isobar package for iTRAQ and TMT protein quantification

Florian P. Breitwieser, Jacques Colinge

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1 Introduction

The isobar package is an extensible and interactive environment for the data analysis and exploration of iTRAQ and TMT data. isobar implements the theory presented in Breitwieser et al., Journal of Proteome Research 2011. Further extensions for the analysis of PTM data are presented in Breitwieser and Colinge, Journal of Proteomics 2013.

1.1 Installation

isobar is part of Bioconductor. To install the latest stable version available for your version of R, start R and enter:

```
> if (!requireNamespace("BiocManager", quietly=TRUE))
```

+ install.packages("BiocManager")

```
> BiocManager::install("isobar")
```

Bioconductor has semi-annual releases, and the releases are tied to specific to R versions. The latest development version of isobar are always available at GitHub (https://github.com/fbreitwieser/isobar). Use the devtools package to install it from the source:

```
> library(devtools)
> install_github("fbreitwieser/isobar")
```

1.2 Loading package

The first thing you need to do is load the package.

> library(isobar) ## load the isobar package

Check the installed version of isobar:

> packageVersion("isobar")

or equivalently for older versions of R:

> packageDescription("isobar")\$Version

2 Loading data

isobar can read identifications and quantifications from tab-separated and MGF files. Perl scripts are supplied to generate a tab-separated version from the vendor formats of Mascot and Phenyx, see appendix C. The "ID.CSV" format is described in appendix A. Rockerbox and MSGF+ CSV file formats can be read, too, and are recognized by their file extension (peptides.csv and msgfp.csv). Experimental support for the mzIdentML format is also available - please contact the maintainer in case of problems.

ID.CSV tab-separated file containing peptide-spectra matches and spectrum meta-information such as retention time, m/z and charge. Generated by parser scripts.

- **MGF** contains peak lists from which quantitative information on reporter tags are extracted. Must be centroided.
- **IBSPECTRA.CSV** tab-separated file containing the same columns as ID.CSV plus quantitative information extracted from MGF file - that means the reporter tag masses and intensities as additional columns.

readIBSpectra is the primary function to generate a IBSpectra object. The first argument is one of iTRAQ4plexSpectra, iTRAQ8plexSpectra, TMT2plexSpectra, TMT6plexSpectra and TMT10plexSpectra denotes the tag type and therefore class.

```
>
    ## generating IBSpectra object from ID.CSV and MGF
>
    ib <- readIBSpectra("iTRAQ4plexSpectra",list.files(pattern=".id.csv"),</pre>
+
            list.files(pattern=".mgf"))
    ## write in tabular IBSPECTRA.CSV format to file
>
    write.table(as.data.frame(ib),sep="\t",row.names=F,
>
            file="myexperiment.ibspectra.csv")
+
    ## generate from saved IBSPECTRA.CSV - MGF does not have to be supplied
>
>
    ib.2 <- readIBSpectra("iTRAQ4plexSpectra", "myexperiment.ibspectra.csv")</pre>
```

In case the MGF file is very big, it can be advanteguous to generate a smaller version containing only meta- and quantitative information before import in R. On Linux, the tool grep is readily available.

egrep '^[A-Z]|^1[12][0-9]\.' BIG.mgf > SMALL.mgf

2.1 ibspiked test samples

The examples presented are based on the dataset ibspiked_set1 which has been designed to test isobar's functionality and searched against the Swissprot human database with Mascot and Phenyx. ibspiked_set1 is an iTRAQ 4-plex data set comprised of a complex background (albumin- and IgG-depleted human plasma) and spiked proteins. MS analysis was performed in ThermoFisher Scientific LTQ Orbitrap HCD instrument with 2D shotgun peptide separation (see original paper for more details). The samples used for each iTRAQ channel are as follows:

- Depleted human plasma background (>150 protein detected);
- Spiked-in proteins with the following ratios
 - CERU_HUMAN (P00450) at concentrations 1:1:1:1;
 - CERU_RAT (P13635) at concentrations 1:2:5:10;
 - CERU_MOUSE (Q61147) at concentrations 10:5:2:1.

A second data set with ratios 1:10:50:100 is available as ibspiked_set2 from http://bininformatics.cemm.oeaw.ac.at/isobar.

The Ceruplasmins have been selected as the share peptides. Hereafter, we load the data pacakage and the ceru protein IDs are identified via the protein.g function, which provides a mean to retrieve data from ProteinGroup objects. ProteinGroup is a slot of IBSpectra objects and contains informations on proteins and their grouping. See 2.2.

```
> data(ibspiked_set1)
```

```
> ceru.human <- protein.g(proteinGroup(ibspiked_set1),"CERU_HUMAN")</pre>
```

```
> ceru.rat <- protein.g(proteinGroup(ibspiked_set1),"CERU_RAT")</pre>
```

```
> ceru.mouse <- protein.g(proteinGroup(ibspiked_set1),"CERU_MOUSE")</pre>
```

> ceru.proteins <- c(ceru.human,ceru.rat,ceru.mouse)</pre>

2.2 Protein information and grouping in ProteinGroup

When an **ibspectra.csv** is read, protein are grouped to identify proteins which have unique peptides. By default, only peptides with unique peptides are grouped.

