# Package 'iq'

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Type Package

```
Title Protein Quantification in Mass Spectrometry-Based Proteomics
Version 1.9.7
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Description An implementation of the MaxLFQ algorithm by
     Cox et al. (2014) <doi:10.1074/mcp.M113.031591> in a comprehensive
     pipeline for processing proteomics data in data-independent acquisition mode
     (Pham et al. 2020 <doi:10.1093/bioinformatics/btz961>).
     It offers additional options for protein quantification using
     the N most intense fragment ions, using all fragment ions, and
     a wrapper for the median polish algorithm by Tukey (1977, ISBN:0201076160).
     In general, the tool can be used to integrate multiple
     proportional observations into a single quantitative value.
Depends R (>= 2.10)
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```

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# Description

Index

For each protein, a numerical matrix is formed where the columns are samples and rows are fragment ions.

## Usage

```
create_protein_list(preprocessed_data)
```

## **Arguments**

preprocessed\_data

A data frame of four components as output of the preprocess function.

## Value

A list where each element contains the quantitative data of a protein. The column names are sample names and the row names fragment ions.

## Author(s)

Thang V. Pham

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## References

Pham TV, Henneman AA, Jimenez CR. iq: an R package to estimate relative protein abundances from ion quantification in DIA-MS-based proteomics. *Bioinformatics* 2020 Apr 15;36(8):2611-2613.

## See Also

```
preprocess
```

## **Examples**

```
data("spikeins")
head(spikeins)
# This example set of spike-in proteins has been 'median-normalized'.
norm_data <- iq::preprocess(spikeins, median_normalization = FALSE, pdf_out = NULL)
protein_list <- iq::create_protein_list(norm_data)</pre>
```

create\_protein\_table Protein quantification for a list of proteins

## Description

Travels through the input list and quantifies all proteins one by one.

# Usage

```
create_protein_table(protein_list, method = "maxLFQ", ...)
```

# Arguments

method Possible values are "maxLFQ", "median\_polish", "topN", and "meanInt".

... Additional parameters for individual quantitation methods.

## Value

A list of two components is returned

estimate A table of protein abundances for all samples.

annotation A vector of annotations, one for each protein.

## Author(s)

Thang V. Pham

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## References

Pham TV, Henneman AA, Jimenez CR. iq: an R package to estimate relative protein abundances from ion quantification in DIA-MS-based proteomics. *Bioinformatics* 2020 Apr 15;36(8):2611-2613.

## See Also

```
create_protein_list, maxLFQ, median_polish, topN, meanInt
```

## **Examples**

```
data("spikeins")
# This example set of spike-in proteins has been 'median-normalized'.
norm_data <- iq::preprocess(spikeins, median_normalization = FALSE, pdf_out = NULL)
protein_list <- iq::create_protein_list(norm_data)
result <- iq::create_protein_table(protein_list)
head(result)</pre>
```

extract\_annotation

Protein annotation extraction

## **Description**

Extracts annotation columns from a long-format input

# Usage

# Arguments

protein\_ids A vector of protein ids.

quant\_table A long-format input table. The input is typically the same as input to the

preprocess function.

primary\_id The column containing protein ids.

annotation\_columns

A vector of columns for annotation.

## Value

A table of proteins and associated annotation extracted from the input.

## Author(s)

Thang V. Pham

fast\_MaxLFQ 5

## References

Pham TV, Henneman AA, Jimenez CR. iq: an R package to estimate relative protein abundances from ion quantification in DIA-MS-based proteomics. *Bioinformatics* 2020 Apr 15;36(8):2611-2613.

#### See Also

preprocess

## **Examples**

fast\_MaxLFQ

The MaxLFQ algorithm

#### **Description**

A fast implementation of the MaxLFQ algorithm.

#### Usage

```
fast_MaxLFQ(norm_data, row_names = NULL, col_names = NULL)
```

# **Arguments**

norm\_data A list of four vectors with equal length protein\_list, sample\_list, id and

quant as prepared by the fast\_preprocess function or the quant\_table component returned by the fast\_read function. Note that quant should contain

log2 intensities.

row\_names A vector of character strings for row names. If NULL, unique values in the

protein\_list component of norm\_data will be used. Otherwise, it should

be the sample component returned by the fast\_read.

col\_names A vector of character strings for column names. If NULL, unique values in the

sample\_list component of norm\_data will be used. Otherwise, it should be

the sample component returned by the fast\_read.

#### Value

A list is returned with two components

estimate A quantification result table.

annotation A vector of strings indicating membership in case of multiple connected com-

ponents for each row of estimate.

fast\_preprocess

## Author(s)

Thang V. Pham

#### References

Pham TV, Henneman AA, Jimenez CR. iq: an R package to estimate relative protein abundances from ion quantification in DIA-MS-based proteomics. *Bioinformatics* 2020 Apr 15;36(8):2611-2613.

#### See Also

