

**Usage**

```
meanSdPlot(x, ranks = TRUE, xlab = ifelse(ranks, "rank(mean)", "mean"),
           ylab = "sd", pch = TRUE, bins = 50, ...)
```

**Arguments**

- x** SummarizedExperiment, Data object.
- ranks** Logical, Whether or not to plot the row means on the rank scale.
- xlab** Character, x-axis label.
- ylab** Character, y-axis label.

pch	Ignored - exists for backward compatibility.
plot	Logical, Whether or not to produce the plot.
bins	Numeric vector, Data object before normalization.
...	Other arguments, Passed to <a href="#">stat_binhex</a> .

**Value**

A scatter plot of row standard deviations versus row means(generated by [stat\\_binhex](#))

**Examples**

```
# Load example
data <- UbiLength
data <- data[data$Reverse != "+" & data$Potential.contaminant != "+",]
data_unique <- make_unique(data, "Gene.names", "Protein.IDs", delim = ";")

# Make SummarizedExperiment
columns <- grep("LFQ.", colnames(data_unique))
exp_design <- UbiLength_ExpDesign
se <- make_se(data_unique, columns, exp_design)

# Filter and normalize
filt <- filter_missval(se, thr = 0)
norm <- normalize_vsn(filt)

# Plot meanSdPlot
meanSdPlot(norm)
```

**normalize\_vsn**      *Normalization using vsn*

**Description**

`normalize_vsn` performs variance stabilizing transformation using the [vsn-package](#).

**Usage**

```
normalize_vsn(se)
```

**Arguments**

se	SummarizedExperiment, Proteomics data (output from <a href="#">make_se()</a> or <a href="#">make_se_parse()</a> ). It is advised to first remove proteins with too many missing values using <a href="#">filter_missval()</a> .
----	---

**Value**

A normalized SummarizedExperiment object.

## Examples

```
# Load example
data <- UbiLength
data <- data[data$Reverse != "+" & data$Potential.contaminant != "+",]
data_unique <- make_unique(data, "Gene.names", "Protein.IDs", delim = ";")

# Make SummarizedExperiment
columns <- grep("LFQ.", colnames(data_unique))
exp_design <- UbiLength_ExpDesign
se <- make_se(data_unique, columns, exp_design)

# Filter and normalize
filt <- filter_missval(se, thr = 0)
norm <- normalize_vsn(filt)
```

### plot\_all

*Visualize the results in different types of plots*

## Description

`plot_all` visualizes the results of the differential protein expression analysis in different types of plots. These are (1) volcano plots, (2) heatmaps, (3) single protein plots, (4) frequency plots and/or (5) comparison plots.

## Usage

```
plot_all(dep, plots = c("volcano", "heatmap", "single", "freq", "comparison"))
```

## Arguments

dep	SummarizedExperiment, Data object which has been generated by <a href="#">analyze_dep</a> or the combination of <a href="#">test_diff</a> and <a href="#">add_rejections</a> .
plots	"volcano", "heatmap", "single", "freq" and/or "comparison",

## Value

Pdfs containg the desired plots.

## Examples

```
# Load datasets
data <- UbiLength
exp_design <- UbiLength_ExpDesign

# Import and process data
se <- import_MaxQuant(data, exp_design)
processed <- process(se)

# Differential protein expression analysis
dep <- analyze_dep(processed, "control", "Ctrl")

## Not run:
# Plot all plots
```

```
plot_all(dep)
## End(Not run)
```

**plot\_cond**

*Plot frequency of significant conditions per protein and the overlap in proteins between conditions*

## Description

`plot_cond` generates a histogram of the number of proteins per condition and stacks for overlapping conditions.

## Usage

```
plot_cond(dep)
```

## Arguments

<code>dep</code>	SummarizedExperiment, Data object for which differentially enriched proteins are annotated (output from <code>test_diff()</code> and <code>add_rejections()</code> ).
------------------	---

## Value

A histogram (generated by `ggplot`)

## Examples

```
# Load example
data <- UbiLength
data <- data[data$Reverse != "+" & data$Potential.contaminant != "+",]
data_unique <- make_unique(data, "Gene.names", "Protein.IDs", delim = ";")

# Make SummarizedExperiment
columns <- grep("LFQ.", colnames(data_unique))
exp_design <- UbiLength_ExpDesign
se <- make_se(data_unique, columns, exp_design)

# Filter, normalize and impute missing values
filt <- filter_missval(se, thr = 0)
norm <- normalize_vsn(filt)
imputed <- impute(norm, fun = "MinProb", q = 0.01)

# Test for differentially expressed proteins
diff <- test_diff(imputed, "control", "Ctrl")
dep <- add_rejections(diff, alpha = 0.05, lfc = 1)

# Plot histogram with overlaps
plot_cond(dep)
```

---

<code>plot_cond_freq</code>	<i>Plot frequency of significant conditions per protein</i>
-----------------------------	---

---

## Description

`plot_cond_freq` generates a histogram of the number of significant conditions per protein.