The algorithm to infer protein groups works as follows:

- 1. Group proteins together which have been seen with exactly the same peptides (indistinguishableProteins) these are the protein.g identifiers.
- 2. Create protein groups (proteinGroupTable):
 - (a) Define proteins with specific peptides as reporters (reporterProteins)
 - (b) Get proteins which are contained ¹ by reporterProteins and group them below.
- 3. Create protein groups for proteins without specific peptides as above.

2.3 Loading data from CID/HCD (or CID/MS3, etc) experiments

A combined CID/HCD approach, in which for each precursor two fragmentation spectra are acquired, has proven useful to increase the number of identified and quantified peptide-spectrum matches. Usually, the reporter intensity information is taken from the HCD spectrum, and the peptide is identified based on the fragment ions in the CID spectrum.

To import these experiments, a comma-separated *mapping file* is needed, which contains the association from the identification to the quantification spectrum title.

Example mapping file (mapping.csv):

```
"hcd","cid"
"spectrum 1","spectrum 2"
"spectrum 3","spectrum 4"
```

By calling readIBSpectra with mapping.file="mapping.csv", the spectra titles in the id.file are matched to those in the peaklist.file. If the column names for the quantification and identification spectrum are not hcd and cid, resp., they can be set with the argument mapping=c(identification.spectrum="column name 1", quantification.spectrum="column name 2").

Example mapping file mapping2.csv:

The argument mapping.file can take multiple files as argument (which are read and concatenated), or a data.frame.

¹That means these proteins have a subset of the peptides of the reporter

2.4 Loading data from CID/HCD experiments with full HCD spectrum

If a full HCD spectrum was acquired, and both the HCD and CID spectrum are searched against a protein database, the argument id.file.domap to readIBSpectra can be used to merge both CID and HCD identifications:

```
> readIBSpectra("TMT6plexSpectra",
+ id.file="cid.identifications.csv",
+ peaklist.file="hcd.peaklist.mgf",
+ id.file.domap="hcd.identifications.csv",
+ mapping.file="mapping.csv",
+ ...
+ )
```

Here, the CID (cid.identifications.csv) and HCD identifications (hcd.identifications.csv) are combined and mapped according to mapping.csv. Diverging identifications are discarded.

2.5 Integrating and thresholding precursor purity measures

Savitzki et al., 2011 describe a measure for the precursor purity of fragment spectra. An implementation of the algorithm for the multiplierz platform is provided at http://sourceforge.net/apps/ trac/multiplierz/wiki/Precursors. The script creates _precursors.txt files for each RAW file. To integrate and threshold based on the s2i measure, use the following code:

```
> precursors <- read.delim("file1_precursors.txt",header=TRUE,sep="\t",stringsAsFactors=FALSE)
> idfile <- read.delim("file1.id.csv",header=TRUE,sep="\t",stringsAsFactors=FALSE)
> idfile <- merge(idfile,precursors,
+ by.x=c("fixed_spectrum","scans.from"),
+ by.y=c("fixed_spectrum","SCAN"),all.x=TRUE) # FIXME see /work/analysis/gwinter/raw-s</pre>
```

2.6 MSnbase integration

NOTE: It has been reported that the conversion functions are not functional for current versions of the package. We will try to resolve this.

MSnbase by Laurent Gatto provides data manipulation and processing methods for MS-based proteomics data. It provides import, representation and analysis of raw MS data stored in mzXML, mzML and mzData using the mzR package and centroided and un-centroided MGF peak lists. It allows to use and preprocess raw data whereas isobar requires centroided peak lists. In the future, the isobar class IBSpectra might be based on or replaced by MSnbase's class MSnSet. For now, methods for coercion are implemented:

```
> as(ibspectra,"MSnSet")
```

```
> as(msnset,"IBSpectra")
```

3 Data Analysis

3.1 Reporter mass precision

The distribution of observed masses from the reporter tags can be used to visualize the precision of the MS setup on the fragment level and used to set the correct window for isolation.

The expected masses of the reporter tags are in the slot reporterTagMasses of the implementations of the IBSpectra class. The experimental masses are in the matrix mass of AssayData; they can also be accessed by the method reporterMasses(x).

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```

```
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```

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reporterMassPrecision provides a plot of the distribution.

```
> print(reporterMassPrecision(ibspiked_set1))
```

4.2 Normalization and isotope impurity correction

Isotope impurity correction factors are supplied by labelling reagent manufacturers. Default values that can be modified by the user are available in **isobar** and corrections are obtained by simple linear algebra.

Due to differences between samples it is advisable to normalize data before further processing. By default, normalize corrects by a factor such that the median intensities in all reporter channels are equal.

See figure 2.

