# Package 'SWATH2stats'

June 18, 2018

Type Package			
Title Transform and Filter SWATH Data for Statistical Packages			
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<b>Description</b> This package is intended to transform SWATH data from the OpenSWATH software into a format readable by other statistics packages while performing filtering, annotation and FDR estimation.			
License GPL-3			
<b>Depends</b> $R(>=2.10.0)$			
Imports data.table, reshape2, grid, ggplot2, stats, grDevices, graphics, utils, biomaRt			
Suggests testthat, aLFQ, knitr, PECA			
Enhances imsbInfer, MSstats			
<b>biocViews</b> Proteomics, Annotation, ExperimentalDesign, Preprocessing, MassSpectrometry			
NeedsCompilation no			
VignetteBuilder knitr			
RoxygenNote 6.0.1			
R topics documented:			
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	set.

## Description

Gather gene symbols from biomart and add them to a swath2stats data set.

## Usage

```
add_genesymbol(data_table, gene.ID.table, column.name = "protein",
   ID1 = "uniprotswissprot", ID2 = "hgnc_symbol", id.separator = "/",
   copy_nonconverted = TRUE)
```

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#### **Arguments**

data\_table A data frame or file name. gene.ID.table A table to match against

column.name The column name where the original protein identifiers are present.

The type of the original protein identifiers (e.g., "uniprotswissprot", "ensembl\_peptide\_id").

The type of the converted protein identifiers (e.g. "hgnc\_symbol", "mgi\_symbol",

"external\_gene\_name").

id. separator Separator between protein identifiers of shared peptides.

copy\_nonconverted

Option defining if the identifiers that cannot be converted should be copied.

#### Value

some gene symbols

#### Note

Protein identifiers from shared peptides should be separated by a forward slash. The host of archived ensembl databases can be introduced as well (e.g. "dec2017.archive.ensembl.org")

#### Author(s)

Peter Blattmann

assess\_decoy\_rate

Query a swath2stats data set for decoy hit rates.

## Description

This looks at the swath2stats table and compares the length of the detected decoy peptides vs non-decoy peptides.

#### Usage

```
assess_decoy_rate(data)
```

## **Arguments**

data A data frame that contains at least a column named "FullPeptideName" and

"decoy".

#### **Details**

A printout is generated to indicate the number of non-decoy, decoy peptides and the rate of decoy vs non-decoy peptides. Unique peptides are counted, so a precursor with different charge states is counted as one peptide. In the column "decoy" the values need to be 1,0 or TRUE and FALSE.

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#### Value

list containing the number of decoys, non-decoys, and the ratio.

#### Author(s)

Peter Blattmann

#### **Examples**

```
## Not run:
  data("OpenSWATH_data", package="SWATH2stats")
  data <- OpenSWATH_data
  assess_decoy_rate(data)
## End(Not run)</pre>
```

assess\_fdr\_byrun

Assess assay, peptide and protein level FDR by run (for each MS\_injection separately) in OpenSWATH output table

## Description

This function estimates the assay, peptide and protein FDR by run in an OpenSWATH result table in dependence of a range of m\_score cutoffs. The results can be visualized and summarized by the associated method plot.fdr\_table(). It counts target and decoy assays (unique transition\_group\_id), peptides (unique FullPeptideName) and proteins (unique ProteinName) in the OpenSWATH output table in dependence of m-score cutoff, the useful m\_score cutoff range is evaluated for each dataset individually on the fly. To arrive from decoy counts at an estimation of the false discovery rate (false positives among the targets remaining at a given mscore cutoff) the ratio of false positives to true negatives (decoys) (FFT) must be supplied. It is estimated for each run individually by pyProphet and contained in the pyProphet statistics [Injection\_name]\_full\_stat.csv. As an approximation, the FFTs of multiple runs are averaged and supplied as argument FFT. For further details see the Vignette Section 1.3 and 4.1. To assess fdr over the entire dataset, please refer to function assess\_fdr\_overall. FDR is calculated as FDR = (TN\*FFT/T); TN=decoys, T=targets, FFT=see above.

## Usage

```
assess_fdr_byrun(data, FFT = 1, n_range = 20, output = "pdf_csv",
plot = TRUE, filename = "FDR_report_byrun",
output_mscore_levels = c(0.01, 0.001))
```

#### **Arguments**

data

Annotated OpenSWATH/pyProphet output table. Refer to function sample\_annotation from this package for further information.

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**FFT** Ratio of false positives to true negatives, q-values from [Injection\_name]\_full\_stat.csv in pyProphet stats output. As an approximation, the q-values of multiple runs are averaged and supplied as argument FFT. Numeric from 0 to 1. Defaults to 1, the most conservative value (1 Decoy indicates 1 False target). Option to set the number of magnitude for which the m\_score threshold is den\_range creased (e.g. n.range = 10, m-score from 0.1 until  $10^{-10}$ ). Choose output type. "pdf\_csv" creates the output as files in the working direcoutput tory, "Rconsole" triggers delivery of the output to the console enabling further computation or custom plotting / output. plot Logical, whether or not to create plots from the results (using the associated method plot.fdr cube() output\_mscore\_levels Define m-score levels to plot and write the estimated FDR results. Optional, modifying the basename of the result files if applicable.

#### Value

Returns an array of target/decoy identification numbers and calculated FDR values at different mscore cutoffs.

#### Author(s)

Moritz Heusel

#### **Examples**

assess\_fdr\_overall

Assess overall FDR in annotated OpenSWATH/pyProphet output table in dependence of m\_score cutoff

## **Description**

This function estimates the assay, peptide and protein FDR over a multi-run OpenSWATH/pyProphet output table. It counts target and decoy assays (unique transition\_group\_id), peptides (unique FullPeptideName) and proteins (unique ProteinName) in dependence of the m-score cutoff (1e-2 to 1e-20). To arrive from decoy counts at an estimation of the false discovery rate (false positives among the targets remaining at a given mscore cutoff) the ratio of false positives to true negatives

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(decoys) (FFT) must be supplied. It is estimated for each run individually by pyProphet and contained in the pyProphet statistics [Injection\_name]\_full\_stat.csv. As an approximation, the FFTs of multiple runs are averaged and supplied as argument FFT. For further details see the Vignette Section 1.3 and 4.1. Protein FDR control on peak group quality level is a very strict filter and should be handled with caution. FDR is calculated as FDR = (TN\*FFT/T); TN=decoys, T=targets, FFT=see above

## Usage

```
assess_fdr_overall(data, FFT = 1, n_range = 20, output = "pdf_csv",
plot = TRUE, filename = "FDR_report_overall")
```

#### **Arguments**

data	Data table that is produced by the OpenSWATH/pyProphet workflow
FFT	Ratio of false positives to true negatives, q-values from [Injection_name]_full_stat.csv in pyProphet stats output. As an approximation, the q-values of multiple runs are averaged and supplied as argument FFT. Numeric from 0 to 1. Defaults to 1, the most conservative value (1 Decoy indicates 1 False target).
n_range	I am also not certain what this is, nor why 20 is the optimal default value, but I think the idea is to set up a series of mscore thresholds.
output	Choose output type. "pdf_csv" creates the output as files in the working directory, "Rconsole" triggers delivery of the output to the console enabling further computation or custom plotting / output.
plot	Logical, whether or not to create plots from the results (using the associated method plot.fdr_table()
filename	Optional, modifying the basename of the result files if applicable.