## Usage

```
plot_cond_freq(dep)
```

## Arguments

<code>dep</code>	SummarizedExperiment, Data object for which differentially enriched proteins are annotated (output from <code>test_diff()</code> and <code>add_rejections()</code> ).
------------------	---

## Value

A histogram (generated by `ggplot`)

## Examples

```
# Load example
data <- UbiLength
data <- data[data$Reverse != "+" & data$Potential.contaminant != "+",]
data_unique <- make_unique(data, "Gene.names", "Protein.IDs", delim = ",")

# Make SummarizedExperiment
columns <- grep("LFQ.", colnames(data_unique))
exp_design <- UbiLength_ExpDesign
se <- make_se(data_unique, columns, exp_design)

# Filter, normalize and impute missing values
filt <- filter_missval(se, thr = 0)
norm <- normalize_vsn(filt)
imputed <- impute(norm, fun = "MinProb", q = 0.01)

# Test for differentially expressed proteins
diff <- test_diff(imputed, "control", "Ctrl")
dep <- add_rejections(diff, alpha = 0.05, lfc = 1)

# Plot frequency of significant conditions
plot_cond_freq(dep)
```

---

<code>plot_cond_overlap</code>	<i>Plot conditions overlap</i>
--------------------------------	--------------------------------

---

## Description

`plot_cond_overlap` generates a histogram of the number of proteins per condition or overlapping conditions.

**Usage**

```
plot_cond_overlap(dep)
```

**Arguments**

**dep** SummarizedExperiment, Data object for which differentially enriched proteins are annotated (output from `test_diff()` and `add_rejections()`).

**Value**

A histogram (generated by `ggplot`)

**Examples**

```
# Load example
data <- UbiLength
data <- data[data$Reverse != "+" & data$Potential.contaminant != "+",]
data_unique <- make_unique(data, "Gene.names", "Protein.IDs", delim = ";")

# Make SummarizedExperiment
columns <- grep("LFQ.", colnames(data_unique))
exp_design <- UbiLength_ExpDesign
se <- make_se(data_unique, columns, exp_design)

# Filter, normalize and impute missing values
filt <- filter_missval(se, thr = 0)
norm <- normalize_vsn(filt)
imputed <- impute(norm, fun = "MinProb", q = 0.01)

# Test for differentially expressed proteins
diff <- test_diff(imputed, "control", "Ctrl")
dep <- add_rejections(diff, alpha = 0.05, lfc = 1)

# Plot condition overlap
plot_cond_overlap(dep)
```

**plot\_cor**

*Plot correlation matrix*

**Description**

`plot_cor` generates a Pearson correlation matrix.

**Usage**

```
plot_cor(dep, significant = TRUE, lower = -1, upper = 1, pal = "PRGn",
         pal_rev = FALSE, indicate = NULL, font_size = 12, ...)
```

### Arguments

dep	SummarizedExperiment, Data object for which differentially enriched proteins are annotated (output from <a href="#">test_diff()</a> and <a href="#">add_rejections()</a> ).
significant	Logical(1), Whether or not to filter for significant proteins.
lower	Integer(1), Sets the lower limit of the color scale.
upper	Integer(1), Sets the upper limit of the color scale.
pal	Character(1), Sets the color panel (from <a href="#">brewer.pal</a> ).
pal_rev	Logical(1), Whether or not to invert the color palette.
indicate	Character, Sets additional annotation on the top of the heatmap based on columns from the experimental design (colData).
font_size	Integer(1), Sets the size of the labels.
...	Additional arguments for Heatmap function as depicted in <a href="#">Heatmap</a>

### Value

A heatmap plot (generated by [Heatmap](#))

### Examples

```
# Load example
data <- UbiLength
data <- data[data$Reverse != "+" & data$Potential.contaminant != "+",]
data_unique <- make_unique(data, "Gene.names", "Protein.IDs", delim = ";")

# Make SummarizedExperiment
columns <- grep("LFQ.", colnames(data_unique))
exp_design <- UbiLength_ExpDesign
se <- make_se(data_unique, columns, exp_design)

# Filter, normalize and impute missing values
filt <- filter_missval(se, thr = 0)
norm <- normalize_vsn(filt)
imputed <- impute(norm, fun = "MinProb", q = 0.01)

# Test for differentially expressed proteins
diff <- test_diff(imputed, "control", "Ctrl")
dep <- add_rejections(diff, alpha = 0.05, lfc = 1)

# Plot correlation matrix
plot_cor(dep)
```

plot\_coverage

*Plot protein coverage*

### Description

plot\_coverage generates a barplot of the protein coverage in all samples.

### Usage

`plot_coverage(se)`

**Arguments**

**se** SummarizedExperiment, Data object for which to plot observation frequency.

**Value**

Barplot of protein coverage in samples (generated by [ggplot](#))

**Examples**

```
# Load example
data <- UbiLength
data <- data[data$Reverse != "+" & data$Potential.contaminant != "+",]
data_unique <- make_unique(data, "Gene.names", "Protein.IDs", delim = ";")

# Make SummarizedExperiment
columns <- grep("LFQ.", colnames(data_unique))
exp_design <- UbiLength_ExpDesign
se <- make_se(data_unique, columns, exp_design)

# Filter and plot coverage
filt <- filter_missval(se, thr = 0)
plot_coverage(filt)
```

**plot\_detect**

*Visualize intensities of proteins with missing values*

**Description**

`plot_detect` generates density and CumSum plots of protein intensities with and without missing values

**Usage**

```
plot_detect(se)
```

**Arguments**

**se** SummarizedExperiment, Data object with missing values.

**Value**

Density and CumSum plots of intensities of proteins with and without missing values (generated by [ggplot](#)).