```
> ib.old <- ibspiked_set1
> ibspiked_set1 <- correctIsotopeImpurities(ibspiked_set1)
> ibspiked_set1 <- normalize(ibspiked_set1)</pre>
```

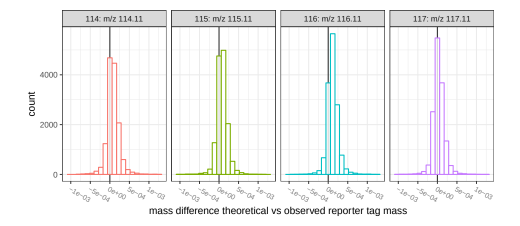


Figure 1: Reporter mass precision plot.

```
> par(mfrow=c(1,2))
> maplot(ib.old,channel1="114",channel2="117",ylim=c(0.5,2),
+ main="before normalization")
> abline(h=1,col="red",lwd=2)
> maplot(ibspiked_set1,channel1="114",channel2="117",ylim=c(0.5,2),
+ main="after normalization")
> abline(h=1,col="red",lwd=2)
```

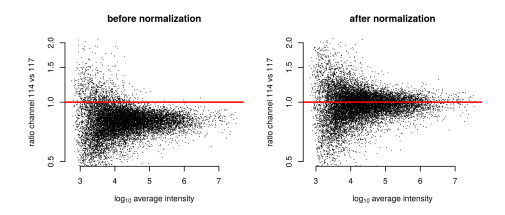


Figure 2: Ratio versus intensity plots ('MA plots') before and after applying normalization.

4.3 Fitting a noise model

A noise model is a approximation of the expected technical variation based on signal intensity. It is stable for a certain experimental setup and thus can be learned once. Noise is observed directly when comparing identical samples in multiple channels (1:1 iTRAQ/TMT sample) and we can use ibspiked_set1 background proteins as a 1:1 sample. Therefore we exclude the ceruplasmins before fitting a noise model using NoiseModel. See figure 3.

> ib.background <- subsetIBSpectra(ibspiked_set1,protein=ceru.proteins,direction="exclude")
> noise.model <- NoiseModel(ib.background)</pre>

[1] 0.03402699 11.26912497 1.42751145

Though only recommended when sufficient data are available, a method exist for the estimation of a noise model without a 1:1 dataset. It takes longer time as it first computes all the protein ratios to shift spectrum ratios to 1:1. To examplify this procedure, we only take rat and mouse CERU proteins from ibspiked_set1, see figure 3. The resultant noise model is a rough approximation only because of the very limitted data, see Breitwieser et al. Supporting Information, submitted, for a real example.

```
> ib.ceru <- subsetIBSpectra(ibspiked_set1,protein=ceru.proteins,
+ direction="include",
+ specificity="reporter-specific")
> nm.ceru <- NoiseModel(ib.ceru,one.to.one=FALSE,pool=TRUE)
3 proteins with more than 10 spectra, taking top 50.
[1] 0.0000000001 0.4177844310 0.1950041267
> maplot(ib.background,noise.model=c(noise.model,nm.ceru),
+ channel1="114",channel2="115",ylim=c(0.2,5),
+ main="95% CI noise model")
```

4.4 Protein and peptide ratio calculation

estimateRatio calculates the relative abundance of a peptide or protein in one tag compared to another. It calculates a weighted average (after outlier removal) of the spectrum ratios. The weights are the inverse of the spectrum ratio variances. It requires a IBSpectra and NoiseModel object and definitions of channel1, channel2, and the protein or peptide. The result is channel2/channel1.

```
> ## Calculate ratio based on all spectra of peptides specific
> ## to CERU_HUMAN, CERU_RAT or CERU_MOUSE. Returns a named
> ## numeric vector.
> 10<sup>estimateRatio</sup>(ibspiked_set1,noise.model,
                   channel1="114", channel2="115",
+
+
                   protein=ceru.proteins)['lratio']
   lratio
0.9397285
> ## If argument 'combine=FALSE', estimateRatio returns a data.frame
> ## with one row per protein
> 10<sup>estimateRatio</sup>(ibspiked_set1,noise.model,
                   channel1="114", channel2="115",
+
                   protein=ceru.proteins,combine=FALSE)[,'lratio']
+
[1] 1.0502867 1.8645614 0.4959669
> ## spiked material channel 115 vs 114:
> ##
                      CERU_HUMAN (P00450): 1:1
> ##
                      CERU_RAT
                                 (P13635): 2:1 = 2
> ##
                      CERU_MOUSE (Q61147): 5:10 = 0.5
```

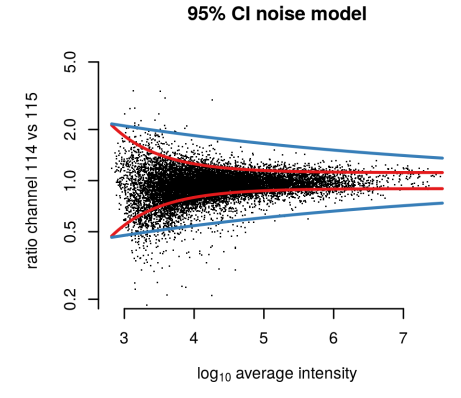


Figure 3: Red lines denote the 95 % confidence interval as estimated by the noise model on background proteins. The blue line is estimated as non 1:1 noise model based on only spectra of CERU proteins.

