

# Package ‘SafeQuant’

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**Type** Package

**Title** SafeQuant: Proteomics Data Analysis

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**Description** The SafeQuant Package includes methods for analysis of quantitative Proteomics data. More documentation to come.

**Imports** affy,  
limma,  
gplots,  
seqinr,  
corrplot,  
optparse,  
data.table,  
epiR

**License** GPL-3

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---

addIdQvalues	<i>Add identification leve q-values to ExpressionSet (calculated based on target-decoy score distribution)</i>
--------------	--

---

## Description

Add identification leve q-values to ExpressionSet (calculated based on target-decoy score distribution)

## Usage

```
addIdQvalues(eset = eset)
```

## Arguments

eset	ExpressionSet
------	---------------

## Details

if ptm column is part if the ExpressionSet q-values are calculated seperately for modified and non-modified features

## Value

ExpressionSet object

## Note

No note

## See Also

[getIdLevelQvals](#)

## Examples

```
print("No examples")
```

---

```
addScaffoldPTMFAnnotations
```

*Add scaffold ptm annotaitons to tmt experiment*

---

### Description

Add scaffold ptm annotaitons to tmt experiment

### Usage

```
addScaffoldPTMFAnnotations(eset, file)
```

### Arguments

eset	ExpressionSet
file	path to Scaffold file

### Value

ExpressionSet object

### Note

No note

### References

No references

### Examples

```
print("No examples")
```

---

```
barplotMSSignal
```

*Barplot of ms-signal per column*

---

### Description

Barplot of ms-signal per column

### Usage

```
barplotMSSignal(eset,
  col = as.character(.getConditionColors(eset)[pData(eset)$condition, ]),
  method = c("sum", "sharedSignal"), cex.lab = 1.25, cex.axis = 1.25,
  labels = rownames(pData(eset)), ...)
```

**Arguments**

method	c("median","sum","sharedSignal")
matrix	matrix of ms-signals
color	color

**Details**

No details

**Note**

No note

**References**

NA

**Examples**

```
print("No examples")
```

---

createCalibrationCurve

*S3 class object describing a calibration curve and storing some figures of merit*

---

**Description**

S3 class object describing a calibration curve and storing some figures of merit

**Usage**

```
createCalibrationCurve(eset, method = "blank")
```

**Arguments**

eset	ExpressionSet
method	to calculate Limit of Detection / Limit of Quantification. c("blank","low")

**Details**

No details

**Value**

calibrationCurve object

**Note**

No note

## References

Statistical characterization of multiple-reaction monitoring mass spectrometry (MRM-MS) assays for quantitative proteomics, Mani et al. (2012), <http://www.ncbi.nlm.nih.gov/pubmed/23176545>

## Examples

```
print("No examples")
```

---

createExpDesign	<i>Create Experimental Design</i>
-----------------	-----------------------------------

---

## Description

Create Experimental Design

## Usage

```
createExpDesign(tag, nbPlex)
```

## Arguments

tag	user input tag e.g. 1,2,3:4,5,6 indicating two condition with 3 reps each
nbPlex	tmt 6 or 10 plex

## Details

The first listed condition is always the control condition

## Value

expDesign data.frame

## Note

No note

## References

NA

## Examples

```
print("No examples")
```

---

`createExpressionDataset`*Create ExpressionSet object*

---

**Description**

Create ExpressionSet object

**Usage**

```
createExpressionDataset(expressionMatrix = expressionMatrix,  
  expDesign = expDesign, featureAnnotations = featureAnnotations)
```

**Arguments**

expressionMatrix	matrix of expression signals per feature and sample
expDesign	experimental design data.frame
featureAnnotations	data.frame including e.g: Protein Description, Id score etc.

**Details**

No details

**Value**

ExpressionSet object

**Note**

No note

**References**

NA

**See Also**

[ExpressionSet](#)

**Examples**

```
print("No examples")
```

---

cvBoxplot	<i>C.V. boxplot</i>
-----------	---------------------

---

**Description**

C.V. boxplot

**Usage**

```
cvBoxplot(eset,  
  col = as.character(.getConditionColors(eset)[unique(pData(eset)$condition),  
    ]), cex.names = 0.9, cex.axis = 1.25, cex.lab = 1.25,  
  ylab = "C.V. (%)", ...)
```

**Arguments**

eset                      ExpressionSet

**Details**

No details

**Note**

No note

**References**

NA

**Examples**

```
print("No examples")
```

---

expDesignTagToExpDesign	<i>Create experimental design data.frame from user input string</i>
-------------------------	---

---

**Description**

Create experimental design data.frame from user input string

**Usage**

```
expDesignTagToExpDesign(tag, expDesignDefault)
```

**Arguments**

string	tag
data.frame	default expDesign



**Details**

tag: 1,2:3:4,5,6 condition isControl 1 Condition 1 TRUE 2 Condition 1 TRUE 3 Condition 1 TRUE  
4 Condition 2 FALSE 5 Condition 2 FALSE 6 Condition 2 FALSE

**Value**

data.frame describing experimental design

**Note**

No note

**References**

NA

**Examples**

```
print("No examples")
```

---

export.safeQuantAnalysis

*Export content of safeQuantAnalysis object*

---

**Description**

Export content of safeQuantAnalysis object

**Usage**

```
## S3 method for class 'safeQuantAnalysis'  
export(sqa, nbRows = nrow(sqa$pValue),  
       file = NA)
```

