Title  Analysis and quantitation of isobarically tagged MSMS proteomics data

Description  isobar provides methods for preprocessing, normalization, and report generation for the analysis of quantitative mass spectrometry proteomics data labeled with isobaric tags, such as iTRAQ and TMT. Features modules for integrating and validating PTM-centric datasets (isobar-PTM). More information on http://www.ms-isobar.org.

Version  1.22.0

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biocViews  Proteomics, MassSpectrometry, Bioinformatics, MultipleComparisons, QualityControl

Depends  R (>= 2.10.0), Biobase, stats, methods

Imports  distr, plyr, biomaRt, ggplot2

Suggests  MSnbase, OrgMassSpecR, XML, RJJSONIO, Hmisc, gplots, RColorBrewer, gridExtra, limma, boot, DBI, MASS

LazyLoad  yes

License  LGPL-2

URL  https://github.com/fbreitwieser/isobar

BugReports  https://github.com/fbreitwieser/isobar/issues


NeedsCompilation  no

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isobar-package

Analysis and quantitation of isobarically tagged MSMS proteomics data

Description

isobar provides methods for preprocessing, normalization, and report generation for the analysis of quantitative mass spectrometry proteomics data labeled with OA isobaric tags, such as iTRAQ and TMT.

Details

Package: isobar
Version: 1.1.2
biocViews: Proteomics, MassSpectrometry, Bioinformatics, MultipleComparisons, QualityControl
Depends: R (>= 2.9.0), Biobase, stats, methods, ggplot2
Imports: distr, biomaRt
Suggests: MSnbase, XML
LazyLoad: yes
License: LGPL-2
URL: http://bioinformatics.cemm.oeaw.ac.at

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Description

Calculates delta score from raw search engine score by subtracting the best matching hit with the second best matching. Data needs to have not only the best hit per spectrum, but multiple, to be able to calculate the delta score. filterSpectraDeltaScore calls calc.delta.score and filters spectra below a minimum delta score.

Usage

calc.delta.score(my.data)
filterSpectraDeltaScore(my.data, min.delta.score=10, do.remove=FALSE)

Arguments

my.data IBSpectra data frame.
min.delta.score Minimum delta score.
do.remove If TRUE, spectra below the min.prob threshold are not just set as 'use.for.quant=FALSE' but removed.

Value

Returns data with additional column 'delta.score'.

Author(s)

Florian P. Breitwieser
calcPeptidePosition

Recalculate peptide start positions based on protein sequence

Description
Function to recalculate start position of peptide in protein when it is missing or wrong.

Usage
calcPeptidePosition(peptide.info, protein.info, calc.il.peptide)

Arguments
peptide.info  Peptide info object of ProteinGroup.
protein.info  Protein info object of ProteinGroup.
calc.il.peptide  Should the 'real' peptide (I/L difference) be calculated?

calculate-pvalues

Calculate and Adjust Ratio and Sample p-values.

Description
Functions for calculating and adjusting ratios and sample p-values. Usually, these are called by proteinRatios or peptideRatios.

Usage
calculate.ratio.pvalue(lratio, variance, ratiodistr = NULL)
calculate.sample.pvalue(lratio, ratiodistr)
calculate.mult.sample.pvalue(lratio, ratiodistr, strict.pval, lower.tail, n.possible.val, n.observed.val)
adjust.ratio.pvalue(quant.tbl, p.adjust, sign.level, globally = FALSE)

Arguments
lratio  log 10 protein or peptide ratios.
ratiodistr  Fitted ratio distribution/
variance  Variance of lratios.
strict.pval  If FALSE, missing ratios are ignored. If TRUE, missing ratios are penalized by giving them a sample.pval of 0.5.
lower.tail  lower.tail of distribution?
n.possible.val  Number of possible ratios.
n.observed.val  Number of observed ratios.
quant.tbl  Quantification table (from proteinRatios or peptideRatios).
p_adjust  p-value adjustment method (see ?p.adjust).

sign_level  Ratio significance level.

globally  Whether the p-values should be adjusted over all conditions, or individually in each condition.

Author(s)
Florian P. Breitwieser

See Also
proteinRatios, peptideRatios

Examples

lratio <- c(-1,-1,seq(from=-1,to=1,by=.25),1,1)
variance <- c(0,1,rep(0.1,9),0,1)
ratiodistr.precise <- new("Norm",mean=0,sd=.25)
ratiodistr.wide <- new("Norm",mean=0,sd=.5)

# ratio p-value is impacted only by the variance
# sample p-value captures whether the ratio distribution is narrow ('precise')
# or wide
data.frame(lratio, variance,
    ratio.pvalue=calculate.ratio.pvalue(lratio, variance),
    sample.pvalue.precise=calculate.sample.pvalue(lratio,ratiodistr.precise),
    sample.pvalue.wide=calculate.sample.pvalue(lratio,ratiodistr.wide))


calculate.dNSAF  dNSAF approximate abundance calculations.

Description
Distributed normalized spectral abundance factor (dNSAF) is a label free quantitative measure of protein abundance based on spectral counts which are corrected for peptides shared by multiple proteins. Original publication: Zhang Y et al., Analytical Chemistry (2010).

Usage

calculate.dNSAF(protein.group, use.mw = FALSE, normalize = TRUE,
    combine.f = mean)

Arguments

protein.group  ProteinGroup object. Its @proteinInfo slot data.frame must contain a length column.

use.mw  Use MW to account for protein size

normalize  Normalize dSAF to dNSAF?

combine.f  How to handle proteins seen only with shared peptides?
calculate.emPAI

Value

Named numeric vector of dNSAF values.

Author(s)

Florian P Breitwieser

References

Zhang Y et al., Analytical Chemistry (2010)

See Also

proteinInfo, getProteinInfoFromUniprot, calculate.emPAI, ProteinGroup

Examples

data(ibspiked_set1)
protein.group <- proteinGroup(ibspiked_set1)
calculate.dNSAF(protein.group)

Description

The Exponentially Modified Protein Abundance Index (emPAI) is a la
bel free quantitative measure of protein abundance based on pro
tein coverage by peptide matches. The original publication is

Usage

calculate.emPAI(protein.group, protein.g = reporterProteins(protein.group), normalize = FALSE,
observed.pep = c("pep", "mod.charge.pep"), use.mw = FALSE, combine.f = mean,
..., nmc = 0, report.all = FALSE)

Arguments

protein.group ProteinGroup object. Its @proteinInfo slot data.frame must contain a sequence
column to calculate the number of observable peptides per protein.
protein.g Protein group identifiers.
normalize Normalize to sum = 1?.
observed.pep What counts as observed peptide?
report.all TOADD
use.mw Use MW to normalize for protein size
combine.f How to handle proteins seen only with shared peptides?
correct.peptide.ratios

seq          Protein sequence.
nmc          Number of missed cleavages.
min.length   Minimum length of peptide.
min.mass     Minimum mass of peptide.
max.mass     Maximum mass of peptide.
custom       User defined residue for Digest.
...          Further arguments to observable.peptides/Digest.

Details

The formula is:

\[ emPAI = 10^{\frac{N_{\text{observed}}}{N_{\text{observable}}} - 1} \]

\( N_{\text{observed}} \) is the number of observed peptides - we use the count of unique peptide without consideration of charge state. \( N_{\text{observable}} \) is the number of observable peptides. Sequence cleavage is done using Digest.

Value

Named numeric vector of emPAI values.

Author(s)

Florian P Breitwieser

References


See Also

Digest, proteinInfo, getProteinInfoFromUniprot, calculate.dNSAF, ProteinGroup

Examples

```r
data(ibspiked_set1)
protein.group <- proteinGroup(ibspiked_set1)
calculate.emPAI(protein.group, protein.g=protein.g(protein.group,"CERU"))
```

correct.peptide.ratios

Correct peptide ratios with protein ratios from a separate experiment.

Description

Correct peptide ratios with protein ratios from a separate experiment.

Usage

```r
correct.peptide.ratios(ibspectra, peptide.quant.tbl, protein.quant.tbl,
protein.group.combined, adjust.variance = TRUE,
correlation = 0, recalculate.pvalue = TRUE)
```
Arguments

- `ibspectra`: IBSpectra object. Calculated with peptideRatios.
- `peptide.quant.tbl`: Calculated with peptideRatios.
- `protein.quant.tbl`: Calculated with proteinRatios.
- `protein.group.combined`: ProteinGroup object generated on both PTM and protein data.
- `adjust.variance`: Adjust variance of ratios.
- `correlation`: Assumed correlation between peptide and protein ratios for variance adjustment.
- `recalculate.pvalue`: Recalculate p-value after variance adjustment.