## Value

Returns a list of class "fdr\_table". If output "pdf\_csv" and plot = TRUE were chosen, report files are written to the working folder.

## Author(s)

Moritz Heusel

base\_size 7

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The following is used to set the ggplot2 default text size.

## **Description**

The following is used to set the ggplot2 default text size.

## Usage

base\_size

#### **Format**

An object of class numeric of length 1.

convert\_aLFQ

Convert table into the format for aLFQ

## **Description**

This functions selects the columns necessary for the aLFQ R package.

#### Usage

```
convert_aLFQ(data, annotation = TRUE, check_transitions = TRUE)
```

## Arguments

data A data frame containing the SWATH data in transition-level format

annotation Option to indicate if the data has been annotated, i.e. if the columns Condition,

Replicate, Run are present. If option is set to true it will write a new run\_id as a

string of the combination of these three columns.

check\_transitions

if number of transitions should be checked. As input only transition-level data should be used and therefore this is checked. However, this makes the function

slow and herewith be omitted.

## Value

Returns a data frame in the appropriate format for aLFQ.

## Author(s)

Peter Blattmann

8 convert\_mapDIA

#### References

Rosenberger G, Ludwig C, Rost HL, Aebersold R, Malmstrom L. aLFQ: an R-package for estimating absolute protein quantities from label-free LC-MS/MS proteomics data. Bioinformatics. 2014 Sep 1;30(17):2511-3. doi: 10.1093/bioinformatics/btu200.

## **Examples**

```
## Not run:
data("OpenSWATH_data", package="SWATH2stats")
data("Study_design", package="SWATH2stats")
data <- sample_annotation(OpenSWATH_data, Study_design)
data.filtered.decoy <- filter_mscore(data, 0.01)
raw <- disaggregate(data.filtered.decoy)
data.aLFQ <- convert4aLFQ(raw)
## End(Not run)</pre>
```

convert\_mapDIA

Convert table into the format for mapDIA

## Description

This functions selects the columns necessary for mapDIA.

## Usage

```
convert_mapDIA(data, RT = FALSE)
```

#### **Arguments**

data A data frame containing SWATH data.

RT Option to export the retention times.

#### Value

Returns a data frame in the appropriate format for mapDIA.

#### Note

The table must not contain any technical replica, the intensity of technical replica is averaged. This function requires the package reshape2.

#### Author(s)

Peter Blattmann

convert\_MSstats 9

#### References

Teo, G., et al. (2015). "mapDIA: Preprocessing and statistical analysis of quantitative proteomics data from data independent acquisition mass spectrometry." J Proteomics 129: 108-120.

#### **Examples**

```
## Not run:
    data("OpenSWATH_data", package="SWATH2stats")
    data("Study_design", package="SWATH2stats")
    data <- sample_annotation(OpenSWATH_data, Study_design)
    data.filtered.decoy <- filter_mscore(data, 0.01)
    raw <- disaggregate(data.filtered.decoy)
    data.mapDIA <- convert4mapDIA(raw, RT=TRUE)

## End(Not run)

convert_MSstats

Take data from the swath2stats format and get it ready for use by
    MSstats.</pre>
```

#### **Description**

Though the two tools use very similar formats, some coercion is required to Convert table into the format for MSstats.

#### Usage

```
convert_MSstats(data, replace_values = TRUE, replace_colnames = TRUE,
  replace_unimod = TRUE)
```

## **Arguments**

data A data frame containing SWATH data.

replace\_values Option to indicate if negative and 0 values should be replaced with NA. replace\_colnames

Option to indicate if column names should be renamed and columns reduced to the necessary columns for MSstats.

replace\_unimod Option to indicate if Unimod Identifier should be replaced from ":" to "\_".

#### **Details**

This functions selects the columns necessary for MSstats and renames them if necessary.

The necessary columns are selected and three columns renamed: FullPeptideName -> PeptideSequence Charge -> PrecursorCharge filename -> File

#### Value

Returns a data frame in the appropriate format for MSstats.

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#### Author(s)

Peter Blattmann

#### References

Choi M, Chang CY, Clough T, Broudy D, Killeen T, MacLean B, Vitek O. MSstats: an R package for statistical analysis of quantitative mass spectrometry-based proteomic experiments. Bioinformatics. 2014 Sep 1;30(17):2524-6. doi: 10.1093/bioinformatics/btu305.

## **Examples**

```
## Not run:
data("OpenSWATH_data", package="SWATH2stats")
data("Study_design", package="SWATH2stats")
data <- sample_annotation(OpenSWATH_data, Study_design)
data.filtered.decoy <- filter_mscore(data, 0.01)
raw <- disaggregate(data.filtered.decoy)
data.mapDIA <- convert4MSstats(raw)
## End(Not run)</pre>
```

convert\_PECA

Convert table into the format for ROPECA

## **Description**

This functions selects the columns necessary for ROPECA.

## Usage

```
convert_PECA(data)
```

#### **Arguments**

data

A data frame containing SWATH data.

#### Value

Returns a data frame in the appropriate format for ROPECA.

#### Note

The table must not contain any technical replica, the intensity of technical replica is averaged. This function requires the package reshape2.

#### Author(s)

Peter Blattmann

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#### References

Suomi, T. and Elo L.L. (2017). "Enhanced differential expression statistics for data-independent acquisition proteomics" Scientific Reports 7, Article number: 5869.doi:10.1038/s41598-017-05949-y

#### **Examples**

```
## Not run:
data("OpenSWATH_data", package="SWATH2stats")
data("Study_design", package="SWATH2stats")
data <- sample_annotation(OpenSWATH_data, Study_design)
data.filtered.decoy <- filter_mscore(data, 0.01)
data.PECA <- convert4PECA(data.filtered.decoy)
## End(Not run)</pre>
```

convert\_protein\_ids

Convert protein ids

## **Description**

This function renames protein ids in a data frame or file

## Usage

```
convert_protein_ids(data_table, column.name = "Protein",
   species = "hsapiens_gene_ensembl", host = "www.ensembl.org",
   mart = "ENSEMBL_MART_ENSEMBL", ID1 = "uniprotswissprot",
   ID2 = "hgnc_symbol", id.separator = "/", copy_nonconverted = TRUE,
   verbose = FALSE)
```

#### **Arguments**

verbose

data_table	A data frame or file name.
column.name	The column name where the original protein identifiers are present.
species	The species of the protein identifiers in the term used by biomaRt (e.g. "hsapiens_gene_ensembl", "mmusculus_gene_ensembl", "drerio_gene_ensembl", etc.)
host	Path of the biomaRt database (e.g. "www.ensembl.org", "dec2017.archive.ensembl.org").
mart	The type of mart (e.g. "ENSEMBL_MART_ENSEMBL", etc.)
ID1	The type of the original protein identifiers (e.g. "uniprotswissprot", "ensembl_peptide_id").
ID2	The type of the converted protein identifiers (e.g. "hgnc_symbol", "mgi_symbol", "external_gene_name").
id.separator	Separator between protein identifiers of shared peptides.
copy_nonconverted	
	Option defining if the identifiers that cannot be converted should be copied.

