isobar for quantification of PTM datasets

Florian P. Breitwieser, Jacques Colinge

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1 Introduction

isobar [1] version 2 includes modules to facilitate PTM quantification. This vignette describes its parts, and how to use it to generate quantification reports.

> library(isobar) ## load the isobar package

Using isobar, automatic report generation is straight-forward given proper input files using the script report/create_reports.R. When called, it parses the globabl properties file report/properties.R and then the properties.R in the current directory. Below is a small example properties.R for creating a PDF Quality Control and XLSX analysis report:

```
type="iTRAQ4plexSpectra"
## peaklist files for quantitation, by default all mgf file in directory
peaklist=list.files(pattern="*\\.mgf$")

## id files, by default all id.csv files in directory
identifications=list.files(pattern="*\\.id.csv$")

modif="PHOS" # modification to track (eg PHOS, ACET, MET)
ptm.info.f <- getPtmInfoFromNextprot
spreadsheet.format="xlsx"</pre>
```

Reports will be generated calling path_to_isobar/report/create_reports.R -peptide from the directory containing the peaklists, identifications and properties.R.

2 Modification Site Localization

isobar supports PhosphoRS [5] and Delta Score [4] for modification site localization.

PhosphoRS integration The standalone Java version of PhosphoRS can be downloaded from http://cores.imp.ac.at/uploads/media/PhosphoRS.zip. It features a command line interface to a script which rescores localizations of the modification for each peptide-spectrum match. It uses XML files for input and output, which can be generated and parsed by isobar.

After calling PhosphoRS (java -jar phosphoRS.jar phosphors.in.xml phosphors.out.xml), the resulting XML file can be read:

```
> # Read PhosphoRS XML output file
> # simplify: if TRUE, a data.frame is returned, else a list
> # besthit.only: if TRUE, only the best localization per spectrum is returned
> readPhosphoRSOutput("phosphors.out.xml",simplify=TRUE,besthit.only=TRUE)
```

getPhosphoRSProbabilities is a convenience function calling the writer, the script, and the reader in succession.

```
> getPhosphoRSProbabilities("identifications.id.csv", "peaklist.mgf",
+ massTolerance=0.5,activationType="CID",
+ phosphors.cmd="java -jar phosphoRS.jar")
```

Delta Score calculation The Mascot Delta Score can be calculated directly by the parser mascotParser2.pl and thresholded (e. g. -minDeltaScore=10). For CSV identification files which contain all hits for each spectrum (not just the best one), the function calc.delta.score within the R package is provided.

Using PhosphoRS and Delta Score in Report Generation. When generating an IB-Spectra object from peaklist and identifications, via readIBSpectra's argument annotate.spectra.f a function can be plugged in to extend or modify the identification information. This can be used to calculate scores and filter localization scores with filterSpectraDeltaScore) or annotateSpectraPhosphoRS.

```
> # filterSpectraDeltaScore calls calc.delta.score
> # if no column named delta.score is present in the data frame
> # identifications below a min.delta.score are REMOVED
> ib <- readIBSpectra("identifications.id.csv", "peaklist.mgf",</pre>
```

This can be used in report generation, too, where the readIBSpectra.args can be set accordingly in the report properties file properties.R:

```
readIBSpectra.args = list(annotate.spectra.f=filterSpectraDeltaScore)
or
readIBSpectra.args = list(annotate.spectra.f=filterSpectraPhosphoRS)
```

3 Peptide Ratio Calculation

> data(ib_phospho)

All functions which are available to calculate ratios on protein level can also be used for peptides. The same noise model is appropriate for both.

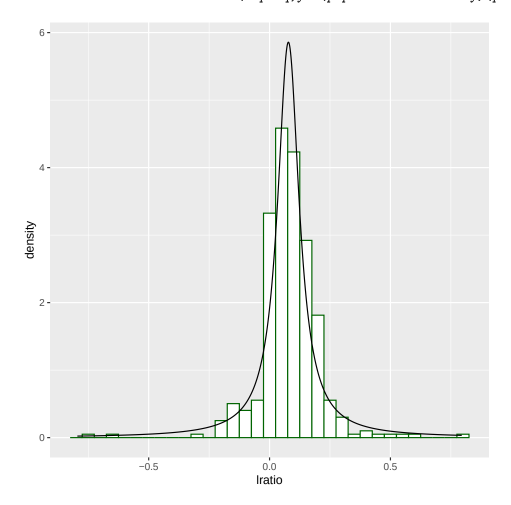
```
> data(noise.model.hcd)
> head(proteinGroup(ib_phospho)@peptideInfo)
     protein
                           peptide start.pos
2072 A1L390-1
                   SPLSPTETFSWPDVR
                                        1037
2074 A1L390-2
                   SPLSPTETFSWPDVR
                                         570
2076 A1L390-3
                   SPLSPTETFSWPDVR
                                         981
1299
                     LLLDLPLQTPHK
                                        1170
      A6NKT7
783
      000264 GDQPAASGDSDDDEPPPLPR
                                          48
2045 014497-1
                         SPFLHSGMK
                                        1604
                                             modif
2072
          iTRAQ4plex_Nterm:PHOS:::PHOS:::::::::
2074
          iTRAQ4plex_Nterm:PHOS:::PHOS:::::::::
2076
          iTRAQ4plex_Nterm:PHOS:::PHOS:::::::::
1299 iTRAQ4plex_Nterm:::::::PHOS:::iTRAQ4plex_K:
783
         iTRAQ4plex_Nterm::::::PHOS:::::::::::
2045
        iTRAQ4plex_Nterm:::::PHOS:::iTRAQ4plex_K:
```