**Arguments**

sqa	safeQuantAnalysis object
nbRows	Number of rows to export. Features are ordred by increasing minimal p.value

**Details**

NA

**Note**

No note

**References**

NA

**See Also**[safeQuantAnalysis](#)**Examples**

```
print("No examples")
```

---

`getAAProteinCoordinates`*Get amino acid coordinates on protein*

---

**Description**

Get amino acid coordinates on protein

**Usage**

```
getAAProteinCoordinates(peptideSeq, proteinSeq, aaRegExpr = "[STY]")
```

**Arguments**

peptideSeq	peptide sequence
proteinSeq	protein sequence

**Details**

NA

**Value**

vector of protein coordinates (mmodification residue number)

**Note**

No note

**References**

NA

**Examples**

```
print("No examples")
```

---

getAllCV	<i>Calculate Coefficient of Variance per feature (Relative standard Deviation) per Condition</i>
----------	--

---

**Description**

Calculate Coefficient of Variance per feature (Relative standard Deviation) per Condition

**Usage**

```
getAllCV(eset)
```

**Arguments**

eset	ExpressionSet
------	---------------

**Details**

$CV = sd / mean$

**Value**

data.frame of CVs per condition

**Note**

No note

**References**

NA

**See Also**

[getCV](#)

**Examples**

```
print("No examples")
```

---

getAllEBayes	<i>Perform statistical test (mderated t-test), comparing all case to control</i>
--------------	--

---

**Description**

Perform statistical test (mderated t-test), comparing all case to control

**Usage**

```
getAllEBayes(eset = eset, adjust = F, log = T, method = "pairwise")
```

**Arguments**

eset	ExpressionSet
adjust	TRUE/FALSE adjust for multiple testing using Benjamini & Hochberg (1995)
method	log T/F log-transform expression values
	c("pairwise","all")

**Details**

No details

**Value**

ExpressionSet object

**Note**

No note

**References**

Empirical Bayes method, Smyth (2004), <http://www.ncbi.nlm.nih.gov/pubmed/16646809>

**See Also**

[eBayes](#)

**Examples**

```
print("No examples")
```

---

getBaselineIntensity	<i>Get signal at zscore x (x standard deviations below mean)</i>
----------------------	--

---

**Description**

Get signal at zscore x (x standard deviations below mean)

**Usage**

```
getBaselineIntensity(intensities, promille = 5)
```

**Arguments**

intensities	refrence run signals
percentile	baseline value set as specified promille

**Value**

baseline value

**Note**

No note

**References**

NA

**Examples**

```
print("No examples")
```

---

getCV	<i>Calculate Coefficient of Variance per feature (Relative standard Deviation)</i>
-------	--

---

**Description**

Calculate Coefficient of Variance per feature (Relative standard Deviation)

**Usage**

```
getCV(data)
```

**Arguments**

data	data.frame of replicate signals
------	---------------------------------

**Details**

$CV = sd / mean$

**Value**

vector of CVs

**Note**

No note

**References**

NA

**Examples**

```
print("No examples")
```

---

```
getExpDesignProgenesisCsv
```

*Parse Experimental Design from Progenesis Csv Export*

---

**Description**

Parse Experimental Design from Progenesis Csv Export

**Usage**

```
getExpDesignProgenesisCsv(file,  
  expressionColIndices = .getProgenesisCsvExpressionColIndices(file))
```

**Arguments**

file                      path to progenesis csv file

**Details**

No details

**Value**

data.frame describing experimental design

**Note**

No note

**References**

NA

**Examples**

```
print("No examples")
```

---

getGlobalNormFactors	<i>Get normalization factors. calculated as summed/median signal per run (column) over summed/median of first run.</i>
----------------------	--

---

**Description**

Get normalization factors. calculated as summed/median signal per run (column) over summed/median of first run.

**Usage**

```
getGlobalNormFactors(eset, method = "sum")
```

**Arguments**

eset	ExpressionSet
------	---------------

**Details**

No details

**Value**

vector of normalization factors

**Note**

No note

**References**

NA

**Examples**

```
print("No examples")
```

---

getIBAQeset	<i>Calculate intensity-based absolute-protein-quantification (iBAQ) metric per protein</i>
-------------	--

---

**Description**

Calculate intensity-based absolute-protein-quantification (iBAQ) metric per protein

**Usage**

```
getIBAQeset(eset, proteinDB = NA, peptideLength = c(5, 36),  
  nbMiscleavages = 0, proteaseRegExp = .getProteaseRegExp("trypsin"))
```

**Arguments**

eset	protein level ExpressionSet
peptideLength	peptide length interval (to get number of peptides used for normalization)
nbMiscleavages	number of mis-cleavages allowed when digesting protein sequences in silico (to get number of peptides used for normalization)
proteaseRegExp	protease Reg Exp cleavage rule
list	protein sequences

**Details**

No details

**Value**

ExpressionSet

**Note**

No note

**References**

Global quantification of mammalian gene expression control, Schwanhauser (2011), <http://www.ncbi.nlm.nih.gov/pubmed/21593866>, Critical assessment of proteome-wide label-free absolute abundance estimation strategies. Ahrne (2013), <http://www.ncbi.nlm.nih.gov/pubmed/23794183>

**Examples**

```
print("No examples")
```

---

getIdLevelQvals	<i>Calculates identification level q-values based on target-decoy score distributions</i>
-----------------	---