Option to write a file containing the version of the database used.

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#### Value

The data frame with an added column of the converted protein identifiers.

#### Note

Protein identifiers from shared peptides should be separated by a forward slash. The host of archived ensembl databases can be introduced as well (e.g. "dec2017.archive.ensembl.org")

#### Author(s)

Peter Blattmann

## **Examples**

convert\_python

Convert data into the format for running a python script.

## **Description**

This functions selects the columns suggested to run a python script to change the data from peptide-level to transition-level.

#### Usage

```
convert_python(data, replace.Unimod = TRUE)
```

## Arguments

```
data A data frame containing SWATH data.

replace.Unimod Option to indicate if Unimod Identifier should be replaced form ":"" to "_".
```

## **Details**

The necessary columns are selected and the run column is renamed to filename for the script. The intensities are taken from the column aggr\_Peak\_Area and therefore the Intensity column is not exported.

#### Value

Returns a data frame in the appropriate format to be used by a custom python script stored in the scripts folder.

count\_analytes 13

#### Author(s)

Peter Blattmann

## **Examples**

```
## Not run:
    data("OpenSWATH_data", package="SWATH2stats")
    data("Study_design", package="SWATH2stats")
    data <- sample_annotation(OpenSWATH_data, Study_design)
    data.filtered.decoy <- filter_mscore(data,0.01)
    data.pythonscript <- convert4pythonscript(data.filtered.decoy)
## End(Not run)</pre>
```

count\_analytes

Counts analytes in different injections

## Description

This functions counts the number of different peakgroups, peptides and proteins in different injections.

## Usage

```
count_analytes(data, column_levels = c("transition_group_id",
   "fullpeptidename", "proteinname"), column_by = "run_id", rm_decoy = TRUE)
```

## **Arguments**

data A data frame containing SWATH data.

column\_by Column for which the different identifiers should be counted for, e.g. for the

different injections.

rm\_decoy Option to not remove decoy before counting.

#### Value

Returns a data frame with the count of the different identifiers per e.g. injection.

#### Author(s)

Peter Blattmann

14 disaggregate

#### **Examples**

```
## Not run:
    data("OpenSWATH_data", package="SWATH2stats")
    data("Study_design", package="SWATH2stats")
    data <- sample_annotation(OpenSWATH_data, Study_design)
    count_analytes(data)
## End(Not run)</pre>
```

disaggregate

Transforms the SWATH data from a peptide- to a transition-level table.

## Description

If the SWATH data should be analyzed on transition-level the data needs to be tranformed from peptide-level table to a transition-level table (one row per transition instead of one row per peptide). The columns "aggr\_Fragment\_Annotation" and "aggr\_Peak\_Area" are disaggregated into the new columns "FragmentIon" and "Intensity". The following columns are renamed if they exist: FullPeptideName -> PeptideSequence, Charge -> PrecursorCharge, Area -> Intensity, Fragment -> FragmentIon, Sequence -> NakedSequence.

#### Usage

```
disaggregate(data, all.columns = FALSE)
```

#### **Arguments**

data A data frame containing SWATH data.

all.columns Option that all columns are processed. Otherwise only the typical columns

needed for downstream analysis are processed.

#### Value

Returns a data frame containing the SWATH data in a transition-level table.

#### Author(s)

Peter Blattmann

```
## Not run:
    data("OpenSWATH_data", package="SWATH2stats")
    data("Study_design", package="SWATH2stats")
    data <- sample_annotation(OpenSWATH_data, Study_design)
    data.filtered.decoy <- filter_mscore(data, 0.01)
    raw <- disaggregate(data.filtered.decoy)

## End(Not run)</pre>
```

filter\_all\_peptides 15

filter\_all\_peptides Select all proteins that are supported by peptides.

## **Description**

This functions counts all proteins that are supported by peptides (including non proteo-typic peptides). All peptides (incl. non proteotypic peptides are selected. For the proteins supproted by proteotypic peptide the "1/" in front of the identifier is removed to facilitate further data processing.

## Usage

```
filter_all_peptides(data, column = "proteinname", n = 6)
```

## **Arguments**

data A data frame containing SWATH data.

column Which column contains the data to modify?

n How many new IDs should we print to show if this worked?

## Value

Returns a data frame with the data from both proteotypic and non-proteotypic peptides.

## Author(s)

Peter Blattmann

```
## Not run:
    data("OpenSWATH_data", package="SWATH2stats")
    data("Study_design", package="SWATH2stats")
    data <- sample_annotation(OpenSWATH_data, Study_design)
    data.filtered.decoy <- filter_mscore(data, 0.01)
    data.all <- filter_all_peptides(data.filtered.decoy)
## End(Not run)</pre>
```

16 filter\_mscore

filter\_mscore

Filter openSWATH output table according to mscore.

#### **Description**

This function filters the SWATH data according to the m\_score value, as well as to the number of occurence in the data (requant) and within a condition (condition).

## Usage

```
filter_mscore(data, mscore = 1, rm.decoy = TRUE)
```

## **Arguments**

data A data frame containing SWATH data.

mscore Value that defines the mscore threshold according to which the data will be

filtered.

rm. decoy Drop decoys from the data?

## Value

A copy of the data which has been mscore-filtered.

#### Author(s)

Peter Blattmann

```
## Not run:
data("OpenSWATH_data", package="SWATH2stats")
data("Study_design", package="SWATH2stats")
data <- sample_annotation(OpenSWATH_data, Study_design)
data.filtered <- filter_mscore(data, 0.01)
data.filtered <- filter_mscore_freqobs(data, 0.01, 0.8)
data.filtered <- filter_mscore_condition(data, 0.01, 3)
## End(Not run)</pre>
```

filter\_mscore\_condition 17

```
filter_mscore_condition
```

Filter openSWATH output table according to mscore.

## Description

This function filters the SWATH data according to the m\_score value, as well as to the number of occurence in the data (requant) and within a condition (condition).

## Usage

```
filter_mscore_condition(data, mscore = 1, n.replica, rm.decoy = TRUE)
```

## **Arguments**

data	A data frame containing SWATH data.
mscore	Value that defines the mscore threshold according to which the data will be filtered.
n.replica	Number of measurements within at least one condition that have to pass the mscore threshold for this transition.
rm.decoy	Drop decoys from the data?

## Value

A copy of the data which has been mscore-filtered.