```
> ## Peptides shared between rat and mouse
> pep.shared <- peptides(proteinGroup(ibspiked_set1),
+ c(ceru.rat,ceru.mouse),set="intersect",
+ columns=c('peptide', 'n.shared.groups'))
> ## remove those which are shared with other proteins
> pep.shared <- pep.shared$peptide[pep.shared$n.shared.groups==2]
> ## calculate ratio: it is between the rat and mouse ratios
> 10^estimateRatio(ibspiked_set1,noise.model,
+ channel1="114",channel2="115",
+ peptide=pep.shared)['lratio']
```

lratio 0.6332743

When examining the global differences and differences in between classes, proteinRatios can be used. It is also suitable to inspect sample variability. The argument cl can be used to define class labels. If combn.method='interclass' or intraclass and summarize=TRUE, proteinRatios return a single summarized ratio across and within classes, resp..

```
> protein.ratios <- proteinRatios(ibspiked_set1,noise.model,cl=c("1","0","0","0"))</pre>
> ## defined class 114 and 115 as class 'T', 116 and 117 as class 'C'
> classLabels(ibspiked_set1) <- c("T","T","C","C")</pre>
>
 proteinRatios(ibspiked_set1, noise.model, protein=ceru.proteins,
+
                cl=classLabels(ibspiked_set1), combn.method="interclass",
                summarize=T)[,c("ac","lratio","variance")]
+
               lratio
                           variance
      ac
1 P00450
         0.007157362 0.0006269188
2 P13635 0.581686951 0.0503315336
3 Q61147 -0.563755457 0.0470624064
```

4.5 Protein ratio distribution and selection

Protein ratio distributions can be calculated ideally on biological replicated. To examine differentially expressed proteins, both sample variability information (random protein ratios) as a *fold-change* constraint, and ratio *precision* can be used. For a experimental setup with biological replicates in the same experiment (but different channels), the distribution of biological variability can be learned by computing the ratios between the replicates. With no replicates available, one has the choice to (a) model the actual protein ratios and just select the most extreme ratios; (b) learn the distribution from a previous experiment; or (c) assume a standard Cauchy distribution with location 0 and scale 0.1, 0.05, and 0.025, which correspond with $\alpha = 0.05$ roughly to fold changes of 4, 2, and 1.5.

A Cauchy distribution fits accurately this type of random protein ratio distribution: Cauchy is displayed in red, Gaussian in blue. In the case of ibspiked_set1, the many 1:1 proteins provide us with adequate data to learn the random protein ratio distribution, however only of the *technical* variation.

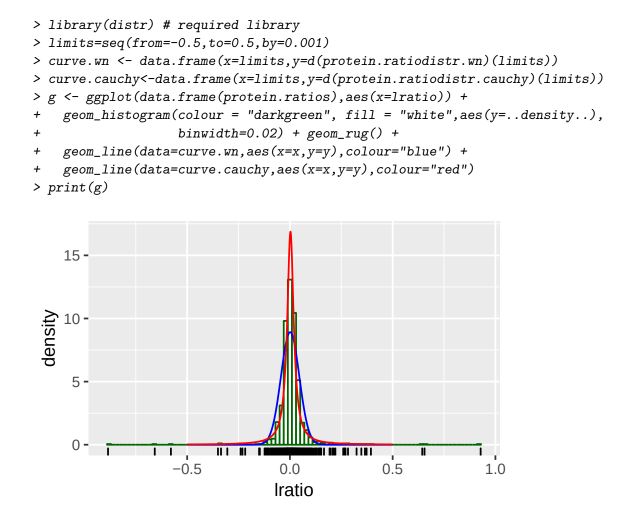


Figure 4: Histogram of all protein ratios in ibspiked_set1. A fit with a Gaussian and Cauchy probability density function is shown in blue and red, respectively.

Now, when supplying a ratiodistr parameter to estimateRatio and proteinRatios, sample and signal p-values are calculated, what we illustrate in the code below

```
> rat.list <-</pre>
    estimateRatio(ibspiked_set1,noise.model=noise.model,channel1="114",channel2="115",
+
                  protein=reporterProteins(proteinGroup(ibspiked_set1)), combine=F,
+
                  ratiodistr=protein.ratiodistr.cauchy)
+
> rat.list[rat.list[,"is.significant"]==1,]
                     variance
                                 var_hat_vk n.spectra n.na1 n.na2 p.value.rat
           lratio
P13635 0.2705767 0.006336962 8.435201e-05
                                                                 0 3.827430e-04
                                                  242
                                                           0
Q61147 -0.3045473 0.001047247 8.678212e-07
                                                                 0 1.105993e-21
                                                  139
                                                           0
       p.value.sample
                         p.value is.significant
P13635
           0.02236936 0.02555809
                                               1
Q61147
           0.01950968 0.01973225
                                               1
```

4.6 Detection of proteins with no specific peptides

It is well known that MS analysis only reveals the presence of so-called protein groups, defined as sets of proteins identified by the same set of peptides. The protein that contains all the peptides is the group reporter (there are possibly several group reporters) and if it has one specific peptide at least then its presence in the sample is certain. The status of the other proteins in the group is in general impossible to determine. When quantitative information is available, there is a potential to elucidate the structure of part of the protein groups.

In the example below, a subset IBSpectra object is created, containing only peptides shared between CERU_RAT and CERU_MOUSE, and those specific to CERU_RAT.