> 10^estimateRatio(ib_phospho,noise.model.hcd,peptide="SPLSPTETFSWPDVR")

```
115
                              116
114 1.0000000 0.3088721 1.4354943 1.641885
115 3.2375859 1.0000000 4.6497966 5.318776
116 0.6966241 0.2150632 1.0000000 1.143867
117 0.6090561 0.1880132 0.8742276 1.000000
   By giving a matrix to estimateRatio, we can calculate ratios for peptides with specific
modifications:
> pep.n.modif <- unique(apply(fData(ib_phospho)[,c("peptide","modif")],2,cbind))
> print(head(pep.n.modif))
     peptide
[1,] "AAATPESQEPQAK"
[2,] "AAEAGGAEEQYGFLTTPTK"
[3,] "AAEEQGDDQDSEK"
[4,] "AAPPPGSPAK"
[5,] "AAVGQESPGGLEAGNAK"
[6,] "AAVLSDSEDEEK"
    modif
[1,] "iTRAQ4plex_Nterm::::::PHOS:::::iTRAQ4plex_K:"
[2,] "iTRAQ4plex_Nterm:::::::PHOS::::::iTRAQ4plex_K:"
[3,] "iTRAQ4plex_Nterm::::::::::::::::::::::::::TRAQ4plex_K:"
[4,] "iTRAQ4plex_Nterm::::::PHOS:::iTRAQ4plex_K:"
[5,] "iTRAQ4plex_Nterm::::::PHOS:::::::iTRAQ4plex_K:"
[6,] "iTRAQ4plex_Nterm:::::PHOS::PHOS::::iTRAQ4plex_K:"
> estimateRatio(ib_phospho, noise.model.hcd, channel1="114", channel2="115",
                  peptide=head(pep.n.modif), combine=FALSE) [, c("lratio", "variance",
                                                                "n.spectra", "p.value.rat")]
+
      lratio
               variance n.spectra p.value.rat
1 -0.6978020 0.01034090
                                 2 3.394310e-12
         {\tt NaN}
3 0.1388425 0.01052788
                                2 8.800084e-02
4 -1.0793665 0.04166971
                                1 6.196545e-08
5 -0.9655771 0.02406589
                                1 2.419553e-10
6 -0.2164083 0.08305527
                                 7 2.263522e-01
   A ratio distribution can be calculated based on peptide ratios:
> suppressPackageStartupMessages(library(distr))
> suppressPackageStartupMessages(library(ggplot2))
> peptide.ratios <- peptideRatios(ib_phospho,noise.model=noise.model.hcd,
```

117

114



Correction with protein ratios. The observed change in concentration of modified peptides in one condition versus another is integrating two separate modes of regulation [6]:

- 1. Protein expression change
- 2. Modification state change

In many cases, it thus can be advisable to conduct separate MS quantification runs of the peptides enriched for the modification of interest, AND the global proteome quantification.

In the report generation, data from other experiments can be integrated using the property compare.to.quant in properties.R:

```
load("../proteome/quant.tbl.rda")  # load proteome quantification table
compare.to.quant=list(proteome=quant.tbl) # set property
rm(quant.tbl)
```