## Author(s)

Peter Blattmann

```
## Not run:
data("OpenSWATH_data", package="SWATH2stats")
data("Study_design", package="SWATH2stats")
data <- sample_annotation(OpenSWATH_data, Study_design)
data.filtered <- filter_mscore(data, 0.01)
data.filtered <- filter_mscore_freqobs(data, 0.01, 0.8)
data.filtered <- filter_mscore_condition(data, 0.01, 3)
## End(Not run)</pre>
```

18 filter\_mscore\_fdr

filter\_mscore\_fdr

Filter annotated OpenSWATH/pyProphet output table to achieve a high FDR quality data matrix with controlled overall protein FDR and quantitative values for all peptides mapping to these high-confidence proteins (up to a desired overall peptide level FDR quality).

#### **Description**

This function controls the protein FDR over a multi-run OpenSWATH/pyProphet output table and filters all quantitative values to a desired overall/global peptide FDR level. It first finds a suitable m-score cutoff to minimally achieve a desired global FDR quality on a protein master list based on the function mscore4protfdr. It then finds a suitable m-score cutoff to minimally achieve a desired global FDR quality on peptide level based on the function mscore4pepfdr. Finally, it reports all the peptide quantities derived based on the peptide level cutoff for only those peptides mapping to the protein master list. It further summarizes the protein and peptide numbers remaining after the filtering and evaluates the individual run FDR qualities of the peptides (and quantitation events) selected.

#### **Usage**

```
filter_mscore_fdr(data, FFT = 1, overall_protein_fdr_target = 0.02,
  mscore_limit = 0.01, upper_overall_peptide_fdr_limit = 0.05,
  rm.decoy = TRUE)
```

## **Arguments**

data Annotated OpenSWATH/pyProphet data table.

FFT Ratio of false positives to true negatives, q-values from [Injection\_name]\_full\_stat.csv

in pyProphet stats output. As an approximation, the q-values of multiple runs are averaged and supplied as argument FFT. Numeric from 0 to 1. Defaults to 1, the most conservative value (1 Decoy indicates 1 False target). For further

details see the Vignette Section 1.3 and 4.1.

overall\_protein\_fdr\_target

FDR target for the protein master list for which quantitative values down to the

less strict peptide\_fdr criterion will be kept/reported. Defaults to 0.02.

mscore\_limit FDR target for the quantitative values kept/reported for all peptides mapping

to the high-confidence protein master list. Defaults to 0.05. If all values up to

 $m_{score} 0.01$  shall be kept, set = 1.

rm. decoy Logical T/F, whether decoy entries should be removed after the analysis. De-

faults to TRUE. Can be useful to disable to track the influence on decoy fraction

by further filtering steps such as requiring 2 peptides per protein.

#### Value

Returns a data frame with the filtered data.

filter\_mscore\_freqobs 19

#### Author(s)

Moritz Heusel

## **Examples**

filter\_mscore\_freqobs Filter openSWATH output table according to mscore.

## **Description**

This function filters the SWATH data according to the m\_score value, as well as to the number of occurence in the data (requant) and within a condition (condition)

## Usage

```
filter_mscore_freqobs(data, mscore = 0.01, percentage = NULL,
    rm.decoy = TRUE, file_column = "filename")
```

## **Arguments**

data A data frame containing SWATH data.

mscore Value that defines the mscore threshold according to which the data will be

filtered.

percentage Percentage in which replicas the transition has to reach the mscore threshold. ??

rm. decoy Option to remove the decoys during filtering.

#### Value

Returns a data frame with the filtered data.

#### Author(s)

Peter Blattmann

## **Examples**

```
## Not run:
data("OpenSWATH_data", package="SWATH2stats")
data("Study_design", package="SWATH2stats")
data <- sample_annotation(OpenSWATH_data, Study_design)
data.filtered <- filter_mscore(data, 0.01)
data.filtered <- filter_mscore_freqobs(data, 0.01, 0.8)
data.filtered <- filter_mscore_condition(data, 0.01, 3)
## End(Not run)</pre>
```

```
filter_on_max_peptides
```

Filter only for the highest intense peptides

## Description

In order to reduce the data, the data is filtered only for the proteins with the highest intensity peptides.

## Usage

```
filter_on_max_peptides(data, n_peptides = 6, rm.decoy = TRUE,
  column = "proteinname")
```

#### Arguments

data A data frame containing SWATH data with the column names: ProteinNames,

PeptideSequence, PrecursorCharge, Intensity.

n\_peptides Maximum number of highest intense peptides to filter the data on.

rm. decoy Option to remove the decoys during filtering.

column which column to use for filtering?

#### Value

Returns a data frame of the filtered data.

#### Author(s)

Peter Blattmann

filter\_on\_min\_peptides

#### **Examples**

```
## Not run:
data("OpenSWATH_data", package="SWATH2stats")
data("Study_design", package="SWATH2stats")
data <- sample_annotation(OpenSWATH_data, Study_design)
data.filtered <- filter_mscore_freqobs(data, 0.01,0.8)
data.max <- filter_on_max_peptides(data.filtered, 5)
## End(Not run)</pre>
```

```
filter_on_min_peptides
```

Filter openSWATH output for proteins that are identified by a minimum of n independent peptides.

## Description

This function removes entries mapping to proteins that are identified by less than n\_peptides. Removing single-hit proteins from an analysis can significantly increase the sensitivity under strict protein fdr criteria, as evaluated by e.g. assess\_fdr\_overall.

#### Usage

```
filter_on_min_peptides(data, n_peptides = 6, rm.decoy = TRUE,
  column = "proteinname")
```

## Arguments

data	Data table that is produced by the openSWATH/iPortal workflow.
n_peptides	Number of minimal number of peptide IDs associated with a protein ID in order to be kept in the dataset.
rm decov	Ontion to remove the decays during filtering

rm. decoy Option to remove the decoys during filtering.

column which column to use for filtering?

#### Value

Returns the filtered data frame with only peptides that map to proteins with  $\geq n_p$  peptides peptides.

#### Author(s)

Moritz Heusel

#### **Examples**

```
## Not run:
data("OpenSWATH_data", package="SWATH2stats")
data("Study_design", package="SWATH2stats")
data <- sample_annotation(OpenSWATH_data, Study_design)
data.filtered <- filter_mscore_freqobs(data, 0.01,0.8)
data.max <- filter_on_max_peptides(data.filtered, 5)
data.min.max <- filter_on_min_peptides(data.max, 3)
## End(Not run)</pre>
```

filter\_proteotypic\_peptides

Filter for proteins that are supported by proteotypic peptides.

## Description

Peptides can match to several proteins. With this function proteotypic peptides, peptides that are only contained in one protein are selected. Additionally the number of proteins are counted and printed.

## Usage

```
filter_proteotypic_peptides(data, rm.decoy = TRUE, column = "proteinname")
```

#### **Arguments**

data A data frame containing SWATH data.

rm. decoy Option to remove the decoys during filtering.

column Which column to query for filtering?

#### Value

Returns a data frame with only the data supported by proteotypic peptides.