---

**Description**

Calculates identification level q-values based on target-decoy score distributions

**Usage**

```
getIdLevelQvals(scores, isDecoy)
```

**Arguments**

scores	peptide/protein identificationscore
isDecoy	vector of TRUE/FALSE



**Details**

$q\text{-value} = (\text{Nb. Decoy Entries at idScore Threshold } S^*) / (\text{Nb. Target Entries at idScore Threshold } S)$ . (\* idScore  $\geq S$ )

**Value**

vector of q.values

**Note**

No note

**References**

NA

**Examples**

```
print("No examples")
```

---

getImpuritiesMatrix	<i>Get Thermo TMT impurity matrix</i>
---------------------	---------------------------------------

---

**Description**

Get Thermo TMT impurity matrix

**Usage**

```
getImpuritiesMatrix(plexNb = 6, test = F)
```

**Arguments**

plexNb                      integer, 6 or 10 plex

**Details**

No details

**Value**

impurity matrix matrix

**Note**

No note

**References**

NA

**Examples**

```
print("No examples")
```

---

getIntSumPerProtein	<i>Sum up raw intensities per protein and channel. keep track of number of summed spectra and unique peptides</i>
---------------------	---

---

### Description

Sum up raw intensities per protein and channel. keep track of number of summed spectra and unique peptides

### Usage

```
getIntSumPerProtein(intData, proteinACs, peptides, minNbPeptPerProt = 1)
```

### Arguments

intData	data.frame of intensities per channel
proteinACs	vector of protein accession numbers
peptides	vector of peptide sequneces
minNbPeptPerProt	minimal number of peptides per protein

### Details

NA

### Value

list containing 3 objects 1) data.frame of channel intensities per protein ac, 2) vector listing number of summed spectra per protein, 3) vector listing number of summed peptides per protein

### Note

No note

### References

NA

### Examples

```
print("No examples")
```

---

getLoocvFoldError	<i>Leave-One-Out Cross Validate Qunatification Model</i>
-------------------	--

---

**Description**

Leave-One-Out Cross Validate Qunatification Model

**Usage**

```
getLoocvFoldError(df)
```

**Arguments**

`data.frame` of two columns 1) "signal" - ms metric 2) "cpc" absolute quantity

**Details**

No details

**Value**

data.frame of fold errors per (left-out) protein

**Note**

No note

**References**

NA

**See Also**

NA

**Examples**

```
print("No examples")
```

---

getMeanCenteredRange    *Get modification coordinates on protein*

---

**Description**

Get modification coordinates on protein

**Usage**

```
getMeanCenteredRange(d, nbSd = 4)
```

**Arguments**

d	numeric vector
nbSd	range spanning number of sd frmo mean

**Details**

NA

**Value**

vector range boundries

**Note**

No note

**References**

NA

**Examples**

```
print("No examples")
```

---

getModifProteinCoordinates  
                          *Get modification coordinates on protein*

---

**Description**

Get modification coordinates on protein

**Usage**

```
getModifProteinCoordinates(modifAnnot, peptideSeq, proteinSeq, format = 1)
```

**Arguments**

modifAnnot	modification as annotated by progenesis. E.g. '[15] Phospho (ST) [30] Phospho (ST)'
peptideSeq	peptide sequence
proteinSeq	protein sequence
numeric	format 1) progenesis 2) scaffold

**Details**

NA

**Value**

vector of protein coordinates (mmodification residue number)

**Note**

No note

**References**

NA

**Examples**

```
print("No examples")
```

---

getMotifX

*Create motif-x peptide annotation*

---

**Description**

Create motif-x peptide annotation

**Usage**

```
getMotifX(modifPos, peptide, proteinSeq, motifLength = 4)
```

**Arguments**

modifPos	vector positions
peptide	peptide sequence
proteinSeq	protein sequence
motifLength	motif flanking sequence

**Details**

motif-x example PGDYS\*TTPG

**Value**

vector of motifs

**Note**

No note

**References**

NA

**Examples**

```
print("No examples")
```

---

```
getNbDetectablePeptides
```

*Get number peptides passing defined length criteria*

---

**Description**

Get number peptides passing defined length criteria

**Usage**

```
getNbDetectablePeptides(peptides, peptideLength = c(5, 36))
```

**Arguments**

peptides	list of peptides
vector	of two integers defining peptide length range

**Details**

No details

**Value**

integer corresponding to number of detectable peptides

**Note**

No note

**Examples**

```
print("No examples")
```

---

getNbMisCleavages	<i>Get number of mis-cleavages perp peptide</i>
-------------------	---

---

**Description**

Get number of mis-cleavages perp peptide

**Usage**

```
getNbMisCleavages(peptide, protease = "trypsin")
```

**Arguments**

peptide	character vector
protease	regular expression

**Details**

NA

**Value**

vector of integers

**Note**

No note

**References**

NA

**Examples**

```
print("No examples")
```

---

getNbPeptidesPerProtein	<i>Get number of peptides per protein</i>
-------------------------	---

