MSnbase, efficient and elegant R-based

processing and visualisation of raw mass

spectrometry data

Laurent Gatto,*,† Sebastian Gibb,‡ and Johannes Rainer¶

†Computational Biology Unit, de Duve Institute, Université catholique de Louvain,

Brussels, Belgium

Department of Anaesthesiology and Intensive Care of the University Medicine Greifswald,

Germany

¶Institute for Biomedicine, Eurac Research, Affiliated Institute of the University of Lübeck,

Bolzano, Italy

E-mail: laurent.gatto@uclouvain.be

Abstract

We present version 2 of the MSnbase R/Bioconductor package. MSnbase provides

infrastructure for the manipulation, processing and visualisation of mass spectrometry

data. We focus on the new on-disk infrastructure, that allows the handling of large raw

mass spectrometry experiment on commodity hardware and illustrate how the package

is used for elegant data processing, method development, and visualisation.

Introduction

Mass spectrometry is a powerful technology to assay chemical and biological samples. It is

used in routine applications with well characterised protocols such as diagnostics, as well as a

1

development platform, with the aim to improve on existing methods protocols and devise new ones. The complexity and diversity of mass spectrometry yield data that is itself complex and often times of considerable size, that require non trivial processing before producing interpretable results. This is particularly relevant, and can The complexity and size of these data constitute a significant challenge for method protocol development: in addition to the development of sample processing and mass spectrometry methods that yield the raw data, it is essential there is a need to process, analyse, interpret and assess these new data to demonstrate the improvement in the technical, analytical and computational workflows.

Practitioners have a diverse catalogue of software tools to explore, process and interpret mass spectrometry data at their disposal. These range from low level software libraries that are aimed at programmers to enable development of new applications, to more user-oriented applications with graphical user interfaces which provide a more limited set of functionalities to address a defined scope. Examples of software libraries include Java-based jmzML¹ or C/C++-based ProteoWizard.² ProteomeDiscoverer (Thermo Scientific) Thermo Scientific Proteome Discoverer (Thermo Fisher Scientific), MaxQuant³ and PeptideShaker⁴ are among the most widely used user-centric applications.

In this software note, we present version 2 of the MSnbase⁵ software, available from the Bioconductor⁶ project. The package, like other software such as Python-based pyOpenMS,⁷ spectrum_utils⁸ or Pyteomics,⁹ offers a platform that lies between low level libraries and enduser software. MSnbase provides a flexible R¹⁰ command-line environment for metabolomics and proteomics mass spectrometry-based applications. It lays out a sound infrastructure to work with raw mass spectrometry data from MS files in mzML, mzXML, mzData or ANDI-MS/netCDF format as well as quantitative and proteomics identification data. The package enables manipulation (for example subsetting, filtering, or accessing specific parts thereof), detailed step-by-step processing (for example smoothing and centroiding of raw profile-mode MS data, or normalisation and imputation of quantitative data), analysis and visualisation of these data and the development of novel computational mass spectrometry methods.¹¹

Extensive documentation and use cases are provided in *package vignettes* ¹² and workflows. ¹³ Here, we focus on the new developments pertaining to raw mass spectrometry data handling and processing.

Infrastructure for raw data

In MSnbase, mass spectrometry experiments are handled as MSnExp objects. While the implementation is more complex, it is useful to schematise a raw data experiment as being composed of raw data, i.e. a collection of individual spectra, as well as spectra-level metadata (Figure 1). Each spectrum is composed of m/z values and associated intensities. The metadata are represented by a single table with variables along the columns and each row associated to a spectrum. Among the metadata available for each spectrum, there are MS level, acquisition number, retention time, precursor m/z and intensity (for MS level 2 and above), and many more. MSnbase relies on the mzR package 2 to import raw mass spectrometry data from one of the many community-maintained open standards formats (mzML, mzXML, mzData or ANDI-MS/netCDF) and provides a rich interface to manipulate such objects. The code chunk below illustrates such an object as displayed in the R console and an enumeration of the metadata fields.

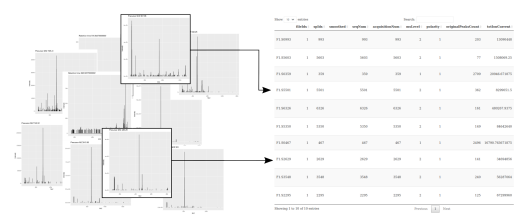


Figure 1: Schematic representation of what is referred to by *raw data*: a collection of mass spectra and a table containing spectrum-level annotations along the lines. Raw data is contained in one of the many community-maintained open standards formats (mzML, mzXML, mzData or ANDI-MS/netCDF).

```
> show(ms)
MSn experiment data ("OnDiskMSnExp")
Object size in memory: 0.54 Mb
- - - Spectra data - - -
MS level(s): 1 2 3
Number of spectra: 994
MSn retention times: 45:27 - 47:6 minutes
- - - Processing information - - -
Data loaded [Sun Apr 26 15:40:58 2020]
MSnbase version: 2.13.6
- - - Meta data - - -
phenoData
  rowNames: MS3TMT11.mzML
 varLabels: sampleNames
  varMetadata: labelDescription
Loaded from:
 MS3TMT11.mzML
protocolData: none
featureData
  featureNames: F1.S001 F1.S002 ... F1.S994 (994 total)
  fvarLabels: fileIdx spIdx ... spectrum (35 total)
  fvarMetadata: labelDescription
experimentData: use 'experimentData(object)'
> fvarLabels(ms)
 [1] "fileIdx"
                                  "spIdx"
 [3] "smoothed"
                                  "seqNum"
 [5] "acquisitionNum"
                                  "msLevel"
```

```
[7] "polarity"
                                   "originalPeaksCount"
 [9] "totIonCurrent"
                                   "retentionTime"
[11] "basePeakMZ"
                                   "basePeakIntensity"
[13] "collisionEnergy"
                                   "ionisationEnergy"
[15] "lowMZ"
                                   "highMZ"
                                   "precursorMZ"
[17] "precursorScanNum"
[19] "precursorCharge"
                                   "precursorIntensity"
    "mergedScan"
                                   "mergedResultScanNum"
[21]
                                   "mergedResultEndScanNum"
    "mergedResultStartScanNum"
[25]
                                   "filterString"
    "injectionTime"
    "spectrumId"
                                   "centroided"
[27]
    "ionMobilityDriftTime"
                                   "isolationWindowTargetMZ"
[31] "isolationWindowLowerOffset" "isolationWindowUpperOffset"
[33] "scanWindowLowerLimit"
                                   "scanWindowUpperLimit"
[35] "spectrum"
```

