

Using R and Bioconductor for proteomics data analysis

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Introduction

Even well-known and respected leaders in proteomics agree that it lies 10 years behind genomics. There are several valid reasons for this, including the chemical complexity of proteins, the technical complexity of the instrumentation (in particular mass-spectrometry - MS) and the vast possibilities in the study of proteins. An often overseen albeit essential component of this failure is arguably the software that is promoted inside the proteomics community. Computational proteomics researchers who value quality software, comprehensive data analysis and reproducible research ought to illustrate how more flexible and advanced tools can effectively be used and demonstrate their advantages. Here, we illustrate some examples of proteomics data analysis in R, in particular low level **raw MS data** manipulation, labelled and label-free **quantitation** and peptide **identification**, taken from the RforProteomics package [4].

Working with raw data

The proteomics community has developed a range of data standards and formats for MS data (the latest being mzML) to overcome the shortcomings of closed, binary vendor-specific formats.

One of the main projects that implement parsers for the XML-based open formats is the C++ proteowizard project [2], which is interfaced by the mzR Bioconductor package using the Rcpp package.

```
library("mzR")
fname <- dir(system.file(package = "MSnbase", dir = "extdata"),
  full.name = TRUE, pattern = "mzXML$")
ms <- openMSfile(fname)
```

The resulting `ms` object is a file handle that allows fast direct and random access to the individual spectra. `mzR` is used by a variety of other packages like `xcms`, `MSnbase`, `RMassBank` and `TargetSearch`.

Challenges Further improve support of raw MS data and develop the range of supported formats, in particular identification (mzIdentML) and quantitation (mzQuantML) formats.

Labelled quantitation

The same raw data file can be imported in a convenient higher level container and directly processed, plotted, quantified and normalised with the `MSnbase` [5] software.

```
exp <- readMSData(fname, verbose = FALSE)
plot(exp[["X3.1"]], full = TRUE, reporters = iTRAQ4)
set <- quantify(exp, method = "trap", reporters = iTRAQ4,
  verbose = FALSE, parallel = TRUE)
head(exprs(set), n = 3)
```