**Examples**

```
# Load example
data <- UbiLength
data <- data[data$Reverse != "+" & data$Potential.contaminant != "+",]
data_unique <- make_unique(data, "Gene.names", "Protein.IDs", delim = ";")

# Make SummarizedExperiment
columns <- grep("LFQ.", colnames(data_unique))
```

```
exp_design <- UbiLength_ExpDesign
se <- make_se(data_unique, columns, exp_design)

# Filter
filt <- filter_missval(se, thr = 0)

# Plot intensities of proteins with missing values
plot_detect(filt)
```

---

**plot\_frequency***Plot protein overlap between samples*

---

**Description**

plot\_frequency generates a barplot of the protein overlap between samples

**Usage**

```
plot_frequency(se)
```

**Arguments**

se SummarizedExperiment, Data object for which to plot observation frequency.

**Value**

Barplot of overlap of protein identifications between samples (generated by [ggplot](#))

**Examples**

```
# Load example
data <- UbiLength
data <- data[data$Reverse != "+" & data$Potential.contaminant != "+",]
data_unique <- make_unique(data, "Gene.names", "Protein.IDs", delim = ";")

# Make SummarizedExperiment
columns <- grep("LFQ.", colnames(data_unique))
exp_design <- UbiLength_ExpDesign
se <- make_se(data_unique, columns, exp_design)

# Filter and plot frequency
filt <- filter_missval(se, thr = 0)
plot_frequency(filt)
```

**plot\_gsea***Plot enriched Gene Sets***Description**

`plot_gsea` plots enriched gene sets from Gene Set Enrichment Analysis.

**Usage**

```
plot_gsea(gsea_results, number = 10, alpha = 0.05, contrasts = NULL,
          databases = NULL, nrow = 1, term_size = 8)
```

**Arguments**

<code>gsea_results</code>	Data.frame, Gene Set Enrichment Analysis results object. (output from <a href="#">test_gsea()</a> ).
<code>number</code>	Numeric(1), Sets the number of enriched terms per contrast to be plotted.
<code>alpha</code>	Numeric(1), Sets the threshold for the adjusted P value.
<code>contrasts</code>	Character, Specifies the contrast(s) to plot. If 'NULL' all contrasts will be plotted.
<code>databases</code>	Character, Specifies the database(s) to plot. If 'NULL' all databases will be plotted.
<code>nrow</code>	Numeric(1), Sets the number of rows for the plot.
<code>term_size</code>	Numeric(1), Sets the text size of the terms.

**Value**

A barplot of the enriched terms (generated by [ggplot](#)).

**Examples**

```
# Load example
data <- UbiLength
data <- data[data$Reverse != "+" & data$Potential.contaminant != "+",]
data_unique <- make_unique(data, "Gene.names", "Protein.IDs", delim = ";")

# Make SummarizedExperiment
columns <- grep("LFQ.", colnames(data_unique))
exp_design <- UbiLength_ExpDesign
se <- make_se(data_unique, columns, exp_design)

# Filter, normalize and impute missing values
filt <- filter_missval(se, thr = 0)
norm <- normalize_vsn(filt)
imputed <- impute(norm, fun = "MinProb", q = 0.01)

# Test for differentially expressed proteins
diff <- diff <- test_diff(imputed, "control", "Ctrl")
dep <- add_rejections(diff, alpha = 0.05, lfc = 1)

## Not run:
```

```
# Test enrichments
gsea_results <- test_gsea(dep)
plot_gsea(gsea_results)

## End(Not run)
```

**plot\_heatmap***Plot a heatmap***Description**

`plot_heatmap` generates a heatmap of all significant proteins.

**Usage**

```
plot_heatmap(dep, type = c("contrast", "centered"), kmeans = FALSE, k = 6,
             col_limit = 6, indicate = NULL, row_font_size = 6, col_font_size = 10,
             ...)
```

**Arguments**

dep	SummarizedExperiment, Data object for which differentially enriched proteins are annotated (output from <a href="#">test_diff()</a> and <a href="#">add_rejections()</a> ).
type	'contrast' or 'centered', The type of data scaling used for plotting. Either the fold change ('contrast') or the centered log2-intensity ('centered').
kmeans	Logical(1), Whether or not to perform k-means clustering.
k	Integer(1), Sets the number of k-means clusters.
col_limit	Integer(1), Sets the outer limits of the color scale.
indicate	Character, Sets additional annotation on the top of the heatmap based on columns from the experimental design (colData). Only applicable to type = 'centered'.
row_font_size	Integer(1), Sets the size of row labels.
col_font_size	Integer(1), Sets the size of column labels.
...	Additional arguments for Heatmap function as depicted in <a href="#">Heatmap</a>

**Value**

A heatmap (generated by [Heatmap](#))