Author(s)

Florian P.Breitwieser

---

distr-methods

Functions for distribution calculations

Description

calcProbXGreaterThanY calculates the probability that X >= Y. calcProbXDiffNormals calculates the probabilities of a set of normals, defined by the vectors mu_Y and sd_Y are greater or less than the reference distribution Y.

Usage

calcProbXGreaterThanY(X, Y, rel.tol = .Machine$double.eps^0.25, subdivisions = 100L)
calcProbXDiffNormals(X, mu_Y, sd_Y, ..., alternative = c("greater", "less", "two-sided"), progress = FALSE)
#calcCumulativeProbXGreaterThanY(Xs, mu_Ys, sd_Ys, alternative = c("greater", "less", "two-sided"), subdivisions = 100L, rel.tol = .Machine$double.eps^0.25)
distrprint(X, round.digits = 5)
twodistr.plot(X, Y, n.steps = 1000, min.q = 10^-3)

Arguments

- `X`: Object of the class Distribution.
- `Y`: Object of the class Distribution.
- `min.q`: Minimum quantile.
- `n.steps`: Number of steps.
- `mu_Y`: Numeric vector of parameter mu of a Normal.
- `sd_Y`: Numeric vector of parameter sd of a Normal.
- `subdivisions`: The maximum number of subintervals.
- `rel.tol`: Relative accuracy requested.
- `...`: Additional arguments to calcProbXGreaterThanY.
- `alternative`: "less", "greater", or "two-sided".
- `progress`: Show text progress bar?
- `round.digits`: Round digits for printing.
**fit distributions**

**Author(s)**
Florian P. Breitwieser

**Examples**

```r
library(distr)
calcProbXGreaterThanY(Norm(0,.25),Norm(1,.25))
```

**Description**

Functions to fit the probability density functions on ratio distribution.

**Usage**

```r
fitCauchy(x)
fitNorm(x, portion = 0.75)
fitWeightedNorm(x, weights)
fitNormalCauchyMixture(x)
fitGaussianMixture(x, n = 500)
fitTlsd(x)
```

**Arguments**

- `x` Ratios
- `weights` Weights
- `portion` Central portion of data to take for computation
- `n` number of sampling steps

**Value**

`Cauchy, Norm`

**Author(s)**
Florian P Breitwieser, Jacques Colinge.

**See Also**

`proteinRatios`
getPeptideModifContext

**Examples**

```r
library(distr)
data(ibspiked_set1)
data(noise.model.hcd)
# calculate protein ratios of Trypsin and CERU_HUMAN. Note: this is only
# for illustration purposes. For estimation of sample variability, data
# from all protein should be used
pr <- proteinRatios(ibspiked_set1, noise.model=noise.model.hcd,
  cl=as.character(c(1,1,2,2)),combn.method="intraclass",protein=c("136429","P00450"))

# fit a Cauchy distribution
ratiodistr <- fitCauchy(pr$lratio)
plot(ratiodistr)
```

---

**getPeptideModifContext**

*Get context of modification*

**Description**

Gets neighboring amino acids around modification which can be used to find enriched motifs.

**Usage**

```r
getPeptideModifContext(protein.group, modif, n.aa.up = 7, n.aa.down = 7)
```

**Arguments**

- **protein.group**  ProteinGroup object.
- **modif**          Modification of interest.
- **n.aa.up**        Number of AA downstream to report.
- **n.aa.down**      Number of AA upstream to report.

---

**getPhosphoRSProbabilities**

*Generate input files for PhosphoRS, call it, and get modification site probabilities*

**Description**

Get phosphorylation site localization probabilities by calling PhosphoRS and parsing its output. getPhosphoRSProbabilities generates a XML input file for PhosphoRS calling writePhosphoRSInput, then executes phosphoRS.jar with java, and parses the XML result file with readPhosphoRSOutput.
getPhosphoRSProbabilities

Usage

getPhosphoRSProbabilities(id.file, mgf.file, massTolerance, activationType, simplify = FALSE, mapping.file = NULL, mapping = c(peaklist = "even", id = "odd"), pepmodif.sep = "##.##", besthit.only = TRUE, phosphors.cmd = paste("java -jar", system.file("phosphors", "phosphoRS.jar", package = "isobar")), file.basename = tempfile("phosphors."))


readPhosphoRSOutput(phosphoRS.outfile, simplify = FALSE, pepmodif.sep = "##.##", besthit.only = TRUE)

filterSpectraPhosphoRS(id.file, mgf.file, ..., min.prob = NULL, do.remove=FALSE)

Arguments

id.file Database search results file in ibspectra.csv or mzIdentML format. See IBSpectra and isobar vignette for information on converting Mascot dat and Phenyx pidres files into ibspectra format.

mgf.file Peaklist file

massTolerance Fragment ion mass tolerance (in Da)

activationType Activation types of spectra. CID, HCD, or ETD.

simplify If TRUE, returns a data.frame instead of a list.

mapping.file Mapping file. See also readIBSpectra.

mapping Mapping columns.

besthit.only Only show best hit, simplifies result to data.frame instead of list.

phosphors.cmd PhosphoRS script.

file.basename Base name for creating phosphoRS input and output files.

phosphoRS.infile PhosphoRS input XML file name.

phosphoRS.outfile PhosphoRS output XML file name.
### getPtmInfo

**pepmodif.sep** separator of peptide and modification in XML id

**modif.masses** masses and ID used for PhosphoRS

**min.prob** Threshold for PhosphoRS peptide probability to consider it for quantification

**do.remove** If TRUE, spectra below the min.prob threshold are not just set as ’use.for.quant=FALSE’ but removed.

### Details

PhosphoRS is described in Taus et al., 2011. It can be downloaded from [http://cores.imp.ac.at/protein-chemistry/download/](http://cores.imp.ac.at/protein-chemistry/download/) and used as Freeware. Java is required at runtime.

### Value

If `simplify=TRUE`, a data.frame with the following columns: `spectrum`, `peptide`, `modif`, `PepScore`, `PepProb`, `seqpos`

If `simplify=FALSE`, a list (of spectra) of lists (of peptide identifications) of lists (with information about identification and localization). `spectrum` -> peptide 1, peptides 2, ... -> peptide. First level: - spectrum Second level: - peptide identifications for spectrum (might be more than one) Third level: - peptide: vector with peptide sequence and modification string - site.probs: matrix with site probabilities for each phospho site - isoforms: peptide score and probabilities for each isoform

### Author(s)

Florian P Breitwieser

### References

Taus et al., 2011

---

**getPtmInfo**

*Get PTM site information for identified proteins from public databases.*

**Description**

Get PTM site information for identified proteins from public databases.

**Usage**

```r

getPtmInfoFromNextprot(protein.group, nextprot.url = "http://www.nextprot.org/rest/entry/NX_XXX/ptm?format=json", url.wildcard = "XXX")
```
getPtmInfo

Arguments

protein.group ProteinGroup object.
file.name File name to save downloaded data, defaults to the original file name (see mapping).
modif Selects dataset to download (see mapping).
psp.url PhosphoSitePlus main URL for datasets.
mapping Names of PhosphoSitePlus modification datasets, mapped by modif name.
nextprot.url URL for fetching Nextprot results.
url.wildcard wildcard to replace with Uniprot Protein AC in nextprot.url.

Details

PhosphoSitePlus datasets are downloaded and written to the working directory with its original name (see mapping) unless a file with that name exists, which is then parsed into a data.frame of suitable format.

Value

data.frame with (at least) the columns: isoform_ac, description, evidence, position

Note

PhosphoSitePlus is licensed under Creative Commons Attribution-NonCommercial-ShareAlike 3.0 Unported License and is freely available for non-commercial purpose, see http://www.phosphosite.org/staticDownloads.do.

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Please read the conditions and use the data only if you agree.

Author(s)

Florian P. Breitwieser

References


Examples

```r
## Not run:
data(ib_phospho)
ptm.info.np <- getPtmInfoFromNextprot(proteinGroup(ib_phospho))
ptm.info.np <- ptm.info.np[grep("Phospho", ptm.info.np$modification.name),]
ptm.info.psp <- getPtmInfoFromPhosphoSitePlus(proteinGroup(ib_phospho), modif="PHOS")
```
**groupMemberPeptides**

For a given reporter protein group identifier, information on its peptides is returned. It contains information on how the peptides are shared and in which member they occur.

