

Package ‘SafeQuant’

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Type Package

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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

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R topics documented:

addIdQvalues	3
addScaffoldPTMAnnotations	4
barplotMSSignal	4
createCalibrationCurve	5
createExpDesign	6
createExpressionDataset	7
cvBoxplot	8
expDesignTagToExpDesign	9
export.safeQuantAnalysis	9
getAAProteinCoordinates	10
getAllCV	11
getAllEBayes	12
getBaselineIntensity	13
getCV	13
getExpDesignProgenesisCsv	14
getGlobalNormFactors	15
getIBAQEset	15

getIdLevelQvals	16
getImpuritiesMatrix	17
getIntSumPerProtein	18
getLoocvFoldError	19
getMeanCenteredRange	20
getModifProteinCoordinates	20
getMotifX	21
getNbDetectablePeptides	22
getNbMisCleavages	23
getNbPeptidesPerProtein	23
getNbSpectraPerProtein	24
getPeptides	25
getRatios	25
getRTNormFactors	26
getScoreCutOff	27
getSignalPerCondition	27
getTopX	28
getUserOptions	29
globalNormalize	30
hClustHeatMap	31
isCon	31
isDecoy	32
isStrippedACs	33
missinValueBarplot	34
pairsAnnot	34
parseMaxQuantProteinGroupTxt	35
parseProgenesisFeatureCsv	36
parseProgenesisPeptideMeasurementCsv	37
parseProgenesisProteinCsv	38
parseScaffoldPTMReport	39
parseScaffoldRawFile	39
perFeatureNormalization	40
plotAbsEstCalibrationCurve	41
plotExpDesign	42
plotIdScoreVsFDR	42
plotMSSignalDistributions	43
plotNbIdentificationsVsRT	44
plotNbValidDeFeaturesPerFDR	45
plotPrecMassErrorDistrib	46
plotPrecMassErrorVsScore	46
plotQValueVsPValue	47
plotROC	48
plotRTNorm	48
plotRTNormSummary	49
plotScoreDistrib	50
plotVolcano	51
plotXYDensity	52
print.safeQuantAnalysis	52
purityCorrectTMT	53
removeOutliers	54
rollUp	55
rtNormalize	56

<i>addIdQvalues</i>	3
setNbPeptidesPerProtein	57
setNbSpectraPerProtein	57
sqNormalize	58
standardise	59
stripACs	60
Index	61

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 annotations to tmt experiment

Description

Add scaffold ptm annotations 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

eset	expressionSet
col	default condition colors
method	c("median","sum","sharedSignal")
...	

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 *Create Experimental Design*

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

C.V. boxplot

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
col	col
cex.names	default 0.9
cex.axis	default 1.25
cex.lab	default 1.25
ylab	C.V.
...	

Details

No details

Note

No note

References

NA

Examples

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

expDesignTagToExpDesign

Create experimental design data.frame from user input string

Description

Create experimental design data.frame from user input string

Usage

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

Arguments

tag	tag
expDesign	data.frame

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 ordered 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 (modification residue number)

Note

No note

References

NA

Examples

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

getAllCV

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

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

Description

Perform statistical test (moderated 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
method	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 *Get signal at zscore x (x standard deviations below mean)*

Description

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

Usage

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

Arguments

intensities	reference run signals
percentile	baseline value set as specified promille

Value

baseline value

Note

No note

References

NA

Examples

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

getCV *Calculate Coefficient of Variance per feature (Relative standard Deviation)*

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

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

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

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

Details

No details

Value

ExpressionSet

Note

No note

References

Global quantification of mammalian gene expression control, Schwanhausser (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")
```

<code>getIdLevelQvals</code>	<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

<code>scores</code>	peptide/protein identificationscore
<code>isDecoy</code>	vector of TRUE/FALSE

Details

q-value = (Nb. Decoy Entries at idScore Threshold S*) / (Nb. Target Entries at idScore Threshold S). (* idScore >= S)

Value

vector of q.values

Note

No note

References

NA

Examples

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

getImpuritiesMatrix *Get Thermo TMT impurity matrix*

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

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

<code>intData</code>	data.frame of intensities per channel
<code>proteinACs</code>	vector of protein accession numbers
<code>peptides</code>	vector of peptide sequences
<code>minNbPeptPerProt</code>	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 *Leave-One-Out Cross Validate Qunatification Model*

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 from mean

Details

NA

Value

vector range boundaries

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
format	c(1,2) 1. progenesis 2. scaffold

Details

NA

Value

vector of protein coordinates (modification 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

<code>peptides</code>	list of peptides
<code>vector</code>	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      Get number of mis-cleavages per peptide
```