```
> ## peptides shared between CERU_RAT and CERU_MOUSE have been computed before
> pep.shared
```

```
[1] "AGLQAFFQVR"
                        "DNEEFLESNK"
                                           "DTANLFPHK"
                                                              "EMGPTYADPVCLSK"
 [5] "ETFTYEWTVPK"
                                                              "LYHSHVDAPK"
                        "GSLLADGR"
                                           "KGSLLADGR"
 [9] "NMATRPYSLHAHGVK" "RDTANLFPHK"
                                           "VFFEQGATR"
> ## peptides specific to CERU_RAT
>
 pep.rat <- peptides(proteinGroup(ibspiked_set1),protein=ceru.rat,</pre>
                       specificity="reporter-specific")
+
> ## create an IBSpectra object with only CERU_RAT and shared peptides
> ib.subset <- subsetIBSpectra(ibspiked_set1,</pre>
+
                                peptide=c(pep.rat,pep.shared),direction="include")
> ## calculate shared ratios
> sr <- shared.ratios(ib.subset,noise.model,</pre>
                       channel1="114", channel2="117",
+
                       ratiodistr=protein.ratiodistr.cauchy)
+
> sr
       reporter.protein protein2
                                     ratio1 ratio1.var n.spectra.1
                                                                          ratio2
                 P13635
                           Q61147 0.9281151 0.01351388
                                                                 241 0.009259277
Iratio
        ratio2.var n.spectra.2
lratio 0.001560001
                            273
>
```

> ## plot significantly different protein groups where 90% CI does not overlap

> ## CERU_MOUSE and CERU_RAT is detected, as expected.

shared.ratios.sign(sr,z.shared=1.282)

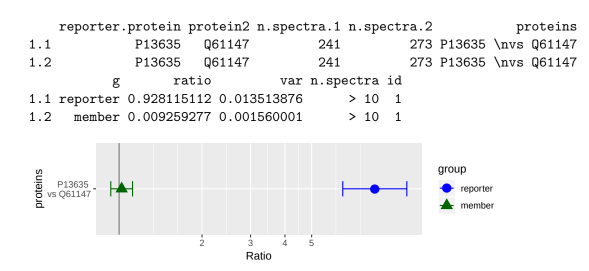


Figure 5: Peptides of spiked ceruplasmins have significantly different ratios between groups. Group *reporter* consists of peptides specific to CERU_RAT (P13635), group *member* are peptides shared between CERU_RAT and CERU_MOUSE (Q61147).

5 Report generation

>

isobar provides a rich interface for creating Excel and PDF reports for further analysis and quality control. The main entry function is create.reports. Alternatively the Rscript create_reports.R can be used. It is located in the report folder of the isobar installation, and reads the properties from a file in the working directory.

The possible values are defined in the report/properties.R file in the isobar installation. To generate a report with standard properties the following code should do the trick:

The properties can also be defined in a **properties.R** file which is located in the working directory. The properties are set in the following order:

- 'global' properties in ISOBAR-DIRECTORY/report/properties.R²
- 'local' properties in WORKING-DIRECTORY/properties.R
- command line arguments to create_reports.R or create.reports function

Appendix B provides a syntax-highlighted version of the properties file supplied with isobar, which sets the default parameters and provides some help in the comments. The number of parameters which can be set may seem a lot at first, however most times only a few are needed.

For successful completion, LAT_EX - for the PDF reports - and Perl - for the Excel reports - need to be installed.

²located in system.file('report', 'properties.R', package='isobar')

5.1 Files used for report generation

> ## execute to find the path and file location in your installation.

> system.file("report",package="isobar") ## path

> list.files(system.file("report",package="isobar")) ## files

create_reports.R R script which can be used to create QC and PDF reports It initializes the environment, reads properties and calls Sweave on QC and DA Sweave files. Additionally it generates a Excel data analysis report by calling tab2xls.pl.

isobar-qc.Rnw Sweave file with quality control plots.

isobar-analysis.Rnw Sweave file for generating a data analysis report with the list of all protein ratios and list of significantly different proteins.

properties.R Default configuration for create_reports.R. It is parsed as R code.

report-utils.R Helper R functions used in Sweave documents.

report-utils.tex Helper LATEX functions used in Sweave documents.

A File formats

A.1 ID CSV file format

The Perl parsers create ID CSV files - identification information for all matched spectra without quantitative information. You can create your own parser, the resulting file should be tab-delimited and contain the following columns. Only bold columns are obligatory. The information is redundant - that means if a peptide may stem from two different proteins the information of the identification is repeated.

Protein AC
Peptide sequence
Peptide modification string
Charge state
Theoretical peptide mass
Experimentally observed mass
Parent intensity
Retention time
Spectrum identifier
Protein search engine and score

A.2 IBSpectra CSV file format

IBSpectra file format has the same columns as the ID CSV format and additionally columns containing the quantitation information, namely Xtagname_mass and Xtagname_ions, for mass and intensity of each tag tagname. Below an example of the further columns for an iTRAQ 4plex IBSpectra.

$X114_mass$	reporter ion mass
$X115_mass$	reporter ion mass
$X116_mass$	reporter ion mass
$X117_mass$	reporter ion mass
$X114_{ions}$	reporter ion intensity
$\rm X115_ions$	reporter ion intensity
$X116_{ions}$	reporter ion intensity
$X117_{-ions}$	reporter ion intensity

B properties.R for report generation