**Usage**

```r
groupMemberPeptides(x, reporter.protein.g, ordered.by.pos = TRUE, only.first.pos = TRUE)
```

**Arguments**

- `x` : ProteinGroup object
- `reporter.protein.g` : group reporter protein
- `ordered.by.pos` : if TRUE, start position of peptides in proteins is exported and peptides are ordered by position
- `only.first.pos` : if TRUE, only first occurrence of peptide in protein is reported

**Value**

list of two: 
- `[1] peptide.info: data.frame peptide specificity n.shared.groups n.shared.proteins start.pos`
- `[2] group.member.peptides: data.frame each column corresponds to a group member, and each row to a peptide`

**Author(s)**

Florian P Breitwieser

**Examples**

```r
data(ibspiked_set1)
protein.group <- proteinGroup(ibspiked_set1)
ceru.rat <- protein.g(protein.group,"CERU_RAT")
groupMemberPeptides(protein.group,ceru.rat)

## find protein groups with members

t <- table(proteinGroupTable(protein.group)$reporter.protein)
t[t>2]
protein.g <- names(t)[t>2][1]
groupMemberPeptides(protein.group,protein.g)
```
Description

Gather human readable information from protein group codes.

Usage

my.protein.info(x, protein.g)

human.protein.names(my.protein.info)

Arguments

x  ProteinGroup object
protein.g  protein
my.protein.info  Return value of function my.protein.info

Author(s)

Florian P Breitwieser

IBSpectra-class  IBSpectra Class for Isobarically Tagged Quantitative MS Proteomics Data

Description

This class represents a quantitative MS proteomics experiment labeled using Isobaric tags (iTRAQ, TMT). IBSpectra is a abstract class which is implemented in the IBSpectraTypes classes iTRAQ4plexSpectra, iTRAQ8plexSpectra, TMT2plexSpectra, TMT6plexSpectra and TMT10plexSpectra.

It contains per-spectrum measurements of the reporter tag intensity and m/z in assayData, and protein grouping in proteinGroup.

Objects from the Class

IBSpectra objects are typically created using the readIBSpectra method or by calls of the form new("iTRAQ4plexSpectra",data=NULL,data.ions=NULL,...).
Slots

IBSpectra extends eSet which is a container for high-throughput assays and experimental metadata. Slots introduced in eSet (for more details on slots and methods refer to eSet help):

- **assayData**: Contains matrices 'ions' and 'mass storing reporter tag intensities and m/z values for each tag and spectrum. Can be accessed by `reporterIntensities` and `reporterMasses`. Class: AssayData
- **phenoData**: Contains experimenter-supplied variables describing phenotypes behind reporter tags. Class: AnnotatedDataFrame-class
- **featureData**: Describes the spectra's retention time, charge, peptide sequence, etc and can be accessed by `fData`. Class: AnnotatedDataFrame
- **experimentData**: Contains details of experimental methods. Class: MIAME
- **annotation**: UNUSED. Label associated with the annotation package used in the experiment. Class: character
- **protocolData**: UNUSED. Contains equipment-generated variables describing reporter tags. Class: AnnotatedDataFrame
- **log**: character matrix logging isotope impurity correction, normalization, etc.

Slots introduced in IBSpectra:

- **proteinGroup**: A ProteinGroup object describing peptide and protein identifications grouped by shared peptides.
- **reporterTagNames**: A character vector denoting the reporter tag labels.
- **reporterMasses**: The 'true' m/z of the reporter tags in the MS/MS spectrum, used to isolate m/z-intensity pairs from peaklist.
- **isotopeImpurities**: Manufacturer supplied isotope impurities, need to be set per batch and used for correction by `correctIsotopeImpurities`.

Constructor

See `readIBSpectra` for creation based on peaklist (e.g. MGF format) and identification files (Mascot and Phenyx output).

new(type, data)

- **type**: Denotes the type of IBSpectra, either 'iTRAQ4plexSpectra','iTRAQ8plexSpectra','TMT2plexSpectra', 'TMT6plexSpectra' or 'TMT10plexSpectra'. Call IBSpectraTypes() to see a list of the implemented types.
- **data**: A 'data.frame' in a ibspectra-csv format.

Coercion

In the code snippets below, x is a IBSpectra object. IBSpectra object can be coerced to

as(x, "data.frame"): Creates a data.frame containing all identification and quantitation information. Peptide matching to multiple proteins produce multiple lines.

ibSpectra.as.concise.data.frame(x): Creates a data.frame containing all identification and quantitation information. Proteins are concatenated - so the resulting data.frame has one line per spectrum.

as(x, "MSnSet"): Coerces to a MSnSet object (package MSnbase).

as(msnset, "IBSpectra"): Coerces a MSnSet to IBSpectra object.
Accessors

In the following code snippets, x is a IBSpectra object.

proteinGroup(x): Gets and sets the ProteinGroup.

isotopeImpurities(x): Gets and sets the isotope impurities of the isobaric tags as defined by the manufacturers per batch.

reporterData(x, element="ions", na.rm=FALSE, na.rm.f='any', ...): Gets and sets the element ('ions' or 'mass') for each tag and spectrum. '...' is handed down to spectrumSel, so it is possible to select for peptides or proteins. If na.rm is TRUE, than spectra missing quantitative information in 'any' or 'all' channels (parameter na.rm.f) are removed.

reporterIntensities(x,...): Convenience function, calls reporterData(...,element="ions")

reporterMasses(x,...): Convenience function, calls reporterData(...,element="mass")

spectrumTitles(x,...): Gets the spectrum titles. '...' is passed down to spectrumSel.

classLabels(x): Gets and sets the class labels in phenoData. Used for summarization, see also estimateRatio and phenoData.

Methods

In the following code snippets, x is a IBSpectra object.

subsetIBSpectra(x, protein=NULL, peptide=NULL, direction="exclude", specificity): Get a 'subset' of IBSpectra: include or exclude proteins or peptides. When selection is based on proteins, it can be defined to exclude only peptides which are specific to the protein ('reporter-specific'), specific to the group ('group-specific') or which are shared with other proteins ('unspecific'). See subsetIBSpectra.

spectrumSel(x,peptide,protein,specificity="reporter-specific"): Gets a boolean vector selecting the corresponding spectra: If peptide is given, all spectra assigned to this peptide. If protein is given, all spectra assigned to peptides of this protein with specificity 'specificity'. See also ProteinGroup.

Author(s)

Florian P. Breitwieser

See Also

ProteinGroup, isobar-preprocessing, isobar-analysis, isobar-plots

Examples

data(ibspiked_set1)
ibspiked_set1
head(reporterIntensities(ibspiked_set1))
head(reporterMasses(ibspiked_set1))
proteinGroup(ibspiked_set1)
isotopeImpurities(ibspiked_set1)

# create new object
set.seed(123)
data <- data.frame(spectrum=letters,
peptide = sample(c("pepA", "pepB", "pepC"), 26, TRUE),
start.pos = 1,
modif = sample(c("::X:::", "::Y:::", "::Z:::"), 26, TRUE),
accession = c("protein1", "protein2")
data.ions <- matrix(rnorm(26*2, 1000, 50),
ncol = 2, dimnames = list(letters, NULL))
data.mass <- matrix(rep(c(126.1, 127.1), 26),
ncol = 2, byrow = TRUE, dimnames = list(letters, NULL))
ib <- new("TMT2plexSpectra", data, data.ions, data.mass)
ib
reporterIntensities(ib)
isotopeImpurities(ib) <- matrix(c(0.8, 0.1, 0.2, 0.9), nrow = 2)
reporterIntensities(correctIsotopeImpurities(ib))

---

**IBSpectra.log**

Log functions for IBSpectra objects

**Description**

The slot log of IBSpectra objects contains a matrix with two columns which contain a timestamp and message. Rownames relate to the item logged.

**Usage**

- `do.log(x, name, msg)`
- `get.log(x, name)`
- `is.logged(x, name)`

**Arguments**

- `x` IBSpectra object
- `name` Name of property to be logged (translates to row name).
- `msg` Message to be logged for name.

**Details**

A warning message will be displayed if a already logged property is logged again.

**Value**

- `do.log`: IBSpectra object with updated log. `get.log`:

**Author(s)**

Florian P Breitwieser

**See Also**

IBSpectra-class
Examples

data(ibspiked_set1)
ib <- normalize(correctIsotopeImpurities(ibspiked_set1))
ib@log

Isobar util functions  Isobar util functions

Description

Utility functions. paste0 as a shorthand to paste(...,sep="") in versions of R pre 2.14.

Usage

paste0(..., sep = "")
a %inrange% b

Arguments

...  Arguments to paste.
sep  Separator.
a  values.
b  range.