Description

Get number of mis-cleavages per 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      Get number of peptides per protein
```

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	<i>Digest protein</i>
-------------	-----------------------

Description

Digest protein

Usage

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

Arguments

proteinSeq protein sequence

Details

No details

Value

vector of peptides

Note

No note

Examples

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

getRatios	<i>Calculate ratios, comparing all case to control</i>
-----------	--

Description

Calculate ratios, comparing all case to control

Usage

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

Arguments

eset ExpressionSet
method median or mean

Details

No details

Value

ExpressionSet object

Note

No note

References

NA

Examples

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

getRTNormFactors

Get retentiontime base normalization factors

Description

Get retentiontime base normalization factors

Usage

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

Arguments

eset ExpressionSet
minFeaturesPerBin

minimum 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

<code>method</code>	median (default), mean, max, min, sd
<code>data</code>	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

<code>entryData</code>	data.frame listing feature intensities of one entry. Typically rows corresponds to Peptide entries of one protein
<code>topX</code>	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*Read User Specified Command Line Options*

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

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")
```

<code>hClustHeatMap</code>	<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

<code>eset</code>	ExpressionSet
<code>conditionColors</code>	data.frame of colors per condition
<code>breaks</code>	default seq(-2,2,length=20)
...	

Details

No details

Note

No note

References

NA

Examples

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

<code>isCon</code>	<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

contaminants 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 *Plot Percentage of Features with with missing values*

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
col	col
cex.axis	cex.axis
cex.lab	cex.lab
...	

Details

No details

Note

No note

References

NA

Examples

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

pairsAnnot *Plot lower triangle Pearsons R^2. Diagonal text, upper triangle all against all scatter plots with lm abline*

Description

Plot lower triangle Pearsons R^2. Diagonal text, upper triangle all against all scatter plots with lm abline

Usage

```
pairsAnnot(data, textCol = rep(1, ncol(data)), diagText = colnames(data),  
           col = rgb(0, 100, 0, 50, maxColorValue = 255), isHeatCol = F,  
           cexTxt = 2, ...)
```

Arguments

data	data.frame
textCol	text color
diagText	diagonal text
col	dot col
isHeatCol	heat colors
cexTxt	cex txt
...	

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

file	path to Progenesis Feature 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")
```

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

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

<code>eset</code>	ExpressionSet
<code>matrix</code>	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, ...)
```

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*Display experimental design, high-lighting the control condition***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*Plot FDR vs. identification score***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-values
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, ylab = "Frequunecy",
                           xlab = "MS-Signal", ...)
```

Arguments

d	matrix of ms-signals
col	color
ylab	default "Frequunecy"
xlab	default "MS-Signal"
...	

Details

No details

Note

No note

References

NA

Examples

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

```
plotNbIdentificationsVsRT
```

Plot the number of identified Features per Retention Time minute.

Description

Plot the number of identified Features per Retention Time minute.

Usage

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

Arguments

eset	ExpressionSet
cex.axis	default 1.25
cex.lab	default 1.25
col	default "blue"
lwd	default 2
...	

Note

No note

References

NA

Examples

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

plotNbValidDeFeaturesPerFDR

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

Description

Plot Total Number of differentially 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 *Plot qValue vs pValue*

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`*Plot Number of Identifications vs. FDR***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

<code>qvals</code>	vector of q-values
<code>qvalueThrs</code>	threshold indicated by vertical line
<code>breaks</code>	see breaks for hist function
...	

Details

No details

Note

No note

References

NA

Examples

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

`plotRTNorm`*Plot all retention time profile overlaying ratios***Description**

Plot all retention time profile overlaying 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 getRT-
                NormFactors
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

Plot all retention time normalization profiles

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 getRT-
                NormFactors
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 *Plot identifications target decoy distribution*

Description

Plot identifications target decoy distribution

Usage

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

Arguments