```
fast_read, fast_preprocess
```

fast\_preprocess

Data filtering and normalization

## **Description**

Filters out low intensities and performs median normalization.

## Usage

## Arguments

quant\_table The quant\_table component as returned by fast\_read.
median\_normalization

A logical value. The default TRUE value is to perform median normalization.

log2\_intensity\_cutoff

Entries lower than this value in log2 space are ignored. Plot a histogram of all

intensities to set this parameter.

pdf\_out A character string specifying the name of the PDF output. A NULL value will

suppress the PDF output.

pdf\_width Width of the pdf output in inches.
pdf\_height Height of the pdf output in inches.

show\_boxplot A logical value. The default TRUE value is to create boxplots of fragment inten-

sities for each sample.

fast\_read 7

#### Value

A list is returned with the same components as input data in which low intensities are filtered out and median normalization is performed if requested.

## Author(s)

Thang V. Pham

#### References

Pham TV, Henneman AA, Jimenez CR. iq: an R package to estimate relative protein abundances from ion quantification in DIA-MS-based proteomics. *Bioinformatics* 2020 Apr 15;36(8):2611-2613.

#### See Also

fast\_read

fast\_read

Reading data from an input file

## Description

A highly efficient reading of a tab-separated text file for iq processing.

## Usage

#### **Arguments**

filename

A long-format tab-separated text file with a primary column of protein identification, secondary columns of fragment ions, a column of sample names, a column for quantitative intensities, and extra columns for annotation.

primary\_id

Unique values in this column form the list of proteins to be quantified.

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secondary\_id A concatenation of these columns determines the fragment ions used for quan-

tification.

sample\_id Unique values in this column form the list of samples.

intensity\_col The column for intensities.

annotation\_col Annotation columns

filter\_string\_equal

A named vector of strings. Only rows satisfying the condition are kept.

filter\_string\_not\_equal

A named vector of strings. Only rows satisfying the condition are kept.

filter\_double\_less

A named vector of strings. Only rows satisfying the condition are kept. Default PG.Qvalue < 0.01 and EG.Qvalue < 0.01.

filter\_double\_greater

A named vector of strings. Only rows satisfying the condition are kept.

intensity\_col\_sep

A separator character when entries in the intensity column contain multiple values

intensity\_col\_id

The column for identities of multiple quantitative values.

na\_string The value considered as NA.

#### **Details**

When entries in the intensity column contain multiple values, this function will replicate entries in other column and the secondary\_id will be appended with corresponding entries in intensity\_col\_id when it is provided. Otherwise, integer values 1, 2, 3, etc... will be used.

## Value

A list is returned with following components

protein A table of proteins in the first column followed by annotation columns.

sample A vector of samples.

ion A vector of fragment ions to be used for quantification.

quant\_table A list of four components: protein\_list (index pointing to protein)), sample\_list

(index pointing to sample), id (index pointing to ion), and quant (intensities).

# Author(s)

Thang V. Pham

# References

maxLFQ 9

maxLFQ	The MaxLFQ algorithm for protein quantification

## **Description**

Estimates protein abundances by aiming to maintain the fragment intensity ratios between samples.

## Usage

maxLFQ(X)

# Arguments

X A matrix of ion intensities in log2 space. Columns are samples and rows are

fragment ions.

#### Value

A list of two components is returned

estimate A vector with length equal to the number of columns of the input containing the

protein abundances.

annotation An empty string if all quantified samples are connected. Otherwise, a string of

membership of the connected components is returned.

## Author(s)

Thang V. Pham

## References

Cox J, Hein MY, Luber CA, et al. Accurate proteome-wide label-free quantification by delayed normalization and maximal peptide ratio extraction, termed MaxLFQ. *Mol Cell Proteomics*. 2014;13(9):2513–2526.

10 meanInt

meanInt

The meanInt algorithm for protein quantification

# Description

Estimates protein abundances by averaging all associated ion intensities

## Usage

```
meanInt(X, aggregation_in_log_space = TRUE)
```

## **Arguments**

A matrix of ion intensities in log2 space. Columns are samples and rows are fragment ions.

aggregation\_in\_log\_space

A logical value. If FALSE, the data aggregation is performed in the original intensity space.

#### Value

A list of two components is returned

estimate A vector with length equal to the number of columns of the input containing the

protein abundances.

annotation Reserved, currently an empty string.

## Author(s)

Thang V. Pham

## References

median\_polish 11

median_polish	A wrapper for the R implementation of the median polish algorithm

## **Description**

Estimates protein abundances using the Tukey median polish algorithm.

## Usage

```
median_polish(X)
```

## **Arguments**

X A matrix of ion intensities in log2 space. Columns are samples and rows are

fragment ions.

#### Value

A list of two components is returned

estimate A vector with length equal to the number of columns of the input containing the

protein abundances.

annotation Reserved, currently an empty string

# Author(s)

Thang V. Pham

## References

Pham TV, Henneman AA, Jimenez CR. iq: an R package to estimate relative protein abundances from ion quantification in DIA-MS-based proteomics. *Bioinformatics* 2020 Apr 15;36(8):2611-2613.

Tukey JW. Exploratory Data Analysis, Reading Massachusetts: Addison-Wesley, 1977.

plot_protein	Plotting the underlying quantitative data for a protein
•	

## **Description**

Displays the underlying data for a protein.