Author(s)

Florian P Breitwieser

Examples

1:10

IBSpectra analysis: Protein and peptide ratio calculation

Description

Calculates the relative abundance of a peptide or protein in one tag compared to another.
Usage

estimateRatio(ibspectra, noise.model = NULL, channel1, channel2, protein, peptide, ...)
estimateRatioForPeptide(peptide, ibspectra, noise.model, channel1, channel2, combine = TRUE, ...)
estimateRatioForProtein(protein, ibspectra, noise.model, channel1, channel2, combine = TRUE, method = "isobar", specificity = REPORTERSPECIFIC, quant.w.grouppeptides = NULL, ...)

## S4 method for signature 'numeric,numeric,missing'
estimateRatioNumeric(channel1,channel2,summarize.f=median, ...)

## S4 method for signature 'numeric,numeric,NoiseModel'
estimateRatioNumeric(channel1,channel2,noise.model,ratiodistr=NULL,variance.function="maxi",
    sign.level=0.05,sign.level.rat=sign.level,sign.l.
    remove.outliers=TRUE,outliers.args=list(method = "isobar",fc.threshold=1.3,
    channel1.raw=NULL,channel2.raw=NULL,use.na=FALSE)

## S4 method for signature 'IBSpectra,ANY,character,character,character,missing'
estimateRatio(ibspectra,noise.model,channel1,channel2,
    protein,peptide,...)

## S4 method for signature 'IBSpectra,ANY,character,character,character,NULL'
estimateRatio(ibspectra,noise.model,channel1,channel2,
    protein,peptide=NULL,...)

## S4 method for signature 'IBSpectra,ANY,character,character,missing,character'
estimateRatio(ibspectra,noise.model,channel1,channel2,protein,peptide,...)

## S4 method for signature 'IBSpectra,ANY,character,character,missing,character'
estimateRatio(ibspectra,noise.model,channel1,channel2,protein=NULL,peptide,...)

Arguments

ibspectra  IBSpectra object.
noise.model NoiseModel object.
channel1  Tag channel 1. Can either be a character denoting a 'reporter name' or a numeric vector whose value should be summarized. Ratio is calculated as channel2/channel1.
channel2  Tag channel 2. Can either be a character denoting a 'reporter name' or a numeric vector whose value should be summarized. Ratio is calculated as channel2/channel1.
protein  Protein(s) of interest. If present, channel1 and channel2 must be reporter names. Provide either proteins or peptides.
peptide  Peptide(s) of interest. If present, channel1 and channel2 must be reporter names. Provide either proteins or peptides.
combine  If true, a single ratio is returned even for multiple peptides/spectra. If false, a data.frame with a row for each peptide/protein is returned.
specificity  See specificities.
quant.w.grouppeptides  Proteins which should be quantified with group specific peptides. Normally, only reporter specific peptides are used.
ratiodistr distr object of ratio distribution.

variance.function

Defines how the variance for ratio is calculated. ’ev’ is the estimator variance and thus 1/sum(1/variances). ’wsv’ is the weighted sample variance. ’maxi’ method takes the maximum of the former two variances.

sign.level Significance level.

sign.level.rat Signal p-value significance level.

sign.level.sample Sample p-value significance level.

remove.outliers Should outliers be removed?

outliers.args Arguments for outlier removal, see OUTLIERS function (TODO).

method method taken for ratio computation and selection: one of ’isobar’, ’libra’, ’multiq’, ’pep’, ’ttest’ and ’compare.all’.

fc.threshold When method equals fc, takes this as fold change threshold.

summarize.f A method for summarizing spectrum ratios when no other information is available. For example median or mean.

channel1.raw When given, noise estimation is based on channel1.raw and channel2.raw. These are the intensities of the channels before normalization.

channel2.raw See channel1.raw.

use.na Use NA values to calculate ratio. Experimental feature - use with caution.

preweights Specifies weights for each spectrum. Experimental feature - use with caution.

Passed down to estimateRatioNumeric methods.

Value

In general, a named character vector with the following elements: - lratio: log ratio - variance - n.spectra: number of spectra available in the ratio calculation - p.value.rat: Signal p-value. NA if called w/o ratiodistr - p.value.sample: Sample p-value. NA if called w/o ratiodistr - is.significant: NA if called w/o ratiodistr

If combine=FALSE, estimateRatio returns a data.frame, with columns as described above.

Author(s)

Florian P. Breitwieser, Jacques Colinge

See Also

ProteinGroup, IBSpectra, isobar-preprocessing, isobar-plots proteinRatios

Examples

data(ibspiked_set1)
data(noise.model.hcd)
ceru.human <- protein.g(proteinGroup(ibspiked_set1),"CERU_HUMAN")
ceru.rat <- protein.g(proteinGroup(ibspiked_set1),"CERU_RAT")
ceru.mouse <- protein.g(proteinGroup(ibspiked_set1),"CERU_MOUSE")
ceru.proteins <- c(ceru.human,ceru.rat,ceru.mouse)
## Calculate ratio based on all spectra of peptides specific to CERU_HUMAN, CERU_RAT or CERU_MOUSE. Returns a named numeric vector.

```
10^estimateRatio(ibspiked_set1,noise.model.hcd,
    channel1="114",channel2="115",
    protein=ceru.proteins)['lratio']
```

## If argument 'combine=FALSE', estimateRatio returns a data.frame with one row per protein

```
10^estimateRatio(ibspiked_set1,noise.model.hcd,
    channel1="114",channel2="115",
    protein=ceru.proteins,combine=FALSE)['lratio']
```

```
## spiked material channel 115 vs 114:
## CERU_HUMAN (P00450): 1
## CERU_RAT (P13635): 2
## CERU_MOUSE (Q61147): 0.5
```
id.file Database search results file in ibspectra.csv or mzIdentML format. See identifications.format. See the vignette for information on converting Mascot dat and Phenyx pidres files into ibspectra format.

peaklist.file Peaklist file, typically in MGF format, see peaklist.format. MGF must be centroid!

mapping.file If defined, spectrum titles from the peaklist file are linked to the identifications via this file. This can be used when running HCD runs for quantification and CID runs for identification. See Koecher et al., 2009 for details.

mapping Named character vector defining the names of columns in mapping.file. The names must be 'peaklist' and 'id', and the values must correspond to colnames of the mapping files.

id.file.domap When using HCD-CID or a method akin and every spectrum is used for identification, the ID result files of the HCD run can be specified in id.file.domap. Then, the results are merged after mapping the identification results.

annotate.spectra.f Function which changes or annotates the spectra feature data before it is written to IBspectra object. This can be used to calculate and threshold additional scores, for example localization scores of post-translational modifications such as Delta Score (filterSpectraDeltaScore) or PhosphoRS site localization probabilities (annotateSpectraPhosphoRS).

peaklist.format "mgf" (Mascot Generic format) or "mcn" (iTracker Machine Readable output). When NULL, it detects the format on file name extension.

identifications.format "ibspectra.csv" or "mzid" (PSI MzIdentML format). When NULL, file format is guessed based on extension.

fragment.precision Fragment precision for extraction of reporter tags: for each tag and spectrum the m/z-intensity pair with it’s mass closest to the known reporter tag mass is extracted within the window true_mass +/- fragment.precision/2.

fragment.outlier.prob Fragment outlier probability filter: After all m/z-intensity pairs have been extracted, those pairs with the fragment.outlier.prob/2 most unprecise m/z values are filtered out.

decode.titles Boolean. Decode spectrum titles in identification file using URLdecode. When extracting the DAT file from Mascot web interface, the spectrum titles are encoded - %20 instead of space, etc. Set decode.titles to TRUE to map these titles to the unescaped MGF titles.

scan.lines Read files sequentially scan.lines lines at a time. Can help in case of memory issues, set to 10000 or higher, for example.

sep sep argument of read.table

... Further arguments handed down to initialize.

Author(s)

Florian P. Breitwieser, Jacques Colinge

See Also

ProteinGroup, IBSpectra, isobar-preprocessing, isobar-analysis, isobar-plots
Examples

data(ibspiked_set1)

# get identifier for Ceruplasmin proteins
ceru.acs <- protein.g(proteinGroup(ibspiked_set1),"CERU")
# create a smaller ibspectra w/ only Ceruplasmins
ib.ceru <- subsetIBSpectra(ibspiked_set1,protein=ceru.acs,direction="include")

# write it to a file
tf <- tempfile("isobar")
write.table(as.data.frame(ib.ceru),sep="\t",file=tf,quote=FALSE)

# read it again into an IBSpectra object
ib.ceru2 <- readIBSpectra("iTRAQ4plexSpectra",tf,identifications.format="ibspectra")
ib.ceru2

unlink(tf)

isobar-plots         IBSpectra plots

Description

Various plots are implement to assure data quality, and accompany preprocessing and analysis.

reporterMassPrecision

reporterMassPrecision(x): Calculates and displays the deviation from the 'true' tag mass - as specified in the IBSpectra object - of each channel.

reporterIntensityPlot

reporterIntensityPlot(x): Displays boxplots of intensity of channels before and after normalization - useful to check the result of normalization.

raplot

raplot(x,...): Ratio-Absolute intensity plot - will be deprecated by maplot
  x  IBSpectra object
  ... Parameters to plot function.

plotRatio

plotRatio(x,channel1,channel2,protein,...): Plots abundances of one protein
  x  IBSpectra object
  channel1
  channel2
  protein
  ... Parameters to plot function.
maplot

maplot(x, channel1, channel2, ...): Creates a ratio-versus-intensity plot.