**Examples**

```
# Load example
data <- UbiLength
data <- data[data$Reverse != "+" & data$Potential.contaminant != "+",]
data_unique <- make_unique(data, "Gene.names", "Protein.IDs", delim = ",")

# Make SummarizedExperiment
columns <- grep("LFQ.", colnames(data_unique))
exp_design <- UbiLength_ExpDesign
se <- make_se(data_unique, columns, exp_design)
```

```

# Filter, normalize and impute missing values
filt <- filter_missval(se, thr = 0)
norm <- normalize_vsn(filt)
imputed <- impute(norm, fun = "MinProb", q = 0.01)

# Test for differentially expressed proteins
diff <- test_diff(imputed, "control", "Ctrl")
dep <- add_rejections(diff, alpha = 0.05, lfc = 1)

# Plot heatmap
plot_heatmap(dep)
plot_heatmap(dep, 'centered', kmeans = TRUE, k = 6, row_font_size = 3)
plot_heatmap(dep, 'contrast', col_limit = 10, row_font_size = 3)

```

---

**plot\_imputation**      *Visualize imputation*

---

## Description

`plot_imputation` generates density plots for all conditions before and after imputation

## Usage

```
plot_imputation(raw, imp)
```

## Arguments

<code>raw</code>	SummarizedExperiment, Data object before imputation (output from <a href="#">normalize_vsn()</a> ).
<code>imp</code>	SummarizedExperiment, Data object after imputation (output from <a href="#">impute()</a> ).

## Value

Density plots for all conditions before and after imputation (generated by [ggplot](#)).

## Examples

```

# Load example
data <- UbiLength
data <- data[data$Reverse != "+" & data$Potential.contaminant != "+",]
data_unique <- make_unique(data, "Gene.names", "Protein.IDs", delim = ";")

# Make SummarizedExperiment
columns <- grep("LFQ.", colnames(data_unique))
exp_design <- UbiLength_ExpDesign
se <- make_se(data_unique, columns, exp_design)

# Filter, normalize and impute missing values
filt <- filter_missval(se, thr = 0)
norm <- normalize_vsn(filt)
imputed <- impute(norm, fun = "MinProb", q = 0.01)

# Plot imputation
plot_imputation(norm, imputed)

```

---

plot_missval	<i>Plot a heatmap of proteins with missing values</i>
--------------	---

---

## Description

plot\_missval generates a heatmap of proteins with missing values to discover whether values are missing by random or not.

## Usage

```
plot_missval(se)
```

## Arguments

se                   SummarizedExperiment, Data object with missing values.

## Value

A heatmap indicating whether values are missing (0) or not (1) (generated by [Heatmap](#)).

## Examples

```
# Load example
data <- UbiLength
data <- data[data$Reverse != "+" & data$Potential.contaminant != "+",]
data_unique <- make_unique(data, "Gene.names", "Protein.IDs", delim = ";")

# Make SummarizedExperiment
columns <- grep("LFQ.", colnames(data_unique))
exp_design <- UbiLength_ExpDesign
se <- make_se(data_unique, columns, exp_design)

# Filter, normalize and impute missing values
filt <- filter_missval(se, thr = 0)

# Plot missing values heatmap
plot_missval(filt)
```

---

plot_normalization	<i>Visualize normalization</i>
--------------------	--------------------------------

---

## Description

plot\_normalization generates boxplots for all conditions before and after normalization.

## Usage

```
plot_normalization(raw, norm)
```

**Arguments**

- `raw` SummarizedExperiment, Data object before normalization (output from [make\\_se\(\)](#) or [make\\_se\\_parse\(\)](#)).  
`norm` SummarizedExperiment, Data object after normalization (output from [normalize\\_vsn](#)).

**Value**

Boxplots for all conditions before and after normalization (generated by [ggplot](#)). Adding components and other plot adjustments can be easily done using the [ggplot2](#) syntax (i.e. using '+')

**Examples**

```
# Load example
data <- UbiLength
data <- data[data$Reverse != "+" & data$Potential.contaminant != "+",]
data_unique <- make_unique(data, "Gene.names", "Protein.IDs", delim = ";")

# Make SummarizedExperiment
columns <- grep("LFQ.", colnames(data_unique))
exp_design <- UbiLength_ExpDesign
se <- make_se(data_unique, columns, exp_design)

# Filter and normalize
filt <- filter_missval(se, thr = 0)
norm <- normalize_vsn(filt)

# Plot normalization
plot_normalization(filt, norm)
```

**plot\_numbers** *Plot protein numbers*

**Description**

`plot_numbers` generates a barplot of the number of identified proteins per sample.

**Usage**

```
plot_numbers(se)
```

**Arguments**

- `se` SummarizedExperiment, Data object for which to plot protein numbers (output from [make\\_se\(\)](#) or [make\\_se\\_parse\(\)](#)).

**Value**

Barplot of the number of identified proteins per sample (generated by [ggplot](#))

## Examples

```
# Load example
data <- UbiLength
data <- data[data$Reverse != "+" & data$Potential.contaminant != "+",]
data_unique <- make_unique(data, "Gene.names", "Protein.IDs", delim = ";")

# Make SummarizedExperiment
columns <- grep("LFQ.", colnames(data_unique))
exp_design <- UbiLength_ExpDesign
se <- make_se(data_unique, columns, exp_design)

# Filter and plot numbers
filt <- filter_missval(se, thr = 0)
plot_numbers(filt)
```

plot\_pca

*Plot PCA*

## Description

`plot_pca` generates a PCA plot using the top variable proteins.

