isobar for developers

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1 Introduction

This documents highlights the structure of the S4 classes and methods in the isobar pacakge.

> library(isobar)

getClass("IBSpectra")

2 Classes

2.1 IBSpectra

Virtual Class "IBSpectra" [package "isobar"] Slots: Name: proteinGroup reporterTagNames reporterTagMasses Class: ProteinGroup character numeric isotopeImpurities assayData Name: log Class: matrix matrix AssayData Name: phenoData featureData experimentData Class: AnnotatedDataFrame AnnotatedDataFrame MIAxE

```
protocolData . classVersion
Name:
              annotation
Class:
               character AnnotatedDataFrame
                                                       Versions
Extends:
Class "eSet", directly
Class "VersionedBiobase", by class "eSet", distance 2
Class "Versioned", by class "eSet", distance 3
Known Subclasses:
Class "iTRAQSpectra", directly
Class "TMTSpectra", directly
Class "iTRAQ4plexSpectra", by class "iTRAQSpectra", distance 2
Class "iTRAQ8plexSpectra", by class "iTRAQSpectra", distance 2
Class "TMT2plexSpectra", by class "TMTSpectra", distance 2
Class "TMT6plexSpectra", by class "TMTSpectra", distance 2
Class "TMT6plexSpectra2", by class "TMTSpectra", distance 2
Class "TMT10plexSpectra", by class "TMTSpectra", distance 2
```

identifications and quantitative values. Spectrums are identified as stemming from distinct peptides, and quantitative information of each spectrum are extracted from a certain m/z region.

IBSpectra class holds this qualitative and quantitative information. It is a virtual class. It extends eSet from Biobase to store meta-information of spectrum identifications and quantitative information (m/z and intensity) of reporter tags. eSet is extended by slots for protein grouping, tag names, tag masses and isotope impurity correction matrix.

ProteinGroup objects store the mapping and grouping of peptide level identifications to protein identifications.

IBSpectra is a virtual class. Currently used isobaric tagging kits iTRAQ 4plex and 8plex, and TMT 2plex and 6plex are implemented in the iTRAQ4plexSpectra, iTRAQ8plexSpectra, TMT2plexSpectra, TMT6plexSpectr and TMT10plexSpectr, respectively. These are subclasses of iTRAQSpectra and TMTSpectra, resp. which in turn are virtual subclasses of IBSpectra.

2.2 ProteinGroup

```
Name: indistinguishableProteins
                                         proteinGroupTable
Class:
                                                 data.frame
                       character
                                      isoformToGeneProduct
             overlappingProteins
Name:
                                                data.frame
Class:
                          matrix
                     proteinInfo
Name:
                                               peptideInfo
Class:
                      data.frame
                                                data.frame
Name:
               .__classVersion__
Class:
                        Versions
Extends:
Class "VersionedBiobase", directly
Class "Versioned", by class "VersionedBiobase", distance 2
```

mapped back to proteins. This mapping leads to protein groups, which explain the observed peptides according to the parsimony law.

A ProteinGroup object is generated when a IBSpectra object is created by readIBSpectra. Protein to peptide to spectrum mapping is extracted from a suitable identication format¹

2.3 NoiseModel

```
getClass("NoiseModel")
Virtual Class "NoiseModel" [package "isobar"]
Slots:
Name:
               na.region
                             low.intensity
                                                                     parameter
Class:
                                   numeric
                                                    function
               numeric
                                                                       numeric
Name: .__classVersion__
Class:
               Versions
Extends:
Class "VersionedBiobase", directly
Class "Versioned", by class "VersionedBiobase", distance 2
Known Subclasses: "ExponentialNoANoiseModel", "ExponentialNoiseModel", "InverseNo:
"InverseNoANoiseModel", "GeneralNoiseModel"
```

in the spectrum-level ratios of a certain experimental setup.

¹IBSpectra CSV, and MzIdentML format. Mascot DAT and Phenyx pidres.xml format converters to IBSpectra format are provided.

3 Session Information

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

- R version 4.3.1 (2023-06-16), x86_64-pc-linux-gnu
- Locale: LC_CTYPE=en_US.UTF-8, LC_NUMERIC=C, LC_TIME=en_GB, 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
- Time zone: America/New_York
- TZcode source: system (glibc)
- Running under: Ubuntu 22.04.3 LTS
- Matrix products: default
- BLAS: /home/biocbuild/bbs-3.18-bioc/R/lib/libRblas.so
- LAPACK: /usr/lib/x86_64-linux-gnu/lapack/liblapack.so.3.10.0
- Base packages: base, datasets, grDevices, graphics, methods, stats, utils
- Other packages: Biobase 2.62.0, BiocGenerics 0.48.0, isobar 1.48.0
- Loaded via a namespace (and not attached): AnnotationDbi 1.64.0, BiocFileCache 2.10.0, Biostrings 2.70.0, DBI 1.1.3, GenomeInfoDb 1.38.0, GenomeInfoDbData 1.2.11, IRanges 2.36.0, KEGGREST 1.42.0, MASS 7.3-60, R6 2.5.1, RCurl 1.98-1.12, RSQLite 2.3.1, Rcpp 1.0.11, S4Vectors 0.40.0, XML 3.99-0.14, XVector 0.42.0, biomaRt 2.58.0, bit 4.0.5, bit64 4.0.5, bitops 1.0-7, blob 1.2.4, cachem 1.0.8, cli 3.6.1, colorspace 2.1-0, compiler 4.3.1, crayon 1.5.2, curl 5.1.0, dbplyr 2.3.4, digest 0.6.33, distr 2.9.2, dplyr 1.1.3, fansi 1.0.5, fastmap 1.1.1, filelock 1.0.2, generics 0.1.3, ggplot2 3.4.4, glue 1.6.2, grid 4.3.1, gtable 0.3.4, hms 1.1.3, httr 1.4.7, lifecycle 1.0.3, magrittr 2.0.3, memoise 2.0.1, munsell 0.5.0, pillar 1.9.0, pkgconfig 2.0.3, plyr 1.8.9, png 0.1-8, prettyunits 1.2.0, progress 1.2.2, rappdirs 0.3.3, rlang 1.1.1, scales 1.2.1, sfsmisc 1.1-16, startupmsg 0.9.6, stats4 4.3.1, stringi 1.7.12, stringr 1.5.0, tibble 3.2.1, tidyselect 1.2.0, tools 4.3.1, utf8 1.2.4, vctrs 0.6.4, xml2 1.3.5, zlibbioc 1.48.0