Peptide ratios can also be corrected with proteome ratios of a separate experiment, when giving as **peptide** argument a **matrix** or **data.frame** with columns for 'peptide', 'modif', and 'correct.ratio'. 'correct.ratio' is a log_{10} ratio which will be used to adjust the one calculated for the specific modified peptide.

```
> peptides <- pep.n.modif[1:5,]</pre>
> orig.ratio <- estimateRatio(ib_phospho,noise.model.hcd,channel1="114",channel2="115",
                               peptide=peptides,combine=FALSE)[,c("lratio","variance")]
> peptides.c <- cbind(peptides,correct.ratio=c(0,-1,1,2,-2))
> corr.ratio <- estimateRatio(ib_phospho,noise.model.hcd,channel1="114",channel2="115",
                               peptide=peptides.c,combine=FALSE)[,c("lratio","variance")]
> data.frame(peptides.c, orig.ratio, corr.ratio)
             peptide
                                                                   modif
1
       AAATPESQEPQAK
                           iTRAQ4plex_Nterm::::::PHOS:::::iTRAQ4plex_K:
2 AAEAGGAEEQYGFLTTPTK iTRAQ4plex_Nterm:::::::::::::::::::::::TRAQ4plex_K:
       AAEEQGDDQDSEK
                           3
                              iTRAQ4plex_Nterm::::::PHOS:::iTRAQ4plex_K:
4
          AAPPPGSPAK
5
   AAVGQESPGGLEAGNAK
                       iTRAQ4plex_Nterm::::::PHOS:::::::iTRAQ4plex_K:
  correct.ratio
                   lratio
                            variance
                                      lratio.1 variance.1
1
             0 -0.6978020 0.01034090 -0.3736099 0.1962761
                                 Inf -0.3736099 0.1962761
2
            -1
                      NaN
3
             1 0.1388425 0.01052788 -0.3736099
                                                0.1962761
             2 -1.0793665 0.04166971 -0.3736099
4
                                                0.1962761
            -2 -0.9655771 0.02406589 -0.3736099 0.1962761
```

As appearent, the variance stays the same also for corrected ratios. If a fourth column variance of the peptide argument reports the variance of the correction ratio, it is added to the calculated ratio's variance (assuming independence).

4 Harvesting public PTM databases

neXtProt [3] and PhosphoSitePlus [2] provide information on experimentally determined post-translational modifications. neXtProt focuses on man, and PhosphoSitePlus on man and mouse. Both are manually curated and annotate thousands of residues of post-translationally modified proteins.

isobar provides functions to gather their information on identified proteins.

```
> ptm.info <- getPtmInfoFromPhosphoSitePlus(proteinGroup(ib_phospho),modif="PHOS")
> ptm.info <- getPtmInfoFromNextprot(proteinGroup(ib_phospho))</pre>
```

> head(ptm.info)

	.id	isoform_ac	quality	description	evidence	first_position	last_position
1	A1L390	A1L390-1	GOLD	Phosphoserine	EXP	76	76
2	A1L390	A1L390-1	SILVER	Phosphoserine	EXP	433	433
3	A1L390	A1L390-1	GOLD	Phosphoserine	Curated	533	533
4	A1L390	A1L390-1	GOLD	Phosphoserine	EXP	576	576
5	A1L390	A1L390-1	GOLD	Phosphoserine	EXP	577	577
6	A1L390	A1L390-1	SILVER	Phosphoserine	EXP	614	614
	modific	cation.name	modifica	ation.accession	position	ı	
1	Phosphoserine			PTM-0253	76	3	
2	Phosphoserine			PTM-0253	433	3	
3	Phosphoserine		PTM-0253	533	3		
4	Phosphoserine		PTM-0253	576	3		
5	Phosphoserine		PTM-0253	577	7		
6	Phosphoserine		PTM-0253	614	<u>l</u>		

For reports, the function can be selected via the property ptm.info.f in properties.R:

protein.info.f = getPtmInfoFromNextprot

For PhosphoSitePlus, define the modification to get the correct dataset:

```
ptm.info.f <- function(...) getPtmInfoFromPhosphoSitePlus(...,modification="PHOS")</pre>
```

PhosphoSitePlus datasets will be downloaded from their website to 'Phosphorylation_site_dataset.gz' or 'Acetylation_site_dataset.gz', etc (see mapping property of getPtmInfoFromPhosphoSitePlus) unless a file with that name exists.

References

- [1] F. P. Breitwieser, A. Müller, L. Dayon, T. Köcher, A. Hainard, P. Pichler, U. Schmidt-Erfurth, G. Superti-Furga, J.-C. Sanchez, K. Mechtler, K. L. Bennett, and J. Colinge. General statistical modeling of data from protein relative expression isobaric tags. *J Proteome Res*, 10(6):2758–2766, Jun 2011.
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- [4] M. M. Savitski, S. Lemeer, M. Boesche, M. Lang, T. Mathieson, M. Bantscheff, and B. Kuster. Confident phosphorylation site localization using the mascot delta score. *Mol Cell Proteomics*, 10(2):M110.003830, Feb 2011.

- [5] T. Taus, T. Köcher, P. Pichler, C. Paschke, A. Schmidt, C. Henrich, and K. Mechtler. Universal and confident phosphorylation site localization using phosphors. J Proteome Res, Nov 2011.
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