---

**Description**

Get number of peptides per protein

**Usage**

```
getNbPeptidesPerProtein(eset)
```

**Arguments**

eset	ExpressionSet
------	---------------

**Details**

NA

**Value**

table

**Note**

No note

**References**

NA

**Examples**

```
print("No examples")
```

---

```
getNbSpectraPerProtein
```

*Get number of spectra per protein*

---

**Description**

Get number of spectra per protein

**Usage**

```
getNbSpectraPerProtein(eset)
```

**Arguments**

eset	ExpressionSet
------	---------------

**Details**

NA

**Value**

table

**Note**

No note

**References**

NA

**Examples**

```
print("No examples")
```



---

`getPeptides`*Digest protein*

---

**Description**

Digest protein

**Usage**

```
getPeptides(proteinSeq, proteaseRegExp = .getProteaseRegExp("trypsin"),  
            nbMiscleavages = 0)
```

**Arguments**

<code>proteinSeq</code>	protein sequence
-------------------------	------------------

**Details**

No details

**Value**

vector of peptides

**Note**

No note

**Examples**

```
print("No examples")
```

---

`getRatios`*Calculate ratios, comparing all case to control*

---

**Description**

Calculate ratios, comparing all case to control

**Usage**

```
getRatios(eset, method = "median", log2 = T)
```

**Arguments**

<code>eset</code>	ExpressionSet
<code>method</code>	median or mean

**Details**

No details

**Value**

ExpressionSet object

**Note**

No note

**References**

NA

**Examples**

```
print("No examples")
```

---

getRTNormFactors	<i>Get retentiontime base normalization factors</i>
------------------	---

---

**Description**

Get retentiontime base normalization factors

**Usage**

```
getRTNormFactors(eset, minFeaturesPerBin = 100)
```

**Arguments**

eset	ExpressionSet
minFeaturesPerBin	minumum number of features per bin. If nb. features are < minFeaturesPerBin -> include neighbouring bins.

**Details**

No details

**Value**

data.frame normalization factors per retention time bin (minute)

**Note**

No note

**References**

In Silico Instrumental Response Correction Improves Precision of Label-free Proteomics and Accuracy of Proteomics-based Predictive Models, Lyutvinskiy et al. (2013), <http://www.ncbi.nlm.nih.gov/pubmed/23589346>

**Examples**

```
print("No examples")
```

---

getScoreCutOff	<i>Get score cutoff for a given fdr cut-off</i>
----------------	---

---

**Description**

Get score cutoff for a given fdr cut-off

**Usage**

```
getScoreCutOff(scores, isDecoy, fdrCutOff = 0.01)
```

**Arguments**

scores	peptide/protein identificationscore
isDecoy	vector of TRUE/FALSE
fdrCutOff	[0,1]

**Details**

NA

**Value**

scoreCutoff

**Note**

No note

**References**

NA

**Examples**

```
print("No examples")
```

---

getSignalPerCondition	<i>Summarize replicate signal per condition (min)</i>
-----------------------	---

---

**Description**

Summarize replicate signal per condition (min)

**Usage**

```
getSignalPerCondition(eset, method = "median")
```

**Arguments**

method	median (default), mean, max, min, sd
data	data.frame of replicate signals

**Details**

No details

**Value**

data.frame of per condition signals

**Note**

No note

**References**

NA

**Examples**

```
print("No examples")
```

---

getTopX

*Calculate Mean of X most intense features*

---

**Description**

Calculate Mean of X most intense features

**Usage**

```
getTopX(entryData, topX = 3)
```

**Arguments**

entryData	data.frame listing feature intensities of one entry. Typically rows corresponds to Peptide entries of one protein
topX	best X flyers

**Details**

No details

**Value**

vector of topX intensities per column (sample)

**Note**

No note

## References

Absolute quantification of proteins by LCMSE: A virtue of parallel MS acquisition, Silva (2006), <http://www.ncbi.nlm.nih.gov/pubmed/16219938>, Critical assessment of proteome-wide label-free absolute abundance estimation strategies. Ahrne (2013), <http://www.ncbi.nlm.nih.gov/pubmed/23794183>

## Examples

```
print("No examples")
```

---

getUserOptions	<i>Read User Specified Command Line Options</i>
----------------	---

---

## Description

Read User Specified Command Line Options

## Usage

```
getUserOptions(version = version)
```

## Arguments

version	Safequant version number
---------	--------------------------

## Details

No details

## Value

user options list

## Note

No note

## References

NA

## Examples

```
print("No examples")
```

globalNormalize	<i>Normalize, Norm factors calculated as median signal per run (column) over median of first run.</i>
-----------------	---

---

**Description**

Normalize, Norm factors calculated as median signal per run (column) over median of first run.

**Usage**

```
globalNormalize(eset, globalNormFactors)
```

**Arguments**

eset	ExpressionSet
------	---------------

**Details**

No details

**Value**

eset ExpressionSet

**Note**

No note

**References**

NA

**See Also**

getGlobalNormFactors

**Examples**

```
print("No examples")
```

---

hClustHeatMap	<i>Hierarchical clustering heat map, cluster by runs intensity, features by ratio and display log2 ratios to control median</i>
---------------	---

---

**Description**

Hierarchical clustering heat map, cluster by runs intensity, features by ratio and display log2 ratios to control median

**Usage**

```
hClustHeatMap(eset, conditionColors = .getConditionColors(eset),  
              breaks = seq(-2, 2, length = 20), ...)
```