## Usage

```
plot_pca(dep, x = 1, y = 2, indicate = c("condition", "replicate"),
          label = FALSE, n = 500, point_size = 6, label_size = 3)
```

## Arguments

<code>dep</code>	SummarizedExperiment, Data object for which differentially enriched proteins are annotated (output from <a href="#">test_diff()</a> and <a href="#">add_rejections()</a> ).
<code>x</code>	Integer(1), Sets the principle component to plot on the x-axis.
<code>y</code>	Integer(1), Sets the principle component to plot on the y-axis.
<code>indicate</code>	Character, Sets the color, shape and facet_wrap of the plot based on columns from the experimental design (colData).
<code>label</code>	Logical, Whether or not to add sample labels.
<code>n</code>	Integer(1), Sets the number of top variable proteins to consider.
<code>point_size</code>	Integer(1), Sets the size of the points.
<code>label_size</code>	Integer(1), Sets the size of the labels

## Value

A scatter plot (generated by [ggplot](#)).

## Examples

```
# Load example
data <- UbiLength
data <- data[data$Reverse != "+" & data$Potential.contaminant != "+",]
data_unique <- make_unique(data, "Gene.names", "Protein.IDs", delim = ";")

# Make SummarizedExperiment
columns <- grep("LFQ.", colnames(data_unique))
exp_design <- UbiLength_ExpDesign
se <- make_se(data_unique, columns, exp_design)

# Filter, normalize and impute missing values
filt <- filter_missval(se, thr = 0)
norm <- normalize_vsn(filt)
imputed <- impute(norm, fun = "MinProb", q = 0.01)

# Test for differentially expressed proteins
diff <- test_diff(imputed, "control", "Ctrl")
dep <- add_rejections(diff, alpha = 0.05, lfc = 1)

# Plot PCA
plot_pca(dep)
plot_pca(dep, indicate = "condition")
```

**plot\_p\_hist**

*Plot a P value histogram*

## Description

`plot_p_hist` generates a p value histogram.

## Usage

```
plot_p_hist(dep, adjusted = FALSE, wrap = FALSE)
```

## Arguments

<code>dep</code>	SummarizedExperiment, Data object for which differentially enriched proteins are annotated (output from <code>test_diff()</code> and <code>add_rejections()</code> ).
<code>adjusted</code>	Logical(1), Whether or not to use adjusted p values.
<code>wrap</code>	Logical(1), Whether or not to display different histograms for the different contrasts.

## Value

A histogram (generated by `ggplot`).

## Examples

```
# Load example
data <- UbiLength
data <- data[data$Reverse != "+" & data$Potential.contaminant != "+",]
data_unique <- make_unique(data, "Gene.names", "Protein.IDs", delim = ";")

# Make SummarizedExperiment
columns <- grep("LFQ.", colnames(data_unique))
exp_design <- UbiLength_ExpDesign
se <- make_se(data_unique, columns, exp_design)

# Filter, normalize and impute missing values
filt <- filter_missval(se, thr = 0)
norm <- normalize_vsn(filt)
imputed <- impute(norm, fun = "MinProb", q = 0.01)

# Test for differentially expressed proteins
diff <- test_diff(imputed, "control", "Ctrl")
dep <- add_rejections(diff, alpha = 0.05, lfc = 1)

# Plot p value histogram
plot_p_hist(dep)
plot_p_hist(dep, wrap = TRUE)
```

**plot\_single**

*Plot values for a protein of interest*

## Description

`plot_single` generates a barplot of a protein of interest.

## Usage

```
plot_single(dep, proteins, type = c("contrast", "centered"))
```

## Arguments

<code>dep</code>	SummarizedExperiment, Data object for which differentially enriched proteins are annotated (output from <code>test_diff()</code> and <code>add_rejections()</code> ).
<code>proteins</code>	Character, The name of the proteins to plot.
<code>type</code>	'contrast' or 'centered', The type of data scaling used for plotting. Either the fold change ('contrast') or the centered log2-intensity ('centered').

## Value

A barplot (generated by `ggplot`).

## Examples

```
# Load example
data <- UbiLength
data <- data[data$Reverse != "+" & data$Potential.contaminant != "+",]
data_unique <- make_unique(data, "Gene.names", "Protein.IDs", delim = ";")

# Make SummarizedExperiment
columns <- grep("LFQ.", colnames(data_unique))
exp_design <- UbiLength_ExpDesign
se <- make_se(data_unique, columns, exp_design)

# Filter, normalize and impute missing values
filt <- filter_missval(se, thr = 0)
norm <- normalize_vsn(filt)
imputed <- impute(norm, fun = "MinProb", q = 0.01)

# Test for differentially expressed proteins
diff <- test_diff(imputed, "control", "Ctrl")
dep <- add_rejections(diff, alpha = 0.05, lfc = 1)

# Plot single proteins
plot_single(dep, 'USP15')
plot_single(dep, 'USP15', 'centered')
plot_single(dep, c('USP15', 'CUL1'))
```

**plot\_volcano**

*Volcano plot*

## Description

`plot_volcano` generates a volcano plot for a specified contrast.