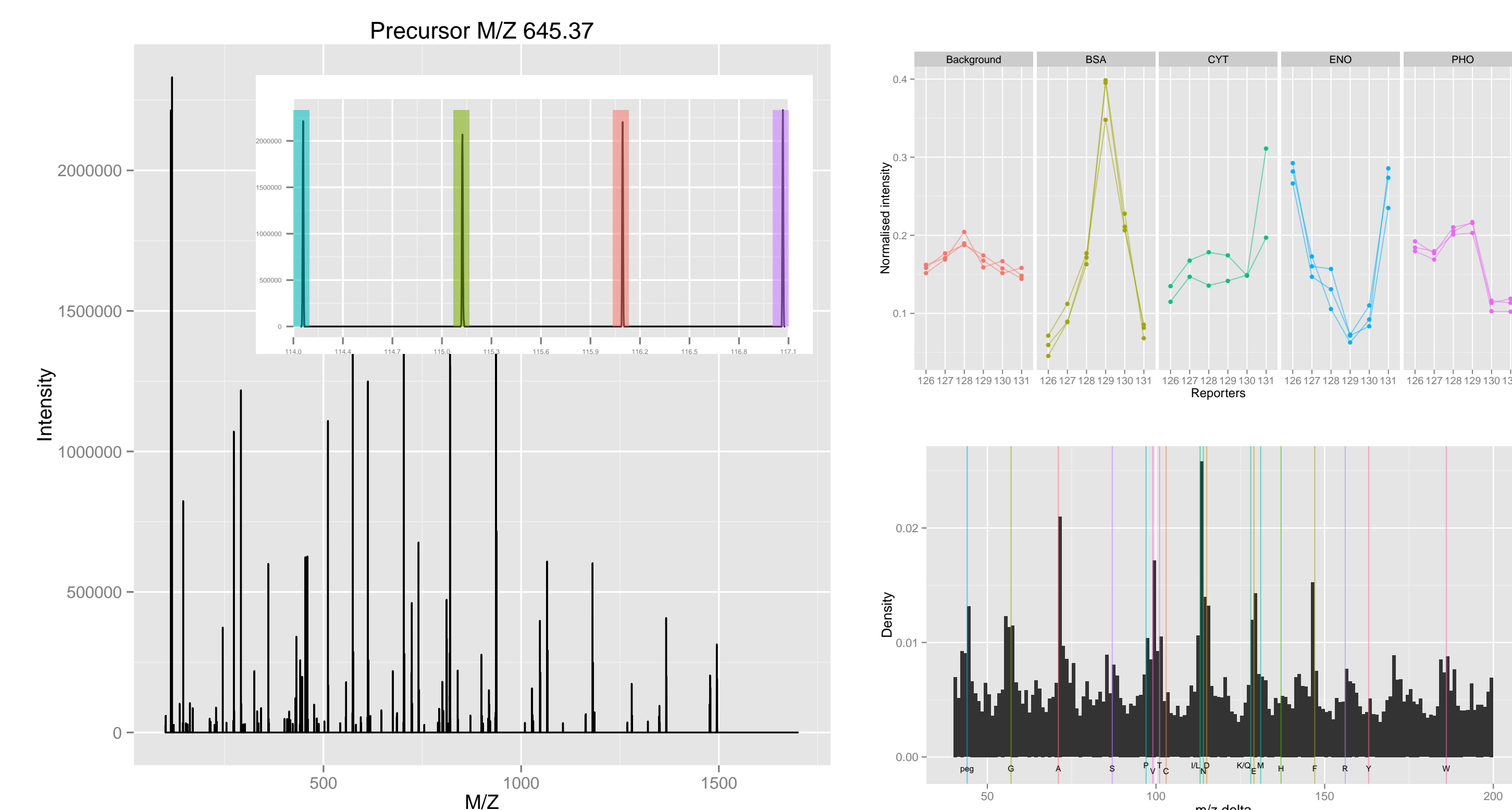


Figure: MS² spectrum of an iTRAQ 4-plex experiment showing the 4 isobaric reporter ions, as produced by `plot` above (left). Peptides of interesting from a spiked-in experiment (top right) and distribution of the *m/z* differences of all MS² spectra from the same experiment, use as a peptide-spectrum matching quality assessment (bottom right).

Challenges Although labelled MS² quantitation is well supported with `MSnbase` and isobar, metabolic labelling techniques like ¹⁵N or SILAC still need to be supported.

Label-free quantitation

Label-free quantitation is available in the `xcms` [8] and `MALDIquant` [7] packages.

The latter provides a complete pipeline, including baseline subtraction, smoothing, peak detection and alignment using warping functions, handling of replicated measurements as well as allowing spectra with different resolutions.