**Arguments**

eset	ExpressionSet
conditionColors	data.frame of colors per condition

**Details**

No details

**Note**

No note

**References**

NA

**Examples**

```
print("No examples")
```

---

isCon	<i>Check if protein is a contaminant entry</i>
-------	--

---

**Description**

Check if protein is a contaminant entry

**Usage**

```
isCon(ac)
```

**Arguments**

ac	vector of protein accession numbers
----	-------------------------------------

**Details**

contaminant proteins are typically annotated as: CON\_P0000

**Value**

vector TRUE/FALSE

**Note**

No note

**References**

NA

**Examples**

```
print("No examples")
```

---

isDecoy

*Check if protein is a decoy entry*

---

**Description**

Check if protein is a decoy entry

**Usage**

```
isDecoy(ac)
```

**Arguments**

ac                      vector of protein accession numbers

**Details**

decoy proteins are typically annotated as: REV\_P0000

**Value**

vector TRUE/FALSE

**Note**

No note

**References**

NA

**Examples**

```
print("No examples")
```



---

isStrippedACs	<i>Check if ACs are in "non-stripped" uniprot format e.g. "sp Q8CHJ2 AQP12_MOUSE"</i>
---------------	---

---

**Description**

Check if ACs are in "non-stripped" uniprot format e.g. "sp|Q8CHJ2|AQP12\_MOUSE"

**Usage**

```
isStrippedACs(acs)
```

**Arguments**

acs	accession numbers
-----	-------------------

**Details**

TRUE if less than 10

**Value**

boolean TRUE/FALSE

**Note**

No note

**References**

NA

**Examples**

```
print("No examples")
```

---

missinValueBarplot	<i>Plot Percentage of Features with with missing values</i>
--------------------	---

---

**Description**

Plot Percentage of Features with with missing values

**Usage**

```
missinValueBarplot(eset,  
  col = as.character(.getConditionColors(eset)[pData(eset)$condition, ]),  
  cex.axis = 1.25, cex.lab = 1.25, ...)
```

**Arguments**

      eset                  ExpressionSet

**Details**

      No details

**Note**

      No note

**References**

      NA

**Examples**

      print("No examples")

---

pairsAnnot	<i>Plot lower triangle Pearson's R<sup>2</sup>. Diagonal text, upper triangle all against all scatter plots with lm abline</i>
------------	--

---

**Description**

Plot lower triangle Pearson's R<sup>2</sup>. Diagonal text, upper triangle all against all scatter plots with lm abline

**Usage**

```
pairsAnnot(data, textCol = rep(1, ncol(data)), diagText = c(),
  col = rgb(0, 100, 0, 50, maxColorValue = 255), isHeatCol = F, ...)
```

**Arguments**

      data                  data.frame

**Details**

      No details

**Note**

      No note

**References**

      NA

**Examples**

      print("No examples")

---

`parseMaxQuantProteinGroupTxt`*Parse MaxQuant Protein Group Txt*

---

**Description**

Parse MaxQuant Protein Group Txt

**Usage**

```
parseMaxQuantProteinGroupTxt(file = file, expDesign = expDesign,  
  method = "auc")
```

**Arguments**

file	path to MaxQuant Protein txt file
expDesign	experimental design data.frame
method	auc (area under curve) or spc (spectral count)

**Details**

No details

**Value**

ExpressionSet object

**Note**

No note

**References**

NA

**See Also**

[ExpressionSet](#)

**Examples**

```
print("No examples")
```

---

`parseProgenesisFeatureCsv`*Parse Progenesis Feature Csv Export*

---

**Description**

Parse Progenesis Feature Csv Export

**Usage**

```
parseProgenesisFeatureCsv(file = file,  
  expDesign = getExpDesignProgenesisCsv(file), method = "auc")
```

**Arguments**

<code>file</code>	path to Progenesis Feature csv file
<code>expDesign</code>	experimental design data.frame
<code>method</code>	auc (area under curve) or spc (spectral count)

**Details**

No details

**Value**

ExpressionSet object

**Note**

No note

**References**

NA

**See Also**

[ExpressionSet](#)

**Examples**

```
print("No examples")
```

---

`parseProgenesisPeptideMeasurementCsv`*Parse Progenesis Peptide Measurement Csv Export*

---

## Description

Parse Progenesis Peptide Measurement Csv Export

## Usage

```
parseProgenesisPeptideMeasurementCsv(file, expDesign = expDesign,  
  method = "auc",  
  expressionColIndices = .getProgenesisCsvExpressionColIndices(file, method =  
    method))
```

## Arguments

file	path to Progenesis Peptide Measurement csv file
expDesign	experimental design data.frame
method	auc (area under curve) or spc (spectral count)

## Details

No details

## Value

ExpressionSet object

## Note

No note

## References

NA

## See Also

[ExpressionSet](#)

## Examples

```
print("No examples")
```

parseProgenesisProteinCsv

*Parse Progenesis Protein Csv*

---

## Description

Parse Progenesis Protein Csv

## Usage

```
parseProgenesisProteinCsv(file = file, expDesign = expDesign,  
  method = "auc")
```

## Arguments

file	path to Progenesis Protein csv file
expDesign	experimental design data.frame
method	auc (area under curve) or spc (spectral count)

## Details

No details

## Value

ExpressionSet object

## Note

No note

## References

NA

## See Also

[ExpressionSet](#)