#### Author(s)

Peter Blattmann

```
## Not run:
data("OpenSWATH_data", package="SWATH2stats")
data("Study_design", package="SWATH2stats")
data <- sample_annotation(OpenSWATH_data, Study_design)
data.filtered.decoy <- filter_mscore(data, 0.01)
data.all <- filter_proteotypic_peptides(data.filtered.decoy)
## End(Not run)</pre>
```

import\_data 23

import_data	Transforms the column names from a data frame to the required format.

#### **Description**

This functions transforms the column names from a data frame from another format to a data frame with column names used by the OpenSWATH output and required for these functions. During executing of the function the corresponding columns for each column in the data need to be selected. For columns that do not correspond to a certain column 'not applicable' needs to be selected and the column names are not changed.

## Usage

import\_data(data)

## **Arguments**

data

A data frame containing the SWATH-MS data (one line per peptide precursor quantified) but with different column names.

#### Value

Returns the data frame in the appropriate format.

#### Note

List of column names of the OpenSWATH data: ProteinName: Unique identifier for protein or proteingroup that the peptide maps to. Proteotypic peptides should be indicated by 1/ in order to be recognized as such by the function filter proteotypic peptides. FullPeptideName: Unique identifier for the peptide. Charge: Charge of the peptide precursor ion quantified. Sequence: Naked peptide sequence without modifications, aggr Fragment Annotation; aggregated annotation for the different Fragments quantified for this peptide. In the OpenSWATH results the different annotation in OpenSWATH are concatenated by a semicolon. aggr\_Peak\_Area: aggregated Intensity values for the different Fragments quantified for this peptide. In the OpenSWATH results the aggregated Peak Area intensities are concatenated by a semicolon. transition\_group\_id: A unique identifier for each transition group used. decoy: Indicating with 1 or 0 if this transition group is a decoy. m\_score: Column containing the score that is used to estimate FDR or filter. M-score values of identified peak groups are equivalent to a q-value and thus typically are smaller than 0.01, depending on the confidence of identification (the lower the m-score, the higher the confidence). Column containing the score that is used to estimate FDR or filter. RT: Column containing the retention time of the quantified peak. filename: Column containing the filename or a unique identifier for each injection. Intensity: column containing the intensity value for each quantified peptide. Columns needed for FDR estimation and filtering functions: ProteinName, FullPeptideName, transition\_group\_id, decoy, m score Columns needed for conversion to transition-level format (needed for MSStats and mapDIA input): aggr\_Fragment\_Annotation, aggr\_Peak\_Are

24 load\_mart

## Author(s)

Peter Blattmann

#### **Examples**

```
## Not run:
  data('Spyogenes', package = 'SWATH2stats')
head(data)
  str(data)
## End(Not run)
```

load\_mart

Gather some data from biomaRt. Convert protein ids

## **Description**

This function renames protein ids in a data frame or file.

## Usage

```
load_mart(species, ensembl.path, mart, verbose = FALSE)
```

#### **Arguments**

species The species of the protein identifiers in the term used by biomaRt (e.g. "hsapi-

ens\_gene\_ensembl", "mmusculus\_gene\_ensembl", "drerio\_gene\_ensembl", etc.)

ensembl.path ensembl host to query.

mart The type of mart (e.g. "ENSEMBL\_MART\_ENSEMBL", etc.)

verbose print a summary of the ensembl connection.

## Value

ensembl connection for performing future queries.

## Author(s)

Peter Blattmann

mscore4assayfdr 25

mscore4assayfdr	Find m_score cutoff to reach a desired FDR on assay level (over the entire OpenSWATH/pyProphet output table)
	· ····································

## Description

This function estimates the m\_score cutoff required in a dataset to reach a given overall assay level FDR. It counts target and decoy assays at high resolution across the m\_score cutoffs and reports a useful m\_score cutoff - assay FDR pair close to the supplied fdr\_target level over the entire dataset. The m\_score cutoff is returned by the function and can be used in the context of the filtering functions, e.g.: data.assayFDR1pc<-filter\_mscore(data, mscore4assayfdr(data, fdr\_target=0.01)) To arrive from decoy counts at an estimation of the false discovery rate (false positives among the targets remaining at a given mscore cutoff) the ratio of false positives to true negatives (decoys) (FFT) must be supplied. It is estimated for each run individually by pyProphet and contained in the pyProphet statistics [Injection\_name]\_full\_stat.csv. As an approximation, the FFTs of multiple runs are averaged and supplied as argument FFT. For further details see the Vignette Section 1.3 and 4.1. For FDR evaluations on peptide and protein level, please refer to functions mscore4pepfdr and mscore4protfdr.

## Usage

```
mscore4assayfdr(data, FFT = 1, fdr_target = 0.01)
```

## **Arguments**

data	Annotated OpenSWATH/pyProphet data table. See function sample_annotation from this package.
FFT	Ratio of false positives to true negatives, q-values from [Injection_name]_full_stat.csv in pyProphet stats output. As an approximation, the q-values of multiple runs are averaged and supplied as argument FFT. Numeric from 0 to 1. Defaults to 1, the most conservative value (1 Decoy indicates 1 False target).
fdr_target	Assay FDR target, numeric, defaults to 0.01. An m_score cutoff achieving an FDR < fdr_target will be selected. Calculated as FDR = (TN*FFT/T); TN=decoys, T=targets, FFT=see above.

#### Value

Returns the m\_score cutoff selected to arrive at the desired FDR

#### Author(s)

Moritz Heusel

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#### **Examples**

```
## Not run:
    data("OpenSWATH_data", package="SWATH2stats")
    data("Study_design", package="SWATH2stats")
    data <- sample_annotation(OpenSWATH_data, Study_design)
    mscore4assayfdr(data, FFT=0.7, fdr_target=0.01)
## End(Not run)</pre>
```

mscore4pepfdr

Find m\_score cutoff to reach a desired FDR on peptide level (over the entire OpenSWATH/pyProphet output table)

#### **Description**

This function estimates the m\_score cutoff required in a dataset to reach a given overall peptide level FDR. It counts target and decoy peptides (unique FullPeptideName) at high resolution across the m\_score cutoffs and reports a useful m\_score cutoff - peptide FDR pair close to the supplied fdr\_target level over the entire dataset. The m\_score cutoff is returned by the function and can be used in the context of the filtering functions, e.g.: data.pepFDR2pc<-filter\_mscore(data, mscore4pepfdr(data, fdr\_target=0.02)) To arrive from decoy counts at an estimation of the false discovery rate (false positives among the targets remaining at a given mscore cutoff) the ratio of false positives to true negatives (decoys) (FFT) must be supplied. It is estimated for each run individually by pyProphet and contained in the pyProphet statistics [Injection\_name]\_full\_stat.csv. As an approximation, the FFTs of multiple runs are averaged and supplied as argument FFT. For further details see the Vignette Section 1.3 and 4.1. For FDR evaluations on assay and protein level, please refer to functions mscore4assayfdr and mscore4protfdr

#### **Usage**

```
mscore4pepfdr(data, FFT = 1, fdr_target = 0.01)
```

## **Arguments**

data Annotated OpenSWATH/pyProphet data table. See function sample\_annotation

from this package.