```
targetScores  
decoyScores  
xlab           default "Identification Score"  
ylab           default "Counts"  
...  
.
```

Details

No details

Note

No note

References

NA

Examples

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

plotVolcano

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

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
ratioThrs	default 1
pValueThreshold	default 0.01
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*Scatter plot with density coloring***Description**

Scatter plot with density coloring

Usage

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

Arguments

<code>x</code>	number vector
<code>y</code>	number vector
<code>isFitLm</code>	fit linear model
<code>disp</code>	c("abline","R","Rc") display options
...	

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

<code>sqa</code>	safeQuantAnalysis object
------------------	--------------------------

Details

NA

Note

No note

References

NA

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

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

purityCorrectTMT	<i>Correct channel intensities based on Reporter ion Isotopic Distributions</i>
------------------	---

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
impurityMatrix	correction matrix
method	to deal with NA and negative values c("", "allZero", "allNA", "allOrg")

Details

Same method as MSnbbase, 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

na.rm	logical indicating whether missing values should be removed.
vector	numeric

Details

No details

Value

vector numeric

Note

No note

References

NA

See Also

NA

Examples

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

rollUp*Roll up feature intensites per unique colum combination*

Description

Roll up feature intensites per unique colum 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 peptie modification charge state

Value

ExpressionSet object

Note

No note

References

No references

See Also

[topX](#)

Examples

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

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

Description

Normalization data per retention time bin

Usage

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

Arguments

<code>eset</code>	ExpressionSet
<code>rtNormFactors</code>	obtained using <code>getRTNormFactors</code>

Details

Normalize for variations in elelctrospray 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 column of featureData

Description

Set nbPeptides coulmn of featureData

Usage

```
setNbPeptidesPerProtein(eset)
```

Arguments

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

Details

NA

Value

eset

Note

No note

References

NA

Examples

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

```
setNbSpectraPerProtein
```

Set nbPeptides column of featureData

Description

Set nbPeptides coulmn of featureData

Usage

```
setNbSpectraPerProtein(eset)
```

Arguments

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

Details

NA

Value

eset

Note

No note

References

NA

Examples

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

sqNormalize*Normalize*

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`

Standardise data

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 *strip uniprot format e.g. "sp|Q8CHJ2|AQP12_MOUSE" -> Q8CHJ2*

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")
```

Index

*Topic **normalization**
 getGlobalNormFactors, 15
 getLoocvFoldError, 19
 globalNormalize, 30
 removeOutliers, 54
 sqNormalize, 58

 addIdQvalues, 3
 addScaffoldPTMAnnotations, 4

 barplotMSSignal, 4

 createCalibrationCurve, 5
 createExpDesign, 6
 createExpressionDataset, 7
 cvBoxplot, 8

 eBayes, 12
 expDesignTagToExpDesign, 9
 export.safeQuantAnalysis, 9
 ExpressionSet, 7, 36–38, 40

 getAAProteinCoordinates, 10
 getAllCV, 11
 getAllEBayes, 12
 getBaselineIntensity, 13
 getCV, 11, 13
 getExpDesignProgenesisCsv, 14
 getGlobalNormFactors, 15
 getIBAQEset, 15
 getIdLevelQvals, 3, 16
 getImpuritiesMatrix, 17
 getIntSumPerProtein, 18
 getLoocvFoldError, 19
 getMeanCenteredRange, 20
 getModifProteinCoordinates, 20
 getMotifX, 21
 getNbDetectablePeptides, 22
 getNbMisCleavages, 23
 getNbPeptidesPerProtein, 23
 getNbSpectraPerProtein, 24
 getPeptides, 25
 getRatios, 25
 getRTNormFactors, 26, 49, 50, 56
 getScoreCutOff, 27

 getSignalPerCondition, 27
 getTopX, 28
 getUserOptions, 29
 globalNormalize, 30

 hClustHeatMap, 31

 isCon, 31
 isDecoy, 32
 isStrippedACs, 33

 missinValueBarplot, 34

 pairsAnnot, 34
 parseMaxQuantProteinGroupTxt, 35
 parseProgenesisFeatureCsv, 36
 parseProgenesisPeptideMeasurementCsv, 37
 parseProgenesisProteinCsv, 38
 parseScaffoldPTMReport, 39
 parseScaffoldRawFile, 39
 perFeatureNormalization, 40
 plotAbsEstCalibrationCurve, 41
 plotExpDesign, 42
 plotIdScoreVsFDR, 42
 plotMSSignalDistributions, 43
 plotNbIdentificationsVsRT, 44
 plotNbValidDeFeaturesPerFDR, 45
 plotPrecMassErrorDistrib, 46
 plotPrecMassErrorVsScore, 46
 plotQValueVsPValue, 47
 plotROC, 48
 plotRTNorm, 48
 plotRTNormSummary, 49
 plotScoreDistrib, 50
 plotVolcano, 51
 plotXYDensity, 52
 print.safeQuantAnalysis, 52
 purityCorrectTMT, 53

 removeOutliers, 54
 rollUp, 55
 rtNormalize, 56

 safeQuantAnalysis, 10, 53

setNbPeptidesPerProtein, [57](#)

setNbSpectraPerProtein, [57](#)

sqNormalize, [58](#)

standardise, [59](#)

stripACs, [60](#)

topX, [41](#), [55](#)