## Examples

```
print("No examples")
```

---

`parseScaffoldPTMReport`*Parse scaffold PTM Spectrum Report*

---

**Description**

Parse scaffold PTM Spectrum Report

**Usage**

```
parseScaffoldPTMReport(file)
```

**Arguments**

file	path to Scaffold file
------	-----------------------

**Details**

No details

**Value**

data.frame

**Note**

No note

**References**

NA

**Examples**

```
print("No examples")
```

---

`parseScaffoldRawFile` *Parse scaffold output .xls file (RAW export)*

---

**Description**

Parse scaffold output .xls file (RAW export)

**Usage**

```
parseScaffoldRawFile(fileName, expDesign = expDesign,  
  keepFirstAcOnly = FALSE, isPurityCorrect = T)
```

**Arguments**

expDesign	experimental design data.frame
keepFirstAcOnly	TRUE/FALSE If multiple ACs in Accession.Numbers filed. Then keep the first one only
file	path to Scaffold file

**Details**

No details

**Value**

ExpressionSet object

**Note**

No note

**References**

NA

**See Also**

[ExpressionSet](#)

**Examples**

```
print("No examples")
```

---

perFeatureNormalization

*Per Feature Normalization*

---

**Description**

Per Feature Normalization

**Usage**

```
perFeatureNormalization(eset, normFactors)
```

**Arguments**

eset	ExpressionSet
matrix	normalization factors (logged) (row names are proteins)

**Details**

Example Usage: Normalize phospho peptide signals for Protein Changes



**Value**

ExpressionSet object

**Note**

No note

**References**

No references

**See Also**

[topX](#)

**Examples**

```
print("No examples")
```

---

plotAbsEstCalibrationCurve

*Plot absolut Estimation calibration Curve*

---

**Description**

Plot absolut Estimation calibration Curve

**Usage**

```
plotAbsEstCalibrationCurve(fit, dispElements = c("formula", "lowess",
  "stats"), xlab = "Conc. (CPC) ", ylab = "Pred. Conc. (CPC) ",
  predictorName = paste("log10(", names(coef(fit))[2], ")", sep = ""),
  text = F, cex.lab = 1, cex.axis = 1, cex.text = 1, cex.dot = 1, ...,
  main = "")
```

**Arguments**

fit	simple log-linear model
dispElements	c("formula","lowess","stats")
cex.lab=	expansion factor for axis labels
cex.axis=	expansion factor for axis
cex.text=	expansion factor for legend
cex.dot=	expansion factor for plotted dots

**Note**

No note

**References**

NA

**Examples**

```
print("No examples")
```

---

plotExpDesign	<i>Display experimental design, high-lighting the control condition</i>
---------------	---

---

**Description**

Display experimental design, high-lighting the control condition

**Usage**

```
plotExpDesign(eset, condColors = .getConditionColors(eset), version = "X")
```

**Arguments**

eset	ExpressionSet
------	---------------

**Details**

No details

**Note**

No note

**References**

NA

**Examples**

```
print("No examples")
```

---

plotIdScoreVsFDR	<i>Plot FDR vs. identification score</i>
------------------	--

---

**Description**

Plot FDR vs. identification score

**Usage**

```
plotIdScoreVsFDR(idScore, qvals, qvalueThrs = 0.01,
  ylab = "False Discovery Rate", xlab = "Identification Score", lwd = 1.5,
  ...)
```

**Arguments**

idScore	vector of identification scores
qvals	vector of q-valres
qvalueThrs	threshold indicated by horizontal line

**Details**

No details

**Note**

No note

**References**

NA

**Examples**

```
print("No examples")
```

---

```
plotMSSignalDistributions
```

*Plot ms.signal distributions*

---

**Description**

Plot ms.signal distributions

**Usage**

```
plotMSSignalDistributions(d, col = 1:100, cex.axis = 1, cex.lab = 1,  
  ylab = "Frequency", xlab = "MS-Signal", ...)
```

**Arguments**

matrix	matrix of ms-signals
color	color

**Details**

No details

**Note**

No note

**References**

NA

**Examples**

```
print("No examples")
```

---

```
plotNbIdentificationsVsRT
```

*Plot the number of identified Features per Reteintion Time minute.*

---

**Description**

Plot the number of identified Features per Reteintion Time minute.

**Usage**

```
plotNbIdentificationsVsRT(eset, cex.axis = 1.25, cex.lab = 1.25,
  col = "blue", lwd = 2, ...)
```

**Arguments**

eset	ExpressionSet
------	---------------

**Note**

No note

**References**

NA

**Examples**

```
print("No examples")
```

---

```
plotNbValidDeFeaturesPerFDR
```

*Plot Total Number of diffrentially Abundant Features (applying ratio cutoff) vs. qValue/pValue for all conditions*

---

**Description**

Plot Total Number of diffrentially Abundant Features (applying ratio cutoff) vs. qValue/pValue for all conditions

**Usage**

```
plotNbValidDeFeaturesPerFDR(sqa, upRegulated = T, log2RatioCufOff = log2(1),
  pvalRange = c(0, 0.3), pvalCutOff = 1, isLegend = T, isAdjusted = T,
  ylab = "Nb. Features", ...)
```

**Arguments**

sqa	SafeQuantAnalysis Object
upRegulated	TRUE/FALSE select for upregulated features
log2RatioCufOff	log2 ratio cut-off
pvalRange	pValue/qValue range
pvalCutOff	pValue/qValue cut-off
isLegend	TRUE/FALSE display legend
isAdjusted	TRUE/FALSE qValues/pValue on x-axis