## Usage

```
plot_volcano(dep, contrast, label_size = 3, add_names = TRUE,
adjusted = FALSE)
```

## Arguments

<code>dep</code>	SummarizedExperiment, Data object for which differentially enriched proteins are annotated (output from <code>test_diff()</code> and <code>add_rejections()</code> ).
<code>contrast</code>	Character(1), Specifies the contrast to plot.
<code>label_size</code>	Integer(1), Sets the size of name labels.
<code>add_names</code>	Logical(1), Whether or not to plot names.
<code>adjusted</code>	Logical(1), Whether or not to use adjusted p values.

## Value

A volcano plot (generated by `ggplot`)

## Examples

```
# Load example
data <- UbiLength
data <- data[data$Reverse != "+" & data$Potential.contaminant != "+",]
data_unique <- make_unique(data, "Gene.names", "Protein.IDs", delim = ";")

# Make SummarizedExperiment
columns <- grep("LFQ.", colnames(data_unique))
exp_design <- UbiLength_ExpDesign
se <- make_se(data_unique, columns, exp_design)

# Filter, normalize and impute missing values
filt <- filter_missval(se, thr = 0)
norm <- normalize_vsn(filt)
imputed <- impute(norm, fun = "MinProb", q = 0.01)

# Test for differentially expressed proteins
diff <- test_diff(imputed, "control", "Ctrl")
dep <- add_rejections(diff, alpha = 0.05, lfc = 1)

# Plot volcano
plot_volcano(dep, 'Ubi6_vs_Ctrl', label_size = 5, add_names = TRUE)
plot_volcano(dep, 'Ubi6_vs_Ctrl', label_size = 5,
             add_names = TRUE, adjusted = TRUE)
plot_volcano(dep, 'Ubi6_vs_Ctrl', add_names = FALSE)
plot_volcano(dep, 'Ubi4_vs_Ctrl', label_size = 5, add_names = TRUE)
```

## Description

process performs data processing on a `SummarizedExperiment` object. It (1) filters a proteomics dataset based on missing values, (2) applies variance stabilizing normalization and (3) imputes eventual remaining missing values.

## Usage

```
process(se, thr = 0, fun = c("man", "bpca", "knn", "QRILC", "MLE", "MinDet",
                            "MinProb", "min", "zero", "mixed", "nbavg"), ...)
```

## Arguments

<code>se</code>	SummarizedExperiment, Proteomics data with unique names and identifiers annotated in 'name' and 'ID' columns. The appropriate columns and objects can be generated using the wrapper import functions <code>import_MaxQuant</code> and <code>import_IsobarQuant</code> or the generic functions <code>make_se</code> and <code>make_se_parse</code> .
<code>thr</code>	Integer(1), Sets the threshold for the allowed number of missing values per condition.
<code>fun</code>	"man", "bpca", "knn", "QRILC", "MLE", "MinDet", "MinProb", "min", "zero", "mixed" or "nbavg", Function used for data imputation based on <code>manual_impute</code> and <code>impute</code> .

... Additional arguments for imputation functions as depicted in [manual\\_impute](#) and [impute](#).

### Value

A filtered, normalized and imputed SummarizedExperiment object.

### Examples

```
# Load datasets
data <- UbiLength
exp_design <- UbiLength_ExpDesign

# Import data
se <- import_MaxQuant(data, exp_design)

# Process data
processed <- process(se)
```

**report**

*Generate a markdown report*

### Description

**report** generates a report of the analysis performed by [TMT](#) and [LFQ](#) wrapper functions. Additionally, the results table is saved as a tab-delimited file.

### Usage

```
report(results)
```

### Arguments

<b>results</b>	List of SummarizedExperiment objects obtained from the <a href="#">LFQ</a> or <a href="#">TMT</a> wrapper functions.
----------------	--

### Value

A [rmarkdown](#) report is generated and saved. Additionally, the results table is saved as a tab-delimited txt file.

### Examples

```
## Not run:

data <- UbiLength
expdesign <- UbiLength_ExpDesign

results <- LFQ(data, expdesign, 'MinProb', 'control', 'Ctrl')
report(results)

## End(Not run)
```

---

run\_app

*DEP shiny apps*

---

### Description

run\_app launches an interactive shiny app for interactive differential enrichment/expression analysis of proteomics data.

### Usage

```
run_app(app)
```

### Arguments

app                    'LFQ' or 'TMT', The name of the app.

### Value

Launches a browser with the shiny app

### Examples

```
## Not run:  
# Run the app  
run_app('LFQ')  
  
run_app('TMT')  
  
## End(Not run)
```

---

se2msn

*SummarizedExperiment to MSnSet object conversion*

---

### Description

se2msn generates a MSnSet object from a SummarizedExperiment object.

### Usage

```
se2msn(se)
```

### Arguments

se                    SummarizedExperiment, Object which will be turned into a MSnSet object.

### Value

A MSnSet object.

## Examples

```
# Load example
data <- UbiLength
data <- data[data$Reverse != "+" & data$Potential.contaminant != "+",]
data_unique <- make_unique(data, "Gene.names", "Protein.IDs", delim = ";")

# Make SummarizedExperiment
columns <- grep("LFQ.", colnames(data_unique))
exp_design <- UbiLength_ExpDesign
se <- make_se(data_unique, columns, exp_design)

# Convert to MSnSet
data_msn <- se2msn(se)
```

**test\_diff**

*Differential enrichment test*

## Description

`test_diff` performs a differential enrichment test based on protein-wise linear models and empirical Bayes statistics using [limma](#).