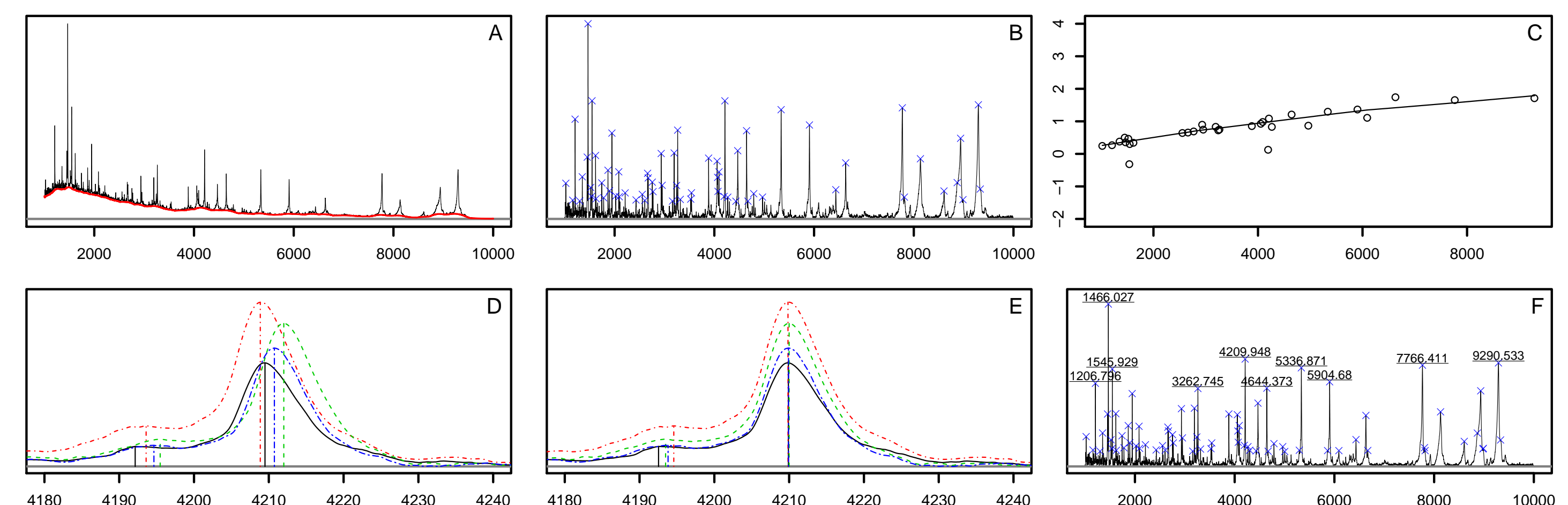


Figure: Illustration of the MALDIquant pipeline: raw spectrum with estimate baseline (A); variance-stabilized, smoothed, baseline-corrected spectrum with detected peaks (B); fitted warping function for peak alignment (C); four unaligned peaks (D); four aligned peaks (E); merged spectrum with discovered and labeled peaks (F).

A complete pipeline for MS^e data independent acquisition, including support for ion mobility separation is available in the `synapter` package [1] that, among other, transfers identification between acquisitions to substantially reduce missing values.

Challenges Application and benchmarking of label-free pipeline on popular Thermo Orbitrap instruments.

Peptide identification

The recently released `rTANDEM` package encapsulates the X!Tandem [3] search engine in R.

It uses the same XML-based parameter files as the native application or dedicated R parameter object. Result files can be directly parsed and mined in R .

```
xmlres <- rtandem(spectra.mgf, taxon = "yeast",
  taxonomy = "taxonomy.xml",
  default.parameters = "default-params.xml")
## or xmlres <- tandem(param)
res <- GetResultsFromXML(xmlres)
proteins <- GetProteins(res) ## data.table objects
peptides <- GetPeptides(res)
```

A complete pipeline with support for identification is welcome at the cost of additional development and maintenance time for developers. With support for `mzIdentML` files, it will become possible to import identification data from most search engines, this facilitating the integration of R based pipelines with existing tools.

Challenges Better integration of identification and raw/quantitation data infrastructures.

Conclusions and perspectives

The flexibility of the R environment and the breath of available packages is sometimes daunting for newcomers and introductory points of entry are welcome. The RforProteomics package [<https://github.com/lgatto/RforProteomics>] ought to assume such a role. For this, RforProteomics should be a collaborative project and contributions through the github repository are encouraged.

Despite well known advantages in terms of statistical analyses of data and some unique software for proteomics and mass-spectrometry data analysis, there remains a lot of efforts and work to be done for R/Bioconductor to become a complete framework for proteomics data processing. These efforts should be tackled by a group of developers. It is our hope that the RforProteomics will be a helpful targeted introduction to new users and motivate collaborative development of package developers.

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