FFT Ratio of false positives to true negatives, q-values from [Injection\_name]\_full\_stat.csv

in pyProphet stats output. As an approximation, the q-values of multiple runs are averaged and supplied as argument FFT. Numeric from 0 to 1. Defaults to 1,

the most conservative value (1 Decoy indicates 1 False target).

fdr\_target FDR target, numeric, defaults to 0.01. An m\_score cutoff achieving an FDR

< fdr\_target will be selected. Calculated as FDR = (TN\*FFT/T); TN=decoys,

T=targets, FFT=see above.

#### Value

Returns the m\_score cutoff selected to arrive at the desired FDR

mscore4protfdr 27

#### Author(s)

Moritz Heusel

#### **Examples**

```
## Not run:
    data("OpenSWATH_data", package="SWATH2stats")
    data("Study_design", package="SWATH2stats")
    data <- sample_annotation(OpenSWATH_data, Study_design)
    mscore4pepfdr(data, FFT=0.7, fdr_target=0.01)
## End(Not run)</pre>
```

mscore4protfdr

Find m\_score cutoff to reach a desired FDR on protein level (over the entire OpenSWATH/pyProphet output table)

#### **Description**

This function estimates the m\_score cutoff required in a dataset to reach a given overall protein level FDR. This filter is to be used with caution as the resulting quantitative matrix is relatively sparse. It can be filled with quantitative values at a lower FDR quality level. It counts target and decoy peptides (unique ProteinName) at high resolution across the m\_score cutoffs and reports a useful m\_score cutoff - peptide FDR pair close to the supplied fdr\_target level over the entire dataset. The m\_score cutoff is returned by the function and can be used in the context of the filtering functions, e.g.: data.protFDR5pc<-filter\_mscore(data, mscore4protfdr(data, fdr\_target=0.02)) To arrive from decoy counts at an estimation of the false discovery rate (false positives among the targets remaining at a given mscore cutoff) the ratio of false positives to true negatives (decoys) (FFT) must be supplied. It is estimated for each run individually by pyProphet and contained in the pyProphet statistics [Injection\_name]\_full\_stat.csv. As an approximation, the FFTs of multiple runs are averaged and supplied as argument FFT. For further details see the Vignette Section 1.3 and 4.1. For FDR evaluations on assay and peptide level, please refer to functions mscore4assayfdr and mscore4pepfdr.

## Usage

```
mscore4protfdr(data, FFT = 1, fdr_target = 0.02)
```

#### **Arguments**

data

Annotated OpenSWATH/pyProphet data table. See function sample\_annotation

from this package.

FFT

Ratio of false positives to true negatives, q-values from [Injection\_name]\_full\_stat.csv in pyProphet stats output. As an approximation, the q-values of multiple runs are averaged and supplied as argument FFT. Numeric from 0 to 1. Defaults to 1, the most conservative value (1 Decoy indicates 1 False target).

fdr\_target FDR target, numeric, defaults to 0.01. An m\_score cutoff achieving an FDR

< fdr\_target will be selected. Calculated as FDR = (TN\*FFT/T); TN=decoys,

T=targets, FFT=see above.

#### Value

Returns the m\_score cutoff selected to arrive at the desired FDR quality.

## Author(s)

Moritz Heusel

#### **Examples**

```
## Not run:
    data("OpenSWATH_data", package="SWATH2stats")
    data("Study_design", package="SWATH2stats")
    data <- sample_annotation(OpenSWATH_data, Study_design)
    mscore4protfdr(data, FFT=0.7, fdr_target=0.01)
## End(Not run)</pre>
```

plot\_correlation\_between\_samples

Plots the correlation between injections.

## Description

This function plots the Pearson's and Spearman correlation between samples. If decoys are present these are removed before plotting.

#### **Usage**

```
plot_correlation_between_samples(data, column.values = "intensity",
    size = 6, comparison = transition_group_id ~ condition + bioreplicate,
    fun.aggregate = NULL, label = TRUE, ...)
```

## **Arguments**

Intensity or Signal, but also the retention time.

size How large should the text in the grid be (smaller is better for rmarkdown html

reports).

comparison The comparison for assessing the variability. Default is to assess the variability

per transition\_group\_id over the different Condition and Replicates. Compari-

son is performed using the dcast() function of the reshape2 package.

plot\_fdr\_cube 29

fun.aggregate	If for the comparison values have to be aggregated one needs to provide the function here.
label	Option to print correlation value in the plot.
	Further arguments passed to methods.

#### Value

Plots in Rconsole a correlation heatmap and returns the data frame used to do the plotting.

#### Author(s)

Peter Blattmann

#### **Examples**

## **Description**

This function creates standard plots from result arrays as produced by e.g. the function assess\_fdr\_byrun(), visualizig assay, peptide and protein level FDR for each run at m-score cutoffs 1e-2 and 1e-3. Furthermore, Target and Decoy ID numbers are visualized.

#### Usage

```
plot_fdr_cube(x, output = "Rconsole", filename = "FDR_report_byrun",
    plot_mscore_levels = c(0.01, 0.001), ...)
```

X	Array of by-run FDR assessment results as produced e.g. by the function assess_fdr_byrun() from this package.	
output	Choose output type. "pdf_csv" creates the output as files in the working directory, "Rconsole" triggers delivery of the output to the console enabling further computation and/or custom plotting / output.	
filename	Basename for output files to be created (if output = "pdf_csv" has been selected).	
plot_mscore_levels		
	Define m-score levels to plot the estimated FDR results.	
	Extra arguments passed on to functions inside this.	

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#### Value

Originally this returned nothing, but now it makes a list of ggplot2 plots which may be passed along and plotted as desired (with that in mind, I would like to remove the explicit plot() calls in this function).

#### Author(s)

Moritz Heusel

#### **Examples**

plot\_fdr\_table

Plot functionality for results of class "fdr\_table" as produced by e.g. the function assess\_fdr\_overall()

#### **Description**

This function created standard plots from results of class "fdr\_table" as produced by e.g. the function assess\_fdr\_overall() visualizig ID numbers in dependence of estimated FDR and also estimated FDR in dependence of m\_score cutoff.

## Usage

```
plot_fdr_table(x, output = "Rconsole", filename = "FDR_report_overall", ...)
```

х	List of class "fdr_table" as produced e.g. by the function assess_fdr_overall() from this package.
output	Choose output type. "pdf_csv" creates the output as files in the working directory, "Rconsole" triggers delivery of the output to the console enabling further computation or custom plotting / output.
filename	Basename for output files to be created (if output = "pdf_csv" has been selected).
	Extra arguments passed on to functions inside this.

plot\_variation 31

## Value

Originally this returned nothing, but now it makes a list of ggplot2 plots which may be passed along and plotted as desired (with that in mind, I would like to remove the explicit plot() calls in this function).