## Usage

```
test_diff(se, type = c("control", "all", "manual"), control = NULL,
          test = NULL, design_formula = formula(~0 + condition))
```

## Arguments

<code>se</code>	SummarizedExperiment, Proteomics data (output from <code>make_se()</code> or <code>make_se_parse()</code> ). It is advised to first remove proteins with too many missing values using <code>filter_missval()</code> , normalize the data using <code>normalize_vsn()</code> and impute remaining missing values using <code>impute()</code> .
<code>type</code>	"control", "all" or "manual", The type of contrasts that will be tested. This can be all possible pairwise comparisons ("all"), limited to the comparisons versus the control ("control"), or manually defined contrasts ("manual").
<code>control</code>	Character(1), The condition to which contrasts are generated if type = "control" (a control condition would be most appropriate).
<code>test</code>	Character, The contrasts that will be tested if type = "manual". These should be formatted as "SampleA_vs_SampleB" or c("SampleA_vs_SampleC", "SampleB_vs_SampleC").
<code>design_formula</code>	Formula, Used to create the design matrix.

## Value

A SummarizedExperiment object containing FDR estimates of differential expression.

## Examples

```
# Load example
data <- UbiLength
data <- data[data$Reverse != "+" & data$Potential.contaminant != "+",]
data_unique <- make_unique(data, "Gene.names", "Protein.IDs", delim = ";")

# Make SummarizedExperiment
columns <- grep("LFQ.", colnames(data_unique))
exp_design <- UbiLength_ExpDesign
se <- make_se(data_unique, columns, exp_design)

# Filter, normalize and impute missing values
filt <- filter_missval(se, thr = 0)
norm <- normalize_vsn(filt)
imputed <- impute(norm, fun = "MinProb", q = 0.01)

# Test for differentially expressed proteins
diff <- test_diff(imputed, "control", "Ctrl")
diff <- test_diff(imputed, "manual",
                  test = c("Ubi4_vs_Ctrl", "Ubi6_vs_Ctrl"))

# Test for differentially expressed proteins with a custom design formula
diff <- test_diff(imputed, "control", "Ctrl",
                  design_formula = formula(~ 0 + condition + replicate))
```

test\_gsea

*Gene Set Enrichment Analysis*

## Description

`test_gsea` tests for enriched gene sets in the differentially enriched proteins. This can be done independently for the different contrasts.

## Usage

```
test_gsea(dep, databases = c("GO_Molecular_Function_2017b",
                            "GO_Cellular_Component_2017b", "GO_Biological_Process_2017b"),
          contrasts = TRUE)
```

## Arguments

<code>dep</code>	SummarizedExperiment, Data object for which differentially enriched proteins are annotated (output from <code>test_diff()</code> and <code>add_rejections()</code> ).
<code>databases</code>	Character, Databases to search for gene set enrichment. See <a href="http://amp.pharm.mssm.edu/Enrichr/">http://amp.pharm.mssm.edu/Enrichr/</a> for available databases.
<code>contrasts</code>	Logical(1), Whether or not to perform the gene set enrichment analysis independently for the different contrasts.

## Value

A data.frame with enrichment terms (generated by `enrichr`)

## Examples

```
# Load example
data <- UbiLength
data <- data[data$Reverse != "+" & data$Potential.contaminant != "+",]
data_unique <- make_unique(data, "Gene.names", "Protein.IDs", delim = ";")

# Make SummarizedExperiment
columns <- grep("LFQ.", colnames(data_unique))
exp_design <- UbiLength_ExpDesign
se <- make_se(data_unique, columns, exp_design)

# Filter, normalize and impute missing values
filt <- filter_missval(se, thr = 0)
norm <- normalize_vsn(filt)
imputed <- impute(norm, fun = "MinProb", q = 0.01)

# Test for differentially expressed proteins
diff <- diff <- test_diff(imputed, "control", "Ctrl")
dep <- add_rejections(diff, alpha = 0.05, lfc = 1)

## Not run:

# Test enrichments
gsea_results_per_contrast <- test_gsea(dep)
gsea_results <- test_gsea(dep, contrasts = FALSE)

gsea_kegg <- test_gsea(dep, databases = "KEGG_2016")

## End(Not run)
```

**theme\_DEP1**

*DEP ggplot theme 1*

## Description

`theme_DEP1` is the default ggplot theme used for plotting in [DEP](#) with horizontal x-axis labels.

## Usage

```
theme_DEP1()
```

## Value

ggplot theme

## Examples

```
data <- UbiLength
data <- data[data$Reverse != '+' & data$Potential.contaminant != '+',]
data_unique <- make_unique(data, 'Gene.names', 'Protein.IDs', delim = ';')

columns <- grep('LFQ.', colnames(data_unique))
exp_design <- UbiLength_ExpDesign
```

---

```
se <- make_se(data_unique, columns, exp_design)

filt <- filter_missval(se, thr = 0)
plot_frequency(filt) # uses theme_DEP1() style
```

---

**theme\_DEP2***DEP ggplot theme 2***Description**

theme\_DEP2 is the ggplot theme used for plotting in DEP with vertical x-axis labels.