**Details**

No details

**Note**

No note

**References**

NA

**Examples**

```
print("No examples")
```

---

plotPrecMassErrorDistrib

*Plot Precursor Mass Error Distribution*

---

**Description**

Plot Precursor Mass Error Distribution

**Usage**

```
plotPrecMassErrorDistrib(eset, pMassTolWindow = c(-10, 10), ...)
```

**Arguments**

eset	ExpressionSet
pMassTolWindow	Precursor Mass Error Tolerance Window

**Details**

No details

**Note**

No note

**References**

NA

**Examples**

```
print("No examples")
```

---

plotPrecMassErrorVsScore

*Plot precursorMass error v.s score highlighting decoy and displaying user specified user specified precursor mass filter*

---

**Description**

Plot precursorMass error v.s score highlighting decoy and displaying user specified user specified precursor mass filter

**Usage**

```
plotPrecMassErrorVsScore(eset, pMassTolWindow = c(-10, 10), ...)
```

**Arguments**

eset                      ExpressionSet  
pMassTolWindow   Precursor Mass Error Tolerance Window

**Details**

No details

**Note**

No note

**References**

NA

**Examples**

```
print("No examples")
```

---

plotQValueVsPValue	<i>Plot qValue vs pValue</i>
--------------------	------------------------------

---

**Description**

Plot qValue vs pValue

**Usage**

```
plotQValueVsPValue(sqa, lim = c(0, 1))
```

**Arguments**

sqa	SafeQuantAnalysis Object
lim	x-axis and y-axis range

**Details**

No details

**Note**

No note

**References**

NA

**Examples**

```
print("No examples")
```

---

plotROC	<i>Plot Number of Identifications vs. FDR</i>
---------	---

---

**Description**

Plot Number of Identifications vs. FDR

**Usage**

```
plotROC(qvals, qvalueThrs = 0.01, xlab = "False Discovery Rate",  
        ylab = "# Valid Identifications", xlim = c(0, 0.1), breaks = 100,  
        col = "blue", lwd = 1.5, ...)
```

**Arguments**

qvals	vector of q-values
qvalueThrs	threshold indicated by vertical line
breaks	see breaks for hist function

**Details**

No details

**Note**

No note

**References**

NA

**Examples**

```
print("No examples")
```

---

plotRTNorm

*Plot all retention time profile overalying ratios*

---

**Description**

Plot all retention time profile overalying ratios

**Usage**

```
plotRTNorm(rtNormFactors, eset, samples = 1:ncol(rtNormFactors), main = "",  
...)
```

**Arguments**

rtNormFactors	data.frame of normalization factor per r.t bin and sample, obtained by getRTNormFactors
eset	ExprssionSet
samples	specify samples (sample numbers) to be plotted

**Details**

No details

**Note**

No note

**References**

In Silico Instrumental Response Correction Improves Precision of Label-free Proteomics and Accuracy of Proteomics-based Predictive Models, Lyutvinskiy et al. (2013), <http://www.ncbi.nlm.nih.gov/pubmed/23589346>

**See Also**

[getRTNormFactors](#)



**Examples**

```
print("No examples")
```

---

plotRTNormSummary	<i>Plot all retention time normalization profiles</i>
-------------------	---

---

**Description**

Plot all retention time normalization profiles

**Usage**

```
plotRTNormSummary(eset,  
  col = as.character(.getConditionColors(eset)[pData(eset)$condition, 1]),  
  ...)
```

**Arguments**

rtNormFactors	data.frame of normalization factor per r.t bin and sample, obtained by getRTNormFactors
condNames	vector of condition names

**Details**

No details

**Note**

No note

**References**

In Silico Instrumental Response Correction Improves Precision of Label-free Proteomics and Accuracy of Proteomics-based Predictive Models, Lyutvinskiy et al. (2013), <http://www.ncbi.nlm.nih.gov/pubmed/23589346>

**See Also**

[getRTNormFactors](#)

**Examples**

```
print("No examples")
```

---

plotScoreDistrib	<i>Plot identifications target decoy distribution</i>
------------------	---

---

**Description**

Plot identifications target decoy distribution

**Usage**

```
plotScoreDistrib(targetScores, decoyScores, xlab = "Identification Score",
  ylab = "Counts", ...)
```

**Arguments**

targetScores  
decoyScores

**Details**

No details

**Note**

No note

**References**

NA

**Examples**

```
print("No examples")
```

---

plotVolcano	<i>Plots volcano, data points colored by max cv of the 2 compared conditions</i>
-------------	--

---

**Description**

Plots volcano, data points colored by max cv of the 2 compared conditions

**Usage**

```
plotVolcano(obj, ratioThrs = 1, pValueThreshold = 0.01, adjusted = T, ...)
```

**Arguments**

obj                    safeQuantAnalysis object or data.frame  
 adjusted               TRUE/FALSE plot qValues or pValues on y-axis  
 ratioCutOffAbsLog2       ratio abline  
 absLog10pValueCutOff    pValue abline

**Details**

data.frame input object should contain 3 columns (ratio,qValue,cv)