```
##
## Isobar properties.R file
##
      for automatic report generation
##
## It is standard R code and parsed using sys.source
**************
## General properties
## Report type: Either 'protein' or 'peptide'
# report.level="peptide"
report.level="protein"
#attr(report.level,"allowed.values") <- c("protein","peptide")</pre>
## Isobaric tagging type. Use one of the following:
# type='iTRAQ4plexSpectra'
# type='iTRAQ8plexSpectra'
# type='TMT2plexSpectra'
# type='TMT6plexSpectra'
type=NULL
#attr(type,"allowed.values") <- IBSpectraTypes()</pre>
isotope.impurities=NULL
correct.isotope.impurities=TRUE
## Name of project, by default the name of working directory
## Will be title and author of the analysis reports.
name=basename(getwd())
author=paste0("isobaruRupackageuv",packageDescription("isobar")$Version)
## specifes the IBSpectra file or object
   - can be a data.frame (e.g. ibspectra=as.data.frame(ibspiked_set1) )
##
##
    - if it is a character string, it is assumed to be a file
##
      - if it ends on .rda, then it is assumed to be a R data object
##
       - if it does not exists, then it is may generated based on
         the peaklist and identifications properties
##
ibspectra=paste(name,"ibspectra.csv",sep=".")
## When replicates or 'samples belonging together' are analyzed, a
## ProteinGroup object based on all data should be constructed
## beforehand. This then acts as a template and a subset is used.
protein.group.template=NULL
## Via database or internet connection, informations on proteins (such
## as gene names and length) can be gathered. protein.info.f defines
## the function which takes a ProteinGroup object as argument
protein.info.f=getProteinInfoFromTheInternet
## Where should cached files be saved? Will be created if it does not
## exist
# cachedir="."
cachedir="cache"
## Regenerate cache files? By default, chache files are used.
```

```
regen=FALSE
```

```
## An ibspectra object can be generated from peaklists and
## identifications.
## peaklist files for quantitation, by default all mgf file in
## directory
peaklist=list.files(pattern="*\\.mgf$")
## id files, by default all id.csv files in directory
identifications=list.files(pattern="*\\.id.csv$")
## mapping files, for data quantified and identified with different but
## correspoding spectra. For example corresponding HCD-CID files.
## masses and intensities which are outside of the 'true' tag mass
## +/- fragment.precision/2 are discarded
fragment.precision=0.01
## filter mass outliers
fragment.outlier.prob=0.001
## Additional arguments of readIBSpectra can be set here
## decode.titles should be set to TRUE for Mascot search results
## as Mascot encodes the spectrum title (e.g. space -> %20)
readIBSpectra.args = list(
    mapping.file=NULL,
    decode.titles=FALSE
)
**************
## Quantification properties
normalize=TRUE
# if defined, normalize.factors will be used for normalization
normalize.factors=NULL
normalize.channels=NULL
normalize.use.protein=NULL
normalize.exclude.protein=NULL
normalize.function=median
normalize.na.rm=FALSE
peptide.specificity=REPORTERSPECIFIC
use.na=FALSE
## the parameter noise.model can be either a NoiseModel object or a file name
data(noise.model.hcd)
noise.model=noise.model.hcd
## If it is a file name, a noise model is estimated as non one-to-one
## and saved into the file. otherwise, the noise model is loaded from
## the file
# noise.model="noise.model.rda"
## Define channels for creation of a noise model, ideally a set of
## channels which are technical replicates.
noise.model.channels=NULL
```