**Usage**

```
theme_DEP2()
```

**Value**

ggplot theme

**Examples**

```
data <- UbiLength
data <- data[data$Reverse != '+' & data$Potential.contaminant != '+',]
data_unique <- make_unique(data, 'Gene.names', 'Protein.IDs', delim = ';')

columns <- grep('LFQ.', colnames(data_unique))
exp_design <- UbiLength_ExpDesign
se <- make_se(data_unique, columns, exp_design)

filt <- filter_missval(se, thr = 0)
plot_numbers(filt) # uses theme_DEP2() style
```

---

**TMT***TMT workflow***Description**

TMT is a wrapper function running the entire differential enrichment/expression analysis workflow for TMT-based proteomics data. The protein table from IsobarQuant is used as direct input.

**Usage**

```
TMT(proteins, expdesign, fun = c("man", "bpca", "knn", "QRILC", "MLE",
"MinDet", "MinProb", "min", "zero", "mixed", "nbavg"), type = c("all",
"control", "manual"), control = NULL, test = NULL, name = "gene_name",
ids = "protein_id", alpha = 0.05, lfc = 1)
```

### Arguments

<code>proteins</code>	Data.frame, The data object.
<code>expdesign</code>	Data.frame, The experimental design object.
<code>fun</code>	"man", "bpca", "knn", "QRILC", "MLE", "MinDet", "MinProb", "min", "zero", "mixed" or "nbavg", Function used for data imputation based on <code>manual_impute</code> and <code>impute</code> .
<code>type</code>	'all', 'control' or 'manual', The type of contrasts that will be generated.
<code>control</code>	Character(1), The sample name to which the contrasts are generated (the control sample would be most appropriate).
<code>test</code>	Character, The contrasts that will be tested if type = "manual". These should be formatted as "SampleA_vs_SampleB" or c("SampleA_vs_SampleC", "SampleB_vs_SampleC").
<code>name</code>	Character(1), Name of the column representing gene names.
<code>ids</code>	'Character(1), Name of the column representing protein IDs.
<code>alpha</code>	Numeric(1), sets the false discovery rate threshold.
<code>lfc</code>	Numeric(1), sets the log fold change threshold.

### Value

A list of 8 objects:

<code>se</code>	SummarizedExperiment object containing the original data
<code>filt</code>	SummarizedExperiment object containing the filtered data
<code>norm</code>	SummarizedExperiment object containing the normalized data
<code>imputed</code>	SummarizedExperiment object containing the imputed data
<code>diff</code>	SummarizedExperiment object containing FDR estimates of differential expression
<code>dep</code>	SummarizedExperiment object annotated with logical columns indicating significant proteins
<code>results</code>	data.frame containing all results variables from the performed analysis
<code>param</code>	data.frame containing the test parameters

### Examples

```
## Not run:
TMT_res <- TMT()

## End(Not run)
```

---

UbiLength	<i>UbiLength - Ubiquitin interactors of different linear ubiquitin lengths (UbIA-MS dataset)</i>
-----------	--

---

## Description

The UbiLength dataset contains label free quantification (LFQ) data for ubiquitin interactors of different linear ubiquitin lengths, generated by Zhang et al 2017. The dataset contains the protein-groups output file from [MaxQuant](#).

## Usage

```
UbiLength
```

## Format

A data.frame with 3006 observations and 35 variables:

**Protein.IDs** Uniprot IDs

**Majority.protein.IDs** Uniprot IDs of major protein(s) in the protein group

**Protein.names** Full protein names

**Gene.names** Gene name

**Fasta.headers** Header as present in the Uniprot fasta file

**Peptides** Number of peptides identified for this protein group

**Razor..unique.peptides** Number of peptides used for the quantification of this protein group

**Unique.peptides** Number of peptides identified which are unique for this protein group

**Intensity columns (12)** Raw mass spectrometry intensity, A.U.

**LFQ.intensity columns (12)** LFQ normalized mass spectrometry intensity, A.U.

**Only.identified.by.site** The protein is only identified by a modification site if marked ('+')

**Reverse** The protein is identified in the decoy database if marked ('+')

**Potential.contaminant** The protein is a known contaminant if marked ('+')

## Value

A data.frame.

## Source

Zhang, Smits, van Tilburg, et al (2017). An interaction landscape of ubiquitin signaling. Molecular Cell 65(5): 941-955. doi: [10.1016/j.molcel.2017.01.004](https://doi.org/10.1016/j.molcel.2017.01.004).

---

**UbiLength\_ExpDesign**    *Experimental design of the UbiLength dataset*

---

**Description**

The `UbiLength_ExpDesign` object annotates 12 different samples of the UbiLength dataset in 4 conditions and 3 replicates.

**Usage**

```
UbiLength_ExpDesign
```

**Format**

A data.frame with 12 observations and 3 variables:

**label** Label names

**condition** Experimental conditions

**replicate** Replicate number

**Value**

A data.frame.

**Source**

Zhang, Smits, van Tilburg, et al (2017). An interaction landscape of ubiquitin signaling. Molecular Cell 65(5): 941-955. doi: [10.1016/j.molcel.2017.01.004](https://doi.org/10.1016/j.molcel.2017.01.004).

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