Author(s)

Florian P. Breitwieser, Jacques Colinge

See Also

IBSpectra, isobar-preprocessing isobar-analysis

Examples

data(ibspiked_set1)
maplot(ibspiked_set1, main="IBSpiked, not normalized")
maplot(normalize(ibspiked_set1), main="IBSpiked, normalized")

isobar-preprocessing  IBSpectra preprocessing

description

Preprocessing is a necessary step prior to analysis of data. In a sequential order, it is often necessary to correct isotope impurities, to normalize, and subtract additive noise.

Isotope impurity correction

correctIsotopeImpurities(x): Returns impurity corrected IBSpectra object by solving a linear system of equations. See also isotopeImpurities.

Normalization

normalize(x, f=median, target="intensity", exclude.protein=NULL, use.protein=NULL, f.doapply=TRUE, log=TRUE, channels=NULL, na.rm=FALSE)

Normalizes the intensities for multiplicative errors. Those changes are most likely produced by pipetting errors, and different hybridization efficiencies, but can also be due to biological reasons. By default, tag intensities are multiplied by a factor so that the median intensity is equal across tags.

f: f is applied to each column, unless f.doapply is FALSE. Then f is supposed to compute column-wise statistics of the matrix of intensities. E.g. colSums and colMeans.

target: One of "intensity" and "ratio".

exclude.protein Spectra of peptides which might come from these proteins are excluded. Use for example for contaminants and proteins depleted in the experiment.

use.protein: If specified, only spectra coming from this protein are used. Use when a protein is spiked-in as normalization control.
f.isglobal: If true, f is applied on each column. If false, f is supposed to compute column-wise statistics of the matrix of intensities. E.g. colSums and colMeans.

log: Used when target=ratio.

Subtract additive noise

subtractAdditiveNoise(x, method="quantile", shared=TRUE, prob=0.01): method 'quantile' method is supported for now. It take’s the prob (0.01) quantile to estimate the noise level. This value is subtracted from all intensities, and all remaining intensities have to be at least that value.

prob See 'method'.

shared If channels are assumed similar in intensity and hence a shared noise level is reasonable. If not, then one level per channel is necessary.

Exclusion of proteins

exclude(x, proteins.to.exclude): Removes spectra which are assigned to proteins in protein.to.exclude from the object. This can be useful to remove contaminants. It create a new grouping based on the data which is left.

proteins.to.exclude Proteins to exclude.

Author(s)

Florian P. Breitwieser, Jacques Colinge

See Also

ProteinGroup, IBSpectra, isobar-analysis, isobar-plots

Examples

data(ibspiked_set1)
maplot(ibspiked_set1, main="IBSpiked, not normalized")
maplot(normalize(ibspiked_set1), main="IBSpiked, normalized")

isobar-reports  Isobar reports

Description

Generation of LaTeX and XLS reports is helped with functions which facilitate the gathering of relevant information and creation of tikz plots. create.reports parses properties (by calling load.properties) and initialize environments and computations (by calling initialize.env) required by the reports, calls Sweave and pdflatex.
Usage

create.reports(properties.file = "properties.R",
global.properties.file = system.file("report","properties.R", package = "isobar")
args = NULL, ..., recreate.properties.env = TRUE, recreate.report.env = TRUE)

load.properties(properties.file = "properties.R",
global.properties.file = system.file("report","properties.R",package="isobar")
args = NULL, ...)

initialize.env(env, properties.env)

Arguments

properties.file  
File which holds the parameters for data analysis and report generation. It is parsed as R code after the global report configuration file global.properties.file and defines peaklists, identification files, significance levels, etc. See the global properties file for the available options and values.
global.properties.file  
system.file("report","properties.R",package="isobar")
args  
Additional (command line) arguments which overrides those in properties.file.
...
recreate.properties.env  
Whether a properties.env existing in the global environment should be used, or it should be recreated.
recreate.report.env  
Whether a report.env existing in the global environment should be used, or it should be recreated.
env  
Item to be initialized.
properties.env  
Environment into which properties are read.

Details

The directory inst in the isobar installation directory system.file("inst",package="isobar") contains R, Sweave, and LaTeX files as examples of how to create XLS and PDF reports using isobar.

create_reports.R  Call with Rscript. It is the main file which
1. parses command line options. --compile and --zip are parsed directly and given as arguments to create.reports. Other arguments are given load.properties.
2. calls a perl script to generate a XLS report
3. generates a LaTeX quality control and analysis report for the XLS report the script pl/tab2xls.pl is used, which concatenates CSV files to a XLS. See Perl requirements. Sweave is called on report/isobar-qc.Rnw and report/isobar-analysis.Rnw. All files are written the working directory.
isobar-qc.Rnw  Quality control Sweave file.
isobar-analysis.Rnw  Data analysis Sweave file.
properties.R  Default configuration for data analysis.
report-utils.tex  LaTeX functions for plotting tikz graphics, etc.
isobar.data

Author(s)

Florian P Breitwieser

See Also

IBSpectra, isobar-preprocessing isobar-analysis

isobar.data  Isobar Data packages

Description

ibspiked_set1 and ibspiked_set2 are objects of class iTRAQ4plexSpectra. It contains over 160 protein groups, over 1600 peptides from about 15,000 spectra each, mainly from background proteins and three spiked-in Ceruplasmins (CERU_HUMAN, CERU_MOUSE, CERU_RAT).

Usage

data(ibspiked_set1)
data(ibspiked_set2)
data(ib_phospho)

Format

iTRAQ4plexSpectra objects.

Source

isobar publication. Acquired on Orbitrap instrument w/ 20 offline-fractions and HCD fragmentation.

Examples

data(ibspiked_set1)
print(ibspiked_set1)

maplot.protein  Ratio intensity plot for individual proteins

Description

Plots ratio-versus-intensity for a selected protein against a reference channel.

Usage

maplot.protein(x, relative.to, protein, noise.model = NULL, channels = NULL, xlim = NULL, ylim = NULL, identify = FALSE, add = FALSE, pchs = NULL, log="xy", legend.pos = "topright", names = NULL, legend.cex = 0.8, cols = pchs, lty = 1, main = protein, xlab = NULL, ylab = NULL, type="ma", show.lm = FALSE, ...)
Arguments

- x: IB Spectra object
- relative.to: a character vector specifying reporter tag names. Either of length 1 or same length as channels.
- protein: Protein group identifier.
- noise.model: NoiseModel object.
- channels: Reporter tag names.
- xlim: See par.
- ylim: See par.
- identify: boolean. If true, identify is called with peptide labels.
- add: a vector of the same length as channels. See pch in plot.default.
- log: a character string which contains x if the x axis is to be logarithmic, y if the y axis is to be logarithmic and xy or yx if both axes are to be logarithmic.
- legend.pos: see pos in legend.
- names: a character string of the same length as channels, legend text.
- legend.cex: see cex in legend.
- cols: a vector of the same length as channels. See col in plot.default.
- lty: a vector of the same length as channels. See lty in plot.default.
- main: a main title for the plot
- xlab: a label for the x axis, defaults to a description of x.
- ylab: a label for the y axis, defaults to a description of y.
- type: type of plot
- ...: passed to plot.
- show.lm: show LM

Author(s)

Florian P. Breitwieser

NoiseModel-class

Description

A NoiseModel represent the technical variation which is dependent on signal intensity.
NoiseModel-class

Constructor

new(type, ibspectra, reporterTagNames=NULL, one.to.one=TRUE, min.spectra=10, plot=FALSE, pool=FALSE):

Creates a new NoiseModel object based on ibspectra object.

type: A non-virtual class deriving from NoiseModel: ExponentialNoiseModel, ExponentialNoANoiseModel, InverseNoiseModel, InverseNoANoiseModel
reporterTagNames: When NULL, all channels from ibspectra are taken (i.e. sampleNames(ibspectra)). Otherwise, specify subset of names, or a matrix which defines the desired combination of channels (nrow=2).
one.to.one: Set to false to learn noise model one a non one-to-one dataset
min.spectra: When one.to.one=FALSE, only take proteins with min.spectra to learn noise model.
plot: Set to true to plot data the noise model is learnt on.
pool: If false, a NoiseModel is estimated on each combination of channels individually, and then the parameters are averaged. If true, the ratios of all channels are pooled and then a NoiseModel is estimated.