```
## If noise.model.is.technicalreplicates is FALSE, the intensities
##
   are normalized for protein means, creating artifical technical
    replicates. For this procedure, only proteins with more than
##
    noise.model.minspectra are considered.
##
noise.model.is.technicalreplicates=FALSE
noise.model.minspectra=50
## class labels. Must by of type character and of same length as
## number of channels I. e. 4 for iTRAQ 4plex, 6 for TMT 6plex Example
## for iTRAQ 4plex:
## Class definitions of the isobaric tag channels.
## A character vector with the same length as channels
##
     (e.g. 4 for iTRAQ 4plex, 6 for TMT 6plex)
## Example for iTRAQ 4plex:
# class.labels=as.character(c(1,0,0,0))
# class.labels=c("Treatment","Treatment","Control","Control")
## Also names are possible - these serves as description in the report
## and less space is used in the rows
# class.labels=c("Treatment"="T","Treatment"="T","Control"="C","Control"="C")
class.labels=NULL
## The following parameters define which ratios are calculated.
## summarize ratios with equal class labels, set to TRUE when replicates are used
summarize=FALSE
## combn.method defines which ratios are calculated - versus a channel or a class,
## all the ratios within or across classes, or all possible combinatioins.
## When summarize=TRUE is set, use "interclass", "versus.class", or "intraclass"
# combn.method="global"
# combn.method="versus.class"
# combn.method="intraclass"
# combn.method="interclass"
combn.method="versus.channel"
vs.class=NULL
cmbn = NULL
## Arguments given to 'proteinRatios' function. See ?proteinRatios
ratios.opts = list(
    sign.level.sample=0.05,
    sign.level.rat=0.05,
    groupspecific.if.same.ac=TRUE)
quant.w.grouppeptides=c()
min.detect=NULL
preselected=c()
### Biological Variability Ratio Distribution options
## ratiodistr can be set to a file or a 'Distribution object. ' If
## NULL, or the specified file is not existent, the biological
## variability of ratios is estimated on the sample at hand and
## written to cachedir/ratiodistr.rda or the specified file.
```

ratiodistr=NULL

```
## Ideally, when the biological variability is estimated for the
## sample at hand, a biological replicate is present (/ie/ same class
## defined in class labels). Classes can also be assigned just for
## estimation of the ratio distribution, /eg/ to choose biologically
## very similar samples as pseudo replicates.
ratiodistr.class.labels=NULL
## Function for fitting. Available: fitCauchy, fitTlsd
ratiodistr.fitting.f=fitCauchy
## Use symetrical ratios - i.e. for every ratio r add a ratio -r
##
   prior to fitting of a distribution
ratiodistr.symmetry=TRUE
## If defined, use z-score instead of ratio distribution
# zscore.threshold=2.5
zscore.threshold=NULL
## PTM properties
## PhosphoSitePlus dataset which can be used to annotate known
## modification sites. Download site:
## http://www.phosphosite.org/staticDownloads.do
phosphosite.dataset <- NULL
## Modification to track. Use 'PHOS' for phosphorylation.
# ptm <- c('ACET','METH','UBI','SUMO', 'PHOS')</pre>
ptm <- NULL
## file name of rda or data.frame with known modification sites
## gathered with ptm.info.f. defaults to 'cachedir/ptm.info.rda'
ptm.info <- NULL
## Function to get PTM modification sites from public datasets
# ptm.info.f <- getPtmInfoFromNextprot</pre>
# ptm.info.f <- function(...)</pre>
#
     getPtmInfoFromPhosphoSitePlus(...,modification="PHOS")
# ptm.info.f <- function(...)</pre>
#
     getPtmInfoFromPhosphoSitePlus(...,modification=ptm)
ptm.info.f <- getPtmInfoFromNextprot</pre>
## A protein quantification data.frame (generated with
## 'proteinRatios'). The ratio and variance are used to correct the
## observed modified peptide ratios Needs to have the experimental
## setup as the modified peptide experiment
correct.peptide.ratios.with <- NULL
## Protein groups to use with correct.peptide.ratios()
correct.peptide.ratios.with_protein.group <- NULL</pre>
## The correlation between peptide and protein ratios defines the
## covariance
```

```
## Var(ratio m) = Var(ratio mp) + Var(ratio p)
##
                                     + 2 * Cov(ratio mp, ratio p),
## Cov(ratio mp, ratio p) = 2 * cor * Sd(ratio mp) * Sd(ratio p),
## with m = modifcation, mp = modified peptide, p = protein
peptide.protein.correlation <- 0</pre>
## quantification table whose columns are attached to the XLS
## quantification table
compare.to.quant <- NULL</pre>
*************
## Report properties
write.qc.report=TRUE
write.report=TRUE
write.xls.report=TRUE
## Use name for report, ie NAME.quant.xlsx instead of
## isobar-analysis.xlsx
use.name.for.report=TRUE
## PDF Analysis report sections: Significant proteins and protein
## details
show.significant.proteins=FALSE
show.protein.details=TRUE
### QC REPORT OPTIONS ###
#qc.maplot.pairs=FALSE # plot one MA plot per tag (versus all others)
qc.maplot.pairs=TRUE # plot MA plot of each tag versus each tag
### XLS REPORT OPTIONS ###
## Spreadsheet format: Either 'xlsx' or 'xls'
# spreadsheet.format="xlsx"
spreadsheet.format="xlsx"
## XLS report format 'wide' or 'long '.
## 'wide' format outputs ratios in separate columns of the same record
## (i.e. one line per protein)
## 'long' format outputs ratios in separate records (i.e. one line per
## ratio)
# xls.report.format="wide"
xls.report.format="long"
## XLS report columns in quantification tab
## possible values: ratio, is.significant, CI95.lower, CI95.upper,
##
                    ratio.minus.sd, ratio.plus.sd,
##
                    p.value.ratio, p.value.sample, n.na1, n.na2,
##
                    log10.ratio, log10.variance,
                    log2.ratio, log2.variance
##
## only for summarize=TRUE: n.pos, n.neg
xls.report.columns <- c("ratio","is.significant","ratio.minus.sd",</pre>
                       "ratio.plus.sd", "p.value.ratio", "p.value.sample",
                       "log10.ratio","log10.variance")
```