#### Author(s)

Moritz Heusel

## **Examples**

```
## Not run:
data("OpenSWATH_data", package="SWATH2stats")
data("Study_design", package="SWATH2stats")
data <- sample_annotation(OpenSWATH_data, Study_design)
x <- assess_fdr_overall(data, FFT=0.7, output = "Rconsole", plot = FALSE)
plot.fdr_table(x, output = "pdf_csv", filename = "Assess_fdr_overall_testplot")
## End(Not run)</pre>
```

plot\_variation

Plots the coefficient of variation for different replicates.

## **Description**

This function plots the coefficient of variation within replicates for a given value. If decoys are present these are removed before plotting.

#### Usage

```
plot_variation(data, column.values = "intensity",
  comparison = transition_group_id + condition ~ bioreplicate,
  fun.aggregate = NULL, label = TRUE, ...)
```

data	Data frame that is produced by the OpenSWATH/pyProphet workflow.
column.values	Indicates the columns for which the variation is assessed. This can be the Intensity or Signal, but also the retention time.
comparison	The comparison for assessing the variability. Default is to assess the variability per transition_group_id and Condition over the different Replicates. Comparison is performed using the dcast() function of the reshape2 package.
fun.aggregate	If for the comparison values have to be aggregated one needs to provide the function here.
label	Option to print value of median cv.
	further arguments passed to method.

## Value

Returns a list with the data and calculated cv and a table that summarizes the mean, median and mode cv per Condition (if Condition is contained in the comparison). In addition it plots in Reconsole a violin plot with the observed coefficient of variations.

## Author(s)

Peter Blattmann

#### **Examples**

```
## Not run:
    data("OpenSWATH_data", package="SWATH2stats")
    data("Study_design", package="SWATH2stats")
    data <- sample_annotation(OpenSWATH_data, Study_design)
    plot_variation(data)
## End(Not run)</pre>
```

```
plot_variation_vs_total
```

Plots the total variation versus variation within replicates

#### **Description**

This function plots the total variation and the variation within replicates for a given value. If decoys are present these are removed before plotting.

## Usage

```
plot_variation_vs_total(data, column.values = "intensity",
   comparison1 = transition_group_id ~ bioreplicate + condition,
   comparison2 = transition_group_id + condition ~ bioreplicate,
   fun.aggregate = NULL, label = TRUE, ...)
```

data	Data table that is produced by the OpenSWATH/pyProphet workflow.
column.values	Indicates the columns for which the variation is assessed. This can be the Intensity or Signal, but also the retention time.
comparison1	The comparison for assessing the total variability. Default is to assess the variability per transition_group_id over the combination of Replicates and different Conditions.
comparison2	The comparison for assessing the variability within the replicates. Default is to assess the variability per transition_group_id and Condition over the different Replicates.

fun.aggregate If depending on the comparison values have to be aggregated one needs to pro-

vide the function here. (I think this should be sum, yesno?)

label Option to print value of median cv.

#### Value

Plots in Rconsole a violin plot comparing the total variation with the variation within replicates. In addition it returns the data frame from which the plotting is done and a table with the calculated mean, median and mode of the cv for the total or replicate data.

#### Author(s)

Peter Blattmann

## **Examples**

```
## Not run:
  data("OpenSWATH_data", package="SWATH2stats")
  data("Study_design", package="SWATH2stats")
  data <- sample_annotation(OpenSWATH_data, Study_design)
  plot_variation_vs_total(data)
## End(Not run)</pre>
```

```
reduce_OpenSWATH_output
```

Reduce columns of OpenSWATH data

#### **Description**

This function selects the columns from the standard OpenSWATH output to column needed for MSstats, aLFQ and mapDIA.

## Usage

```
reduce_OpenSWATH_output(data, column.names = NULL,
  data_file_column = "filename")
```

#### **Arguments**

data A data frame containing SWATH data.

column.names A vector of column names that can be selected.

#### Value

Returns a data frame with the selected columns.

#### Note

A basic set of columns are defined in the function and are used if no column names are indicated.

The column.names can be omitted and then the following columns are selected that are needed for MSstats and mapDIA analysis: ProteinName, FullPeptideName, Sequence, Charge, aggr\_Fragment\_Annotation, aggr\_Peak\_Area, filename, m\_score, decoy, Intensity, RT. This function should be ommitted if the data is analyzed afterwards with the aLFQ or imsbInfer package that needs further columns.

#### Author(s)

Peter Blattmann

## **Examples**

```
## Not run:
  data("OpenSWATH_data", package="SWATH2stats")
  data("Study_design", package="SWATH2stats")
  data <- sample_annotation(OpenSWATH_data, Study_design)
  data.filtered <- reduce_OpenSWATH_output(data)
## End(Not run)</pre>
```

removeDecoyProteins

Removes decoy proteins from the protein group label

## Description

There exist peptides annotated as protein groups with 2/ProteinA/DECOY\_ProteinB. However these are in principal proteotypic peptides and should be annoated 1/ProteinA. This function changes these labels accordingly. The subfunction rmDecoyProt removes the Decoy protein, calling removeDecoyProteins also changes the nubmer before the protein group accordingly.

#### Usage

```
removeDecoyProteins(data, column = "proteinname")
```

#### **Arguments**

data A data frame containing SWATH data.

column which column to query?

#### Value

Returns a data frame with changed protein labels

## Author(s)

Moritz Heusel

rmDecoyProt 35

#### **Examples**

```
## Not run:
data("OpenSWATH_data", package="SWATH2stats")
data("Study_design", package="SWATH2stats")
data <- sample_annotation(OpenSWATH_data, Study_design)
data.filtered.decoy <- filter_mscore(data, 0.01)
data.2 <- removeDecoyProteins(data.filtered.decoy)
## End(Not run)</pre>
```

rmDecoyProt

What string defines decoys?

## Description

What string defines decoys?

#### Usage

```
rmDecoyProt(x, pattern = "DECOY")
```

## Arguments

x proteinname string to query.
pattern chosen string to seek out.

sample\_annotation

Annotate the SWATH data with the sample information

#### **Description**

For statistical analysis and filtering the measurements need to be annotated with Filename, Condition, BioReplicate, and Run. This functions takes this information from a txt file containing this meta-data.

#### Usage

```
sample_annotation(data, sample_annotation, data_type = "OpenSWATH",
   annotation_file_column = "filename", data_file_column = "filename",
   condition_column = "condition", replicate_column = "bioreplicate",
   fullpeptidename_column = c("fullpeptidename", "fullunimodpeptidename"),
   run_id = NULL, run_column = "run", change_run_id = TRUE,
   verbose = FALSE)
```

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## Arguments

data A data frame containing SWATH data.

sample\_annotation

A data frame containing the columns: Filename, Condition, BioReplicate, Run. The values contained in the column filename have to be present in the filename of the SWATH data.

data\_type

Option to specify the format of the table, if the column names from an OpenSWATH output or MSstats table are used.

annotation\_file\_column

Name of the column containing the output file from the experiment metadata for each sample. In my own sample sheet, I keep columns for the mzXML files, tric outputs, raw files, and osw outputs from OpenSwathWorkFlow; and I cannot be relied upon to remember which is which, ergo this option.

data\_file\_column

Option to specify the column name where the injection file is specified. Default is set to "filename".

condition\_column

Which column annotates the experimental condition in the swath data?

replicate\_column

Which column annotates the replicate in the data?

run\_column Which column annotates the separate runs?

change\_run\_id Option to choose if the run\\_id column shall be reassigned to a unique value

combining the values of Condition, BioReplicate and Run. (Option only possi-

ble if data is of format "OpenSWATH")

verbose Option to turn on reporting on which filename it is working on.