Accessor methods

noiseFunction: Gets the noise function.

parameters: Gets and sets the parameters for the noise function.

variance: Gets the variance for data points based on the noise function and parameters.

stddev: Convenience function, sqrt(variance(...)).

lowIntensity: Gets and sets the low intensity slot, denoting the noise region.

naRegion: Gets and sets the na.region slot.

Examples

data(ibsiked_set1)

 ceru.proteins <- protein.g(proteinGroup(ibsiked_set1),"CERU")

 # normalize
 ibspiked_set1 <- normalize(correctIsotopeImpurities(ibsiked_set1))

 # remove spiked proteins
 ibspiked_set1.noceru <- exclude(ibsiked_set1, ceru.proteins)
 ibspiked_set1.justceru <- subsetIBSpectra(ibsiked_set1, protein=ceru.proteins, direction="include")

 # learn noise models
 nm.i <- new("InverseNoiseModel", ibspiked_set1.noceru)
 nm.e <- new("ExponentialNoiseModel", ibspiked_set1.noceru)

 # learn on non-one.to.one data: not normalized, with spiked proteins
 nm.n <- new("ExponentialNoiseModel", ibspiked_set1.justceru, one.to.one=FALSE)

 maplot(ibsiked_set1, noise.model=c(nm.e, nm.i, nm.n), ylim=c(0.1, 10))
number.ranges  
*Helper function to transform number lists to ranges*

**Description**

1,2,3,4,5,8,9,10 -> 1-5,8-10

**Usage**

`number.ranges(numbers)`

**Arguments**

- `numbers` numeric

**Value**

character

**Author(s)**

Florian P Breitwieser

**Examples**

`number.ranges(c(1,2,3,9,3,10,8,11))`

observedKnownSites  
*Observed modification sites.*

**Description**

Functions to display the modification sites observed for each protein isoform and count the number of modified residues per protein.

**Usage**

`observedKnownSites(protein.group, protein.g, ptm.info, modif, modification.name = NULL)`

`modif.site.count(protein.group, protein.g = reporterProteins(protein.group), modif, take = max)`

`modif.sites(protein.group, protein.g = reporterProteins(protein.group), modif)`
peptide.count

Arguments

protein.group  ProteinGroupb object.
protein.g      protein group identifier.
ptm.info      ptm information data.frame, see ?getPtInfo.
modif         Modification to track, e.g. 'PHOS'.
modification.name  Value to filter 'modification.name' column in ptm.info.
take          should be either max or min: When multiple isoforms are present, which value should be taken for the count?

Author(s)

Florian P. Breitwieser

Examples

data(ib_phospho)
data(ptm.info)

# Modification sites of reporter proteins:
# a list of protein groups,
# containing sub-lists of identified sites for each isoform
protein.modif.sites <- sort(modif.site.count(proteinGroup(ib_phospho),modif="PHOS"))

# Details on modification sites of proteins
# detected with most modifications
modif.sites(proteinGroup(ib_phospho),modif="PHOS",protein.g=names(tail(protein.modif.sites)))

# How many sites are known, and how many known sites have been observed?
observedKnownSites(proteinGroup(ib_phospho),modif="PHOS",protein.g=names(tail(protein.modif.sites)),ptm.info=ptm.info)

peptide.count  Peptide counts, spectral counts and sequence coverage for ProteinGroup objects.

Description

Report the peptide count, spectral count and sequence coverage for supplied proteins.

Usage

peptide.count(protein.group, protein.g = reporterProteins(protein.group),
specificity = c("reporter-specific", "group-specific", "unspecific"), ...)
spectra.count(protein.group, protein.g = reporterProteins(protein.group),
specificity = c("reporter-specific", "group-specific", "unspecific"),
modif = NULL, ...)
sequence.coverage(protein.group, protein.g = reporterProteins(protein.group),
specificity = c("reporter-specific", "group-specific", "unspecific"),
simplify = TRUE, ...)
Protein and peptide ratio calculation and summarization

**Arguments**

- `protein.group` ProteinGroup object.
- `protein.g` Protein group identifier.
- `specificity` Specificity of peptides.
- `modif` Only count peptides having a certain modification.
- `simplify` If simplify=TRUE, a named numeric vector is returned, with the mean sequence coverage of the ACs of each `protein.g` supplied. Else, a list with the length of `protein.g` is returned having the sequence coverage for each protein AC.

... Further arguments to `peptides`

**Author(s)**

Florian P Breitwieser

**See Also**

`calculate.emPAI, calculate.dNSAF, ProteinGroup`

**Examples**

```r
data(ibspiked_set1)
sc <- spectra.count(proteinGroup(ibspiked_set1))
pc <- peptide.count(proteinGroup(ibspiked_set1))
plot(jitter(sc), jitter(pc), log="xy")
```

---

Protein and peptide ratio calculation and summarization

*Calculating and Summarizing Protein and Peptide Ratios*

**Description**

A set of functions to create ratios within groups and summarize them. `proteinRatios` serves as hub and calls `combn.matrix`, `combn.protein.tbl` and `summarize.ratios` successively. It can be used to calculate intra-class and inter-class ratios, to assess ratios and variability within and over cases.

**Usage**

```r
proteinRatios(ibspectra, noise.model, reporterTagNames = NULL,
proteins = reporterProteins(proteinGroup(ibspectra)),
peptide = NULL, cl = classLabels(ibspectra),
combn.method = "global", combn.vs = NULL,
symmetry = FALSE, summarize =
FALSE, summarize.method = "mult.pval", min.detect =
NULL, strict.sample.pval = TRUE, strict.ratio.pval =
TRUE, orient.div = 0, sign.level = 0.05,
sign.level.rat = sign.level, sign.level.sample =
sign.level, ratiodistr = NULL, zscore.threshold =
NULL, variance.function = "maxi", combine = FALSE,
p.adjust = NULL, reverse = FALSE, cmbn = NULL,
```

Protein and peptide ratio calculation and summarization

```r
before.summarize.f = NULL, ...)

peptideRatiosNotQuant(ibspectra, ..., peptide = unique(fData(ibspectra)[!fData(ibspectra)["use.for.quant"]])
peptideRatios(ibspectra, ..., peptide = peptides(proteinGroup(ibspectra), columns = c("peptide", "modif")))

combn.matrix(x, method = "global", cl = NULL, vs = NULL)
combn.protein.tbl(cmbn, reverse = FALSE, ...)

summarize.ratios(ratios, by.column = "ac", summarize.method = "mult.pval",
min.detect = NULL, n.combination = NULL,
strict.sample.pval = TRUE, strict.ratio.pval = TRUE,
orient.div = 0, sign.level = 0.05, sign.level.rat =
sign.level, sign.level.sample = sign.level,
variance.function = "maxi", ratiodistr = NULL)
```

**Arguments**

- **ibspectra**
  - IBSpectra object
- **x**
  - for combinatn matrix: reporter names. See reporterTagNames. argument of proteinRatios.
- **ratios**
  - result of combinatn.protein.tbl
- **by.column**
  - Column(s) which are the identifiers. Usually 'ac', 'peptide' or c('peptide','modif')
- **cmbH**
  - result of combinatn.matrix
- **before.summarize.f**
  - Function which is called after calculating ratios before summarizing them.
- **noise.model**
  - NoiseModel for spectra variances
- **reporterTagNames**
  - Reporter tags to use. By default all reporterTagNames of ibspectra object.
- **proteins**
  - proteins for which ratios are calculated - defaults to all proteins with peptides specific to them.
- **peptide**
  - peptides for which ratios are calculated.
- **cl**
  - Class labels. See also ?classLabels.
- **vs**
  - Class label or reporter tag name. When combinatn.method is "versus.class", all combinations against class vs are computed, when combinatn.method is "versus.channel", all combinations against channel vs.
- **combinatn.method**
  - "global", "interclass", "intra-class", "versus.class" or "versus.channel". Defines which ratios are computed, based on class labels cl
- **method**
  - See combinatn.method
- **combinatn.vs**
  - vs argument for combinatn, if combinatn.method is "versus.class" or "versus.channel".
- **symmetry**
  - If true, reports also the inverse ratio
- **summarize**
  - If true, ratios for each protein are summarized.
- **summarize.method**
  - "isobar", for now.
- **min.detect**
  - How many times must a ratio for a protein be present when summarizing? When NULL, defaults to the maximum number of combinations.
 Protein and peptide ratio calculation and summarization

strict.sample.pval
If true, missing ratios are penalized by giving them a sample.pval of 0.5.