In the following sections, we describe MSnbase's ability to efficiently handle very large mass spectrometry data and experiments and how it how MSnbase can be used for data processing and visualisation. An example of its ability to efficiently handle also very large mass spectrometry experiments (in this case with 5,773,464 spectra in 1,182 mzXML files) is provided as supplementary information. We will also illustrate how it makes use of the forward-pipe operator (%>%) defined in the magrittr package. This operator has proved useful to develop non-trivial analyses by combining individual functions into easily readable and elegant pipelines.

On-disk backend

The main feature in version 2 of the MSnbase package was the addition of different backends for raw data storage, namely *in-memory* and *on-disk*. The following code chunk demonstrates how to create two MSnExp objects importing data from an mzML file and storing data them either in-memory or on-disk.

```
library("MSnbase")
raw_mem <- readMSData("file.mzML", mode = "inMemory")
raw_dsk <- readMSData("file.mzML", mode = "onDisk")</pre>
```

Both modes rely on the mzR² package to access the spectra (using the mzR::peaks() function) and the metadata (using the mzR::header() function) in the data files. The former is the legacy storage mode, implemented in the first version of the package, that loads all the raw data and the metadata into memory upon creation of the in-memory MSnExp object. This solution doesn't scale for modern large dataset, and was complemented by the on-disk backend. The on-disk backend only loads the metadata into memory when the on-disk MSnExp is created and accesses the spectra data (i.e. m/z and intensity values) in the original files on disk only when needed (see below and Figure 2 (d)), such as for example for plotting. There are two direct benefits using the on-disk backend, namely faster reading and reduced memory footprint. Figure 2 shows 5-fold faster reading times (a) and over a 10-fold reduction in memory usage (b).

Because the on-disk backend does not hold all the data in memory, direct data manipulations are not possible. We thus implemented a *lazy processing* mechanism for this backend that caches any data manipulation operations in a processing queue in the object itself. Only when the user accesses m/z or intensity values are these operations are applied. The on-disk backend also offers efficient data manipulation by way of *lazy processing* in the object itself. Operations on the raw data are stored in a processing queue and only effectively applied when the user accesses m/z or intensity values. As an additional advantage, operations

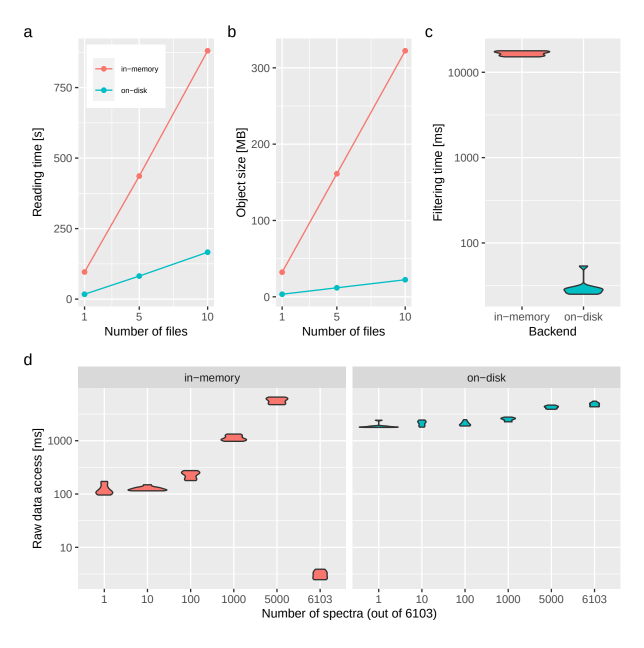


Figure 2: (a) Reading time (in seconds) and (b) data size in memory (in MB) to read/store 1, 5 and 10 files containing 1431 MS1 (on-disk only) and 6103 MS2 (on-disk and in-memory) spectra. (c) Filtering benchmark assessed over 10 interactions on in-memory and on-disk data containing 6103 MS2 spectra. (d) Access time to spectra for the in-memory (left) and on-disk (right) backends for 1, 10, 100 1000, 5000 and all 6103 spectra. Benchmarks were performed on a Dell XPS laptop with an Intel i5-8250U processor 1.60 GHz (4 cores, 8 threads), 7.5 GB RAM running Ubuntu 18.04.4 LTS 64-bit and SSD drive.

on subsets of the data become much faster since data manipulations are applied only to data subsets instead of the full data set at once. Also, data access in on-disk backends is parallelized by data file ensuring a higher performance of this backend over conventional in-memory data representations. As an example, the following short analysis pipeline, that can equally be applied to in-memory or on-disk data, retains MS2 spectra acquired between 1000 and 3000 seconds, extracts the m/z range corresponding to the TMT 6-plex range and focuses on the MS2 spectra with a precursor intensity greater than 11×10^6 (the median precursor intensity).

```
ms <- ms %>%
    filterRt(c(1000, 3000