#### **Details**

Given dataframes of TRIC processed data and sample annotations, mash them together into something appropriate for downstream analyses.

This performs some quick sanity checks on the data and annotations and creates the 'Condition', 'BioReplicate', and 'Run' columns along with other columns expected by MSstats/OpenSWATH.

## Value

Returns a dataframe with each row annotated for the study design

#### Author(s)

Peter Blattman

```
data("OpenSWATH_data", package="SWATH2stats")
data("Study_design", package="SWATH2stats")
data <- SWATH2stats::sample_annotation(OpenSWATH_data, Study_design, verbose=TRUE)
summary(data)</pre>
```

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SWATH2stats

SWATH2stats: a suite of tools to filter, plot, and convert DIA data.

#### **Description**

SWATH2stats: a suite of tools to filter, plot, and convert DIA data.

transform\_MSstats\_OpenSWATH

Transforms column names to OpenSWATH column names

## **Description**

This functions transforms the column names from a data frame in MSstats format to a data frame with column names used by the OpenSWATH output. The original table needs to contain at least the 10 columns defined by MSstats: ProteinName, PeptideSequence, PrecursorCharge, FragmentIon, ProductCharge, IsotopeLabelType, Condition, BioReplicate, Run, Intensity.)

#### Usage

```
transform_MSstats_OpenSWATH(data)
```

## **Arguments**

data

A data frame containing the SWATH data in the MSstats format

#### Value

The data frame in the appropriate format.

## Author(s)

Peter Blattmann

## References

Choi M, Chang CY, Clough T, Broudy D, Killeen T, MacLean B, Vitek O. MSstats: an R package for statistical analysis of quantitative mass spectrometry-based proteomic experiments. Bioinformatics. 2014 Sep 1;30(17):2524-6. doi: 10.1093/bioinformatics/btu305.

```
## Not run:
  data("MSstats_data", package="SWATH2stats")
  transform_MSstats_OpenSWATH(MSstats_data)
## End(Not run)
```

unifyProteinGroupLabels

Unify the protein group labels.

## **Description**

Unify the protein group labels (2/ProteinA/ProteinB and 2/ProteinB/ProteinA) to one common label (e.g. 2/ProteinA/ProteinB)

## Usage

```
unifyProteinGroupLabels(data, column = "proteinname", keep_colnames = FALSE)
```

## **Arguments**

data A data frame containing SWATH data.

column Which column to use for unifying the groups.

keep\_colnames Return the column names to their case-sensitive precursors.

## Value

Returns a data frame with the unififed protein labels.

## Author(s)

Moritz Heusel

```
## Not run:
data("OpenSWATH_data", package="SWATH2stats")
data("Study_design", package="SWATH2stats")
data <- sample_annotation(OpenSWATH_data, Study_design)
data.filtered.decoy <- filter_mscore(data, 0.01)
data.unified <- unifyProteinGroupLabels(data.filtered.decoy)
## End(Not run)</pre>
```

write\_matrix\_peptides 39

write\_matrix\_peptides Writes out an overview matrix of peptides mapping to a FDR quality controlled protein master list at controlled global peptide FDR quality.

#### **Description**

Writes out an overview matrix on peptide level of a supplied (unfiltered or prefiltered) OpenSWATH results data frame. The peptide quantification is achieved by summing the areas under all 6 transitions per precursor and summing all precursors per FullPeptideName. In order to keep the peptide-to-protein association, the FullPeptideName is joined with the ProteinName.

#### Usage

```
write_matrix_peptides(data, write.csv = FALSE, fun.aggregate = sum,
  filename = "SWATH2stats_overview_matrix_peptidelevel.csv",
  rm.decoy = FALSE)
```

## **Arguments**

data A data frame containing annotated OpenSWATH/pyProphet data.

write.csv Option to determine if table should be written automatically into csv file.

fun. aggregate What function to use when aggregating the set of intensities?

filename File base name of the .csv matrix written out to the working folder.

rm.decoy Logical whether decoys will be removed from the data matrix. Defaults to

FALSE. It's sometimes useful to know how decoys behave across a dataset and how many you allow into your final table with the current filtering strategy.

## Value

the peptides as a matrix! also output .csv matrix is written to the working folder.

#### Author(s)

Moritz Heusel

```
## Not run:
    data("OpenSWATH_data", package="SWATH2stats")
    data("Study_design", package="SWATH2stats")
    data <- sample_annotation(OpenSWATH_data, Study_design)
    write_matrix_peptides(data)
## End(Not run)</pre>
```

40 write\_matrix\_proteins

write\_matrix\_proteins Writes out an overview matrix of summed signals per protein identifier (lines) over run\_id(columns).

## **Description**

Writes out an overview matrix on protein level of a supplied (unfiltered or filtered) OpenSWATH results data frame. The protein quantification is achieved by summing the areas under all 6 transitions per precursor, summing all precursors per FullPeptideName and all FullPeptideName signals per ProteinName entry. This function does not select consistently quantified or top peptides but sums all signals availabe that may or may not originate from the same set of peptides across different runs. A more detailed overview can be generated using the function write\_matrix\_peptides(). Peptide selection can be achieved upstream using e.g. the functions filter\_mscore\_requant(), filter\_on\_max\_peptides() and filter\_on\_min\_peptides().

## Usage

```
write_matrix_proteins(data, write.csv = FALSE, fun.aggregate = sum,
  filename = "SWATH2stats_overview_matrix_proteinlevel.csv",
  rm.decoy = FALSE)
```

#### **Arguments**

data A data frame containing annotated OpenSWATH/pyProphet data.

write.csv Option to determine if table should be written automatically into csv file.

fun.aggregate What function to use when aggregating the set of intensities?

File base name of the .csv matrix written out to the working folder

rm. decoy Logical whether decoys will be removed from the data matrix. Defaults to

FALSE. It's sometimes useful to know how decoys behave across a dataset and how many you allow into your final table with the current filtering strategy.

#### Value

the peptides as a matrix, also output .csv matrix is written to the working folder

## Author(s)

Moritz Heusel

```
## Not run:
    data("OpenSWATH_data", package="SWATH2stats")
    data("Study_design", package="SWATH2stats")
    data <- sample_annotation(OpenSWATH_data, Study_design)
    write_matrix_proteins(data)
## End(Not run)</pre>
```

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