strict.ratio.pval
If true, take all ratios into account. If false, only take ratios into account which are in the same direction as the majority of ratios.

orient.div
Number of ratios which might go in the wrong direction.

sign.level
Significance level

sign.level.rat
Significance level on ratio p-value

sign.level.sample
Significance level on sample p-value

ratiodistr
Protein ratio distribution

variance.function
Variance function

zscore.threshold
z-score threshold to apply

... Passed to estimateRatio()

combine
If true, a single ratio for all proteins and peptides, resp., is calculated. See estimateRatio.

p.adjust
Set to one of p.adjust.methods to adjust ratio p-values for multiple comparisions. See p.adjust.

reverse
reverse

n.combination
number of combinations possible

Value
'data.frame': 11 variables:

lratio log ratio
variance variance
n.spectra Number of spectra used for quantification
p.value.rat Signal p-value (NA if ratiodistr is missing)

p.value.sample
Sample p-value (NA if ratiodistr is missing)

is.significant
Is the ratio significant? (NA if ratiodistr is missing)

protein Protein quantified
r1 r1
r2 r2

Author(s)
Florian P Breitwieser, Jacques Colinge

See Also
IBSpectra, isobar-preprocessing isobar-analysis
ProteinGroup-class

Examples

```r
combn.matrix(114:117, method="interclass", cl=as.character(c(1,1,2,2)))
combn.matrix(114:117, method="interclass", cl=as.character(c(1,1,2,2)))
combn.matrix(114:117, method="global")

data(ibsptiked_set1)
data(noise.model.hcd)

ceru.proteins <- c("P13635","Q61147")
proteinRatios(ibsptiked_set1,noise.model=noise.model.hcd,proteins=ceru.proteins,cl=c("T","T","C","C"),combn.method="interclass",summarize=TRUE)
```

---

ProteinGroup-class  ProteinGroup objects

Description

The ProteinGroup class is a container for identified peptides and proteins, and groups them to distinguish proteins with specific peptides.

Usage

```r
ProteinGroup(from, template=NULL, proteinInfo=data.frame())

protein.ac(x, protein.g)
protein.g(x, pattern, variables=c("AC","name"), ...)
```

Arguments

- **from**: data.frame object to create a ProteinGroup from. See Details from column specifications
- **template**: 'template' ProteinGroup object for grouping.
- **x**: ProteinGroup object
- **protein**: character string
- **proteinInfo**: data.frame for proteinInfo slot
- **protein.g**: character string, denoting a 'protein group'.
- **pattern**: character string, see `grep` for details.
- **variables**: `AC` maps a protein accession code to a protein group. `name` maps using protein information from proteinInfo.
- **...**: Passed on to `grep`.

Details

The ProteinGroup class stores spectrum to peptide to protein mapping.

The proteins are grouped by their evidence, i.e. peptides:

- Peptides with changes only from Leucin to Isoleucin are considered the same, as they cannot be distinguished by MS.
ProteinGroup-class

- Proteins which are detected with the same peptides are grouped together to a 'indistinguishable protein'- normally these are splice variants.
- Proteins with specific peptides are 'reporters'.
- Proteins with no specific peptides are grouped under these 'reporters'.

This information is stored in six slots:

- `spectra.n.peptides` a named 'character' vector, names being spectrum identifier and values are peptides.
- `peptide.n.proteins` a 'data.frame' containing the number of proteins the peptides could derive from.
- `peptide.n.protein` a character 'matrix' linking peptides to proteins.
- `indistinguishable.proteins` a 'matrix' contain.

Constructor

```r
ProteinGroup(tbl.prot.pep, template=NULL): Creates a ProteinGroup object.
```

- `template` Optional ProteinGroup object the grouping is based upon.

Coercion

In the code snippets below, `x` is a ProteinGroup object.

```r
as(from, "ProteinGroup"/business): Creates a ProteinGroup object from a data.frame.
```

```r
as.data.frame(x, row.names = NULL, optional = FALSE): Creates a data.frame with columns protein (character), peptide (character), spectrum.
```

```r
as.concise.data.frame(from): Creates a 'concise' data.frame with one spectrum per row, and protein ACs combined
```

Accessors

In the following code snippets, `x` is a ProteinGroup object.

```r
spectrumToPeptide(x): Gets spectrum to peptide assignment.
```

```r
peptideInfo(x): Peptide information such as protein start position.
```

```r
peptideSpecificity(x): Gets a 'data.frame' containing the peptide specificity: they can be reporter-specific, group-specific, or non-specific.
```

```r
peptideNProtein(x): Gets peptide to protein assignment.
```

```r
indistinguishableProteins(x): Gets the proteins which cannot be distinguished based on peptide evidence.
```

```r
proteinGroupTable: Gets the protein grouping, listing reporters and group members.
```

```r
peptides(x, protein=NULL, specificity=c("reporter-specific", "group-specific", "unspecific"), columns=character): Gets all peptides detected, or just those for a protein with the defined specificity. columns might define multiple columns of peptideSpecificity(x). set=union returns the union of peptides of all proteins defined, set=intersect returns the intersection.
```

Author(s)

Florian P. Breitwieser
proteinInfo-methods

See Also

IBSpectra

Examples

tbl <- data.frame(spectrum=1:14,peptide=c(rep(letters[1:3],4),"a","x"),
    modif="",start.pos=1,
    protein=c(rep(c("A","B"),each=6),"C","D"))
pg <- ProteinGroup(tbl)
pg
proteinGroupTable(pg)

data(ibspiked_set1)
pg <- proteinGroup(ibspiked_set1)
ceru.proteins <- protein.g(pg,"CERU")

## all ceru peptides
peptides(pg,ceru.proteins)

## peptides shared by all ceru proteins
peptides(pg,ceru.proteins, set=intersect)

proteinInfo-methods  Methods for Function proteinInfo

Description

proteinInfo slot in Proteingroup objects contains information about proteins. proteinInfo method allows to get and set it.

getProteinInfoFromUniprot downloads information of contained proteins from Uniprot, getProteinInfoFromBiomart from Biomart.

Usage

## S4 method for signature 'ProteinGroup'
proteinInfo(x)

## S4 method for signature 'ProteinGroup,character,missing'
proteinInfo(x, protein.g, select="name", collapse="",
    simplify = TRUE, do.warn = TRUE)

## S4 method for signature 'ProteinGroup,missing,character'
proteinInfo(x, protein.ac, select="name", collapse="",
    simplify = TRUE, do.warn = TRUE)

proteinInfoIsOnSpliceVariants(protein.info)

# getProteinInfoFromUniprot(x, splice.by = 200, fields = c(accession = "id", name
proteinInfo-methods

proteinInfo = "entry%20name", protein_name = "protein%20names",
# gene_name = "genes", organism = "organism", length =
# "length", sequence = "sequence")

getProteinInfoFromTheInternet(x)
getProteinInfoFromNextProt(x)
getProteinInfoFromBiomart(x, database = "Uniprot")
getProteinInfoFromBioDb(x, ..., con = NULL)
getProteinInfoFromEntrez(x, splice.by = 200)

Arguments

x ProteinGroup object
protein.g Protein group identifier. If supplied, only information for these proteins is returned.
protein.ac Protein ACs. If supplied, only information for these proteins is returned.
select indicating columns to select. See Details.
collapse passed to paste to concatenate information of multiple protein in one protein group.
simplify If true, a vector or matrix is returned, with the pasted protein information. If false, a list is returned.
do.warn If true, report diagnostic warning messages.
splice.by Chunk size for query of Uniprot database.
database database from which the ACs stem from. Only Uniprot is supported for now.
con database connection
fields mapping of CSV field names to proteinInfo field names
...
protein.info protein info data.frame

Details

proteinInfo contains columns accession, name, gene_name, protein_name, and possibly length and sequence. accession is mapped with the entry AC is mapped to the entry AC in the database. getProteinInfoFromUniprot is the preferred methods to get the information. getProteinInfoFromBioDb is an example how to implement the query on a local database. Depending on the database, protein information might be available on protein ACs or also on the specific splice variants. This can be queried with the proteinInfoIsOnSpliceVariants function.

See Also

protein.g
Examples

```r
data(ibspiked_set1)
pg <- proteinGroup(ibspiked_set1)

## Not run:
proteinInfo(pg) <- getProteinInfoFromUniprot(pg)
proteinInfo(pg) <- getProteinInfoFromBiomart(pg)

## End(Not run)

proteinInfo(pg,protein.g="P13635")
protein.g(pg,"CERU")
```

---

**proteinNameAndDescription**

*Get protein gene names and description from protein info of protein group.*

Description

Convenience functions to retrieve protein gene names and description for a list of protein group identifiers.