```
## Perl command to be used for Excel report generation
# perl.cmd = "C:/Strawberry/perl/bin/perl.exe"
# perl.cmd = "C:/Perl/bin/perl.exe"
# perl.cmd = "perl5"
perl.cmd = "perl'
**************
## Etc
sum.intensities=FALSE
datbase="Uniprot"
scratch=list()
##
 compile LaTeX reports into PDF files
#
compile=TRUE
# zip final report files into archive
zip=FALSE
# warning level (see 'warn' in ?options)
warning.level=1
## Novel options
shrink.mean=TRUE
use.t.stat=TRUE
```

C Dependencies

C.1 LATEX and PGF/TikZ

 ET_EX is a high-quality typesetting system; it includes features designed for the production of technical and scientific documentation. It is available as free software³. PGF is a T_EX macro package for generating graphics It comes with a user-friedly syntax layer called TikZ⁴.

 IAT_EX is used for creating PDF analysis reports, with the PGF package creating the graphics. Go to http://www.latex-project.org to get information on how to download and install a IAT_EX system and packages.

C.2 Perl

Perl is a high-level, general-purpose, interpreted, dynamic programming language. Perl is required for two tasks:

- Conversion of Pidres XML and Mascot DAT files to ID CSV format;
- Creation of Microsoft Excel format data analysis report.

```
<sup>3</sup>http://www.latex-project.org
```

⁴http://sourceforge.net/projects/pgf

Go to http://www.perl.org to download and get help on the installation of Perl on your Operating System. For file format conversion, perl module Statistics::Lite is required. For Excel export Spreadsheet::WriteExcel. All Perl scripts are in the subdirectory pl of the isobar package installation.

```
> ## execute to find the path and file location in your installation.
```

```
> system.file("pl",package="isobar") ## path
```

```
> list.files(system.file("pl",package="isobar")) ## files
```

mascotParser2.pl and pidresParser2.pl convert from respective protein search outputfiles to a XML file format, which can be converted into a CSV file readable by *isobar* by using psx2tab2.pl.

mascotParser2.pl coverts from Mascot format, and requires the file modifconv.csv as a definition of modification names. pidresParser2.pl converts from Phenyx output and requires the file parsersConfig.xml. tab2xls.pl converts csv file to different sheets of an Excel spreadsheet.

D Session Information

The version number of R and packages loaded for generating the vignette were:

- > toLatex(sessionInfo())
 - R version 3.5.2 (2018-12-20), x86_64-pc-linux-gnu
 - Locale: LC_CTYPE=en_US.UTF-8, LC_NUMERIC=C, LC_TIME=en_US.UTF-8, LC_COLLATE=C, LC_MONETARY=en_US.UTF-8, LC_MESSAGES=en_US.UTF-8, LC_PAPER=en_US.UTF-8, LC_NAME=C, LC_ADDRESS=C, LC_TELEPHONE=C, LC_MEASUREMENT=en_US.UTF-8, LC_IDENTIFICATION=C
 - Running under: Ubuntu 16.04.5 LTS
 - Matrix products: default
 - BLAS: /home/biocbuild/bbs-3.8-bioc/R/lib/libRblas.so
 - LAPACK: /home/biocbuild/bbs-3.8-bioc/R/lib/libRlapack.so
 - Base packages: base, datasets, grDevices, graphics, methods, parallel, stats, utils
 - Other packages: Biobase 2.42.0, BiocGenerics 0.28.0, distr 2.7.0, ggplot2 3.1.0, isobar 1.28.1, sfsmisc 1.1-3, startupmsg 0.9.5
 - Loaded via a namespace (and not attached): AnnotationDbi 1.44.0, DBI 1.0.0, IRanges 2.16.0, MASS 7.3-51.1, R6 2.3.0, RCurl 1.95-4.11, RSQLite 2.1.1, Rcpp 1.0.0, S4Vectors 0.20.1, XML 3.98-1.16, assertthat 0.2.0, bindr 0.1.1, bindrcpp 0.2.2, biomaRt 2.38.0, bit 1.1-14, bit64 0.9-7, bitops 1.0-6, blob 1.1.1, colorspace 1.3-2, compiler 3.5.2, crayon 1.3.4, digest 0.6.18, dplyr 0.7.8, glue 1.3.0, grid 3.5.2, gtable 0.2.0, hms 0.4.2, httr 1.4.0, labeling 0.3, lazyeval 0.2.1, magrittr 1.5, memoise 1.1.0, munsell 0.5.0, pillar 1.3.1, pkgconfig 2.0.2, plyr 1.8.4, prettyunits 1.0.2, progress 1.2.0, purr 0.2.5, rlang 0.3.0.1, scales 1.0.0, stats4 3.5.2, stringi 1.2.4, stringr 1.3.1, tibble 2.0.0, tidyselect 0.2.5, tools 3.5.2, withr 2.1.2