```
plot_protein(X, main = "", col = NULL, split = 0.6, ...)
```

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## **Arguments**

Χ	Protein data matrix.
main	Title of the plot.
col	Colors of the rows of the data matrix.
split	Fraction of the plotting area for the main figure. The remaining one is for legend. Set this parameter to NULL to ignore the legend area.
	Additional parameters for plotting.

#### Value

A NULL value is returned.

## Author(s)

Thang V. Pham

#### References

Pham TV, Henneman AA, Jimenez CR. iq: an R package to estimate relative protein abundances from ion quantification in DIA-MS-based proteomics. *Bioinformatics* 2020 Apr 15;36(8):2611-2613.

# **Examples**

```
data("spikeins")
head(spikeins)
# This example set of spike-in proteins has been 'median-normalized'.
norm_data <- iq::preprocess(spikeins, median_normalization = FALSE, pdf_out = NULL)
protein_list <- iq::create_protein_list(norm_data)
iq::plot_protein(protein_list$P00366, main = "Protein P00366", split = NULL)</pre>
```

preprocess

Data preprocessing for protein quantification

# Description

Prepares a long-format input including removing low-intensity ions and performing median normalization.

preprocess 13

```
median_normalization = TRUE,
log2_intensity_cutoff = 0,
pdf_out = "qc-plots.pdf",
pdf_width = 12,
pdf_height = 8,
intensity_col_sep = NULL,
intensity_col_id = NULL,
na_string = "0")
```

## Arguments

quant\_table A long-format table with a primary column of protein identification, secondary

columns of fragment ions, a column of sample names, and a column for quanti-

tative intensities.

primary\_id Unique values in this column form the list of proteins to be quantified.

secondary\_id A concatenation of these columns determines the fragment ions used for quan-

tification.

sample\_id Unique values in this column form the list of samples.

intensity\_col The column for intensities.

median\_normalization

A logical value. The default TRUE value is to perform median normalization.

log2\_intensity\_cutoff

Entries lower than this value in log2 space are ignored. Plot a histogram of all

intensities to set this parameter.

pdf\_out A character string specifying the name of the PDF output. A NULL value will

suppress the PDF output.

pdf\_width Width of the pdf output in inches.

pdf\_height Height of the pdf output in inches.

intensity\_col\_sep

A separator character when entries in the intensity column contain multiple val-

intensity\_col\_id

The column for identities of multiple quantitative values.

na\_string The value considered as NA.

#### **Details**

When entries in the intensity column contain multiple values, this function will replicate entries in other column and the secondary\_id will be appended with corresponding entries in intensity\_col\_id when it is provided. Otherwise, integer values 1, 2, 3, etc... will be used.

## Value

A data frame is returned with following components

protein\_list A vector of proteins.

sample\_list A vector of samples.

id A vector of fragment ions to be used for quantification.

quant A vector of log2 intensities.

## Author(s)

Thang V. Pham

#### References

Pham TV, Henneman AA, Jimenez CR. iq: an R package to estimate relative protein abundances from ion quantification in DIA-MS-based proteomics. *Bioinformatics* 2020 Apr 15;36(8):2611-2613.

## **Examples**

```
data("spikeins")
head(spikeins)
# This example set of spike-in proteins has been 'median-normalized'.
norm_data <- iq::preprocess(spikeins, median_normalization = FALSE, pdf_out = NULL)</pre>
```

process\_long\_format

Long format to a wide format table using the MaxLFQ algorithm

# Description

A convenient function combining multiple steps to process a long format table using the MaxLFQ algorithm.

process\_long\_format 15

```
normalization = "median",
log2_intensity_cutoff = 0,
pdf_out = "qc-plots.pdf",
pdf_width = 12,
pdf_height = 8,
show_boxplot = TRUE,
peptide_extractor = NULL)
```

#### Arguments