Usage

```r
proteinNameAndDescription(protein.group, protein.g = reporterProteins(protein.group), collapse = FALSE)
proteinGeneName(protein.group, protein.g = reporterProteins(protein.group))
proteinDescription(protein.group, protein.g = reporterProteins(protein.group))
proteinID(protein.group, protein.g = reporterProteins(protein.group))
```

Arguments

- `protein.group`: ProteinGroup object.
- `protein.g`: protein group identifier.
- `collapse`: If TRUE, the information for all protein.gs is combined.

Author(s)

Florian P Breitwieser

Examples

```r
data(ibspiked_set1)
pkg <- proteinGroup(ibspiked_set1)
protein.gs <- protein.g(pkg,"CERU")
protein.gs
proteinNameAndDescription(pkg,protein.gs)
proteinNameAndDescription(pkg,protein.gs,collapse=TRUE)
proteinGeneName(pkg,protein.gs)
proteinDescription(pkg,protein.gs)
proteinID(pkg,protein.gs)
```
**ratiosReshapeWide**  
*Reshape output of proteinRatios into wide format*

### Description

Reshape output of proteinRatios into wide format

### Usage

```r
ratiosReshapeWide(quant.tbl, vs.class = NULL, sep = ".", cmbn = NULL,
short.names = FALSE)
```

### Arguments

- `quant.tbl`: Output of proteinRatios or peptideRatios.
- `vs.class`: Only return ratios where class1 is vs.class
- `sep`: Separator for column names in the reshape.
- `cmbn`: Not functional.
- `short.names`: If vs.class is set and short.names=TRUE, then the comparison name will be i.e. `class2` instead of `class2/class1`.

### Author(s)

Florian P. Breitwieser

---

**reporter.protein-methods**

*Get reporter protein group identifier for protein group identifier*

### Description

Methods for function `reporter.protein` in package `isobar`

### Methods

```r
signature(x = "ProteinGroup", protein.g = "character") Get reporter protein for protein group identifier.
```
sanitize

*Helper function for LaTeX export*

**Description**
Sanitizes strings for LaTeX

**Usage**
sanitize(str, dash = TRUE)

**Arguments**
- **str**: character string to be escaped
- **dash**: should a dash ('-') should be escaped to a `\nobreakdash-`?

**Value**
escaped character

**Author(s)**
iQuantitator, Florian P Breitwieser

**Examples**
sanitize("\textbf{123-123}")

---

**shared.ratios**

*Shared ratio calculation*

**Description**
Calculate ratios of reporter proteins and subset proteins with shared peptides.

**Usage**
shared.ratios(ibspectra, noise.model, channel1, channel2, protein = reporterProteins(proteinGroup(ibspectra)), ...)

**Arguments**
- **ibspectra**: IBspectra object.
- **noise.model**: NoiseModel object.
- **channel1**: channel1 to compare.
- **channel2**: channel2 to compare.
- **protein**: proteins for which the calculation should be made.
- **...**: Additional arguments passed to estimteRatio.
specificities

Value
data.frame

Author(s)
Florian P. Breitwieser

See Also
shared.ratios.sign

shared.ratios.sign  Plot and get significantly shared ratios.

Description
Plot and get significantly shared ratios.

Usage
shared.ratios.sign(ress, z.shared, min.spectra = 1, plot = TRUE)

Arguments
ress  Result of shared.ratios.
z.shared  z.
min.spectra  Minimal number of spectra needed.
plot  plot.

Author(s)
Florian P. Breitwieser

See Also
shared.ratios.

specificities  Peptide specificities

Description
Peptides can appear in multiple proteins and therefore have different specificities.

Details
reporter specific: peptides specific to reporter. group specific: peptides specific to the group. un-specific: peptides shared with other proteins.
spectra.count2

Spectral count for peptides and proteins in ProteinGroup objects.

Description
Spectral count for peptides and proteins in ProteinGroup objects. It can - other than spectra.count - quantify the spectra count on the level of peptides, potentially modified, too.

Usage
spectra.count2(ibspectra, value = reporterProteins(protein.group),
              type = "protein.g", specificity = c("reporter-specific", "group-specific", "unspecific"),
              modif = NULL, combine = FALSE, subset = NULL, require.quant = NULL, ...)

Arguments
ibspectra
  IBSpectra object.
value
  List of protein group identifiers or peptides.
type
  Either 'protein.g' or 'peptide'.
specificity
  Specificity of peptides.
modif
  Only count peptides having a certain modification.
combine
  If TRUE, only one combined result is returned.
subset
  Allows to specify an expression to subset link(featureData) of the ibspectra.
require.quant
  If not NULL, it may be 'any' or 'all' to only consider spectra with quantitative information in at least one or all channels.
...
  Further arguments to peptides

Author(s)
Florian P Breitwieser

See Also
spectra.count, ProteinGroup

Examples
data(ibspiked_set1)
pg <- proteinGroup(ibspiked_set1)
protein.gs <- protein.g(pg,"CERU")
sc <- spectra.count2(ibspiked_set1,protein.gs)
sc.ik <- spectra.count2(ibspiked_set1,protein.gs,modif="iTRAQ4plex.K")
rbind(spectra.counts=sc,spectra.counts_iTRAQk=sc.ik)
subsetIBSpectra  Subset IBSpectra objects

Description

Returns an IBSpectra object which is a subset of the input, excluding or exclusively containing the peptides or proteins supplied.

Usage

subsetIBSpectra(x, protein = NULL, peptide = NULL, direction = "exclude", specificity = c(REPORTERSPECIFIC, GROUPSPECIFIC, UNSPECIFIC), ...)

Arguments

x  IBSpectra object.
protein  Protein group identifiers. Use protein.g to get protein group identifiers from protein database ACs.
peptide  Peptide sequences.
direction  either 'include' or 'exclude'.
specificity  When 'protein' is supplied: Which peptides should be selected? See specificities.
...  Further arguments passed to spectrumSel

Author(s)

Florian P Breitwieser

See Also

protein.g, spectrumSel, specificities

Examples

data(ibspiked_set1)

# get Keratin proteins
keratin.proteins <- protein.g(proteinGroup(ibspiked_set1),"Keratin")

# exclude Keratin proteins
subsetIBSpectra(ibspiked_set1,protein=keratin.proteins,direction="exclude")
Description

Location scale family T distribution, based on the original T function.

Objects from the Class

Objects can be created by calls of the form new("Tlsd", df, location, scale).

Slots

gaps: Object of class "OptionalMatrix" ~
img: Object of class "rSpace" ~
param: Object of class "OptionalParameter" ~
r: Object of class "function" ~
d: Object of class "OptionalFunction" ~
p: Object of class "OptionalFunction" ~
q: Object of class "OptionalFunction" ~
.withSim: Object of class "logical" ~
.withArith: Object of class "logical" ~
.logExact: Object of class "logical" ~
.lowerExact: Object of class "logical" ~
Symmetry: Object of class "DistributionSymmetry" ~

Extends


Methods

No methods defined with class "Tlsd" in the signature.

Author(s)

Florian P. Breitwieser, based on original T distribution class.

Examples

showClass("Tlsd")
TlsParameter-class  

Class "TlsParameter"

Description

The parameter of a location scale t distribution, used by Tlsd-class

Objects from the Class

Objects can be created by calls of the form new("TlsParameter", ...). Usually an object of this class is not needed on its own, it is generated automatically when an object of the class Tlsd is instantiated.

Slots

df: Object of class "numeric" ~~
location: Object of class "numeric" ~~
scale: Object of class "numeric" ~~
name: Object of class "character" ~~

Extends


Methods

No methods defined with class "TlsParameter" in the signature.

Author(s)

Florian P. Breitwieser, based on original TParameter class.

See Also

Tlsd

Examples

showClass("TlsParameter")
writeHscoreData

Write identifications into a format suitable for Hscore.

Description
Write identifications into a format suitable for Hscore.

Usage
writeHscoreData(outfile, ids, massfile = "defs.txt")

Arguments
outfile Output file.
ids IBSpectra identifications data.frame (ie fData).
massfile Definition file for Hscore.

Author(s)
Florian P. Breitwieser

writeIBSpectra
Write IBSpectra file as CSV in a format readable by readIBSpectra.

Description
Write IBSpectra file using write.table with defaults in a format readable by readIBSpectra.

Usage
writeIBSpectra(ibspectra, file, sep = "\t", row.names = FALSE, ...)

Arguments
ibspectra IBSpectra object
file file name.
sep field separator string.
row.names indicates whether row.names should be written.
... further arguments to write.table

Author(s)
Florian P Breitwieser
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