**Note**

No note

**References**

NA

**Examples**

```
print("No examples")
```

---

plotXYDensity	<i>Scatter plot with density coloring</i>
---------------	---

---

**Description**

Scatter plot with density coloring

**Usage**

```
plotXYDensity(x, y, isFitLm = T, legendPos = "bottomright",
  disp = c("abline", "R", "Rc"), ...)
```

**Arguments**

x                    number vector  
 y                    number vector

**Note**

No note

**References**

NA

**Examples**

```
print("No examples")
```

---

```
print.safeQuantAnalysis
```

*Print content of safeQuantAnalysis object*

---

### Description

Print content of safeQuantAnalysis object

### Usage

```
## S3 method for class 'safeQuantAnalysis'  
print(sqa)
```

### Arguments

sqa                      safeQuantAnalysis object

### Details

NA

### Note

No note

### References

NA

### See Also

[safeQuantAnalysis](#)

### Examples

```
print("No examples")
```

---

purityCorrectTMT

*Correct channel intensities based on Reporter ion Isotopic Distributions*

---

### Description

Correct channel intensities based on Reporter ion Isotopic Distributions

### Usage

```
purityCorrectTMT(tmtData, impurityMatrix = impurityMatrix,  
  invalidReplace = "allNA")
```

**Arguments**

tmtData	data.frame containing tmt channel intensities
method	to deal with NA and negative values c("", "allZero", "allNA", "allOrg")

**Details**

Same method as MSnbase, and described in Breitwieser et al. 2012 (Book Chapter)

**Value**

data.frame of corrected tmt intensities

**Note**

No note

**References**

NA

**Examples**

```
print("No examples")
```

---

removeOutliers	<i>Set value to NA if it deviates with more than 1.5 * IQR from lower/upper quantile</i>
----------------	--

---

**Description**

Set value to NA if it deviates with more than 1.5 \* IQR from lower/upper quantile

**Usage**

```
removeOutliers(x, na.rm = TRUE, ...)
```

**Arguments**

vector	numeric
a	logical indicating whether missing values should be removed.

**Details**

No details

**Value**

vector numeric

**Note**

No note

References

NA

See Also

NA

Examples

```
print("No examples")
```

---

rollUp	<i>Roll up feature intensities per unique column combination</i>
--------	--

---

Description

Roll up feature intensities per unique column combination

Usage

```
rollUp(eset, method = "sum", featureDataColumnName = c("proteinName"))
```

Arguments

- eset                      ExpressionSet
- method                  "sum", "mean" or "top3"
- featureDataColumnName  
                            vector of column names e.g. peptide or proteinName

Details

featureDataColumnName = c("peptide","charge","ptm"), method= c("sum"), sums up intensities per peptide modification charge state

Value

ExpressionSet object

Note

No note

References

No references

See Also

[topX](#)

Examples

```
print("No examples")
```

---

rtNormalize	<i>Normalization data per retention time bin</i>
-------------	--

---

**Description**

Normalization data per retention time bin

**Usage**

```
rtNormalize(eset, rtNormFactors)
```

**Arguments**

eset	ExpressionSet
rtNormFactors	obtained using getRTNormFactors

**Details**

Normalize for variations in electrospray ionization current.

**Value**

data.frame normalization factors per retention time bin (minute)

**Note**

No note

**References**

In Silico Instrumental Response Correction Improves Precision of Label-free Proteomics and Accuracy of Proteomics-based Predictive Models, Lyutvinskiy et al. (2013), <http://www.ncbi.nlm.nih.gov/pubmed/23589346>

**See Also**

[getRTNormFactors](#)

**Examples**

```
print("No examples")
```

---

`setNbPeptidesPerProtein`*Set nbPeptides coulumn of featureData*

---

**Description**

Set nbPeptides coulumn of featureData

**Usage**

```
setNbPeptidesPerProtein(eset)
```

**Arguments**

eset	ExpressionSet
------	---------------

**Details**

NA

**Value**

eset

**Note**

No note

**References**

NA

**Examples**

```
print("No examples")
```

---

`setNbSpectraPerProtein`*Set nbPeptides coulumn of featureData*

---

**Description**

Set nbPeptides coulumn of featureData

**Usage**

```
setNbSpectraPerProtein(eset)
```

**Arguments**

eset	ExpressionSet
------	---------------



**Details**

NA

**Value**

eset

**Note**

No note

**References**

NA

**Examples**

```
print("No examples")
```

---

sqNormalize	<i>Normalize</i>
-------------	------------------

---

**Description**

Normalize

**Usage**

```
sqNormalize(eset, method = "global")
```

**Arguments**

eset	ExpressionSet
method	c("global","rt","quantile")

**Details**

No details

**Value**

eset ExpressionSet

**Note**

No note

**References**

NA

See Also

getGlobalNormFactors, getRTNormFactors

Examples

```
print("No examples")
```

---

standardise	<i>Standardise data</i>
-------------	-------------------------

---

Description

Standardise data

Usage

```
standardise(d)
```

Arguments

d                      vector or data.frame or matrix

Details

No details

Value

vector or data.frame or matrix

Note

No note

Examples

```
print("No examples")
```

---

stripACs	<i>strip uniprot format e.g. "sp Q8CHJ2 AQP12_MOUSE" -&gt; Q8CHJ2</i>
----------	---

---

**Description**

strip uniprot format e.g. "sp|Q8CHJ2|AQP12\_MOUSE" -> Q8CHJ2

**Usage**

```
stripACs(acs)
```

**Arguments**

acs	accession numbers
-----	-------------------

**Details**

TRUE if less than 10

**Value**

vector character

**Note**

No note

**References**

NA

**Examples**

```
print("No examples")
```

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