```
input_filename See filename in fast_read.
output_filename
                Output filename.
sample_id
                See sample_id in fast_read.
primary_id
                See primary_id in fast_read.
secondary_id
                See secondary_id in fast_read.
intensity_col
                See intensity_col in fast_read.
annotation_col See annotation_col in fast_read.
filter_string_equal
                See filter_string_equal in fast_read.
filter_string_not_equal
                See filter_string_not_equal in fast_read.
filter_double_less
                See filter_double_less in fast_read.
filter_double_greater
                See filter_double_greater in fast_read.
intensity_col_sep
                See intensity_col_sep in fast_read.
intensity_col_id
                See intensity_col_id in fast_read.
na_string
                See intensity_col_id in fast_read.
normalization
                Normalization type. Possible values are median and none. The default value
                median is for median normalization in fast_preprocess.
log2_intensity_cutoff
                See log2_intensity_cutoff in fast_preprocess.
pdf_out
                See pdf_out in fast_preprocess.
pdf_width
                See pdf_width in fast_preprocess.
pdf_height
                See pdf_height in fast_preprocess.
show_boxplot
                See show_boxplot in fast_preprocess.
peptide_extractor
                A function to parse peptides.
```

process\_wide\_format

## Value

After processing with fast\_read, fast\_preprocess, and fast\_MaxLFQ, the result table is written to output\_filename. A NULL value is returned. If peptide\_extractor is not NULL, fragment statistics for each protein will be calculated based on the result of the extractor function. Counting the number of peptides contributing to a protein is possible using an appropriate extractor function. An example value for peptide\_extractor is function(x) gsub("[0-9].\*\$", "", x), which removes the charge state and fragment descriptors in an ion descriptor to obtain unique peptide sequences. One can examine the ion component returned by the fast\_read function to derive a regular expression to be used in the gsub function above.

## Author(s)

Thang V. Pham

#### References

Pham TV, Henneman AA, Jimenez CR. iq: an R package to estimate relative protein abundances from ion quantification in DIA-MS-based proteomics. *Bioinformatics* 2020 Apr 15;36(8):2611-2613.

#### See Also

```
fast_read, fast_preprocess, fast_MaxLFQ
```

process\_wide\_format

Merging rows with identical values in a particular column in a table

## **Description**

Collapses rows with identical values in a particular column in a table. When the values in each row are proportional such as intensities of multiple fragments of a protein, the MaxLFQ algorithm is recommended.

process\_wide\_format 17

# Arguments

input\_filename Input filename of a tab-separated value text file.

output\_filename

Output filename.

id\_column The column where unique values will be kept. Rows with identical values in

this column are merged. Rows with empty values here are removed.

quant\_columns Columns containing numerical data to be merged.

data\_in\_log\_space

A logical value. If FALSE, the numerical data will be log2-transformed.

annotation\_columns

Columns in the input file apart from id\_column and quant\_columns that will

be kept in the output.

method Method for merging. Default value is "maxLFQ". Possible values are "maxLFQ",

"maxLFQ\_R", "median\_polish", "top3", "top5", "meanInt", "maxInt", "sum", "least\_na" and any function for collapsing a numerical matrix to a row vector.

#### **Details**

Method "maxLFQ\_R" implements the MaxLFQ algorithm pure R. It is slower than "maxLFQ".

Method "maxInt" selects row with maximum intensity (top 1).

Method "sum" sum all intensities.

Method "least\_na" selects row with the least number of missing values.

The value of method can be a function such as  $function(x) log2(colSums(2^x, na.rm = TRUE))$  for summing all intensities in the original space.

## Value

The result table is written to output\_filename. A NULL value is returned.

#### Author(s)

Thang V. Pham

#### References

topN

spikeins

An example dataset of 12 spike-in proteins

# Description

A subset of the Bruderer 2015 dataset containing 12 spike-in proteins. The full dataset was exported from the Spectronaut software. The complete dataset has been median-normalized.

## Usage

```
data("spikeins")
```

#### **Format**

A data frame with 18189 observations on the following 9 variables.

R.Condition Sample names.

PG.ProteinGroups Protein identifiers.

EG. Modified Sequence Sequence of the fragment ions.

FG. Charge Fragment group charge.

F.FrgIon Fragment ions.

F. Charge Fragment charges.

F. PeakArea Quantitative values.

PG. Genes Gene names.

PG. ProteinNames Protein names.

# **Examples**

```
data("spikeins")
head(spikeins)
```

topN

The topN algorithm for protein quantification

# Description

Estimates protein abundances using the N most intense ions.

```
topN(X, N = 3, aggregation_in_log_space = TRUE)
```

topN

## **Arguments**

X A matrix of ion intensities in log2 space. Columns are samples and rows are

fragment ions.

N The number of top ions used for quantification.

aggregation\_in\_log\_space

A logical value. If FALSE, data aggregation is performed in the original intensity

space.

## Value

A list of two components is returned

estimate A vector with length equal to the number of columns of the input containing the

protein abundances.

annotation Reserved, currently an empty string.

## Author(s)

Thang V. Pham

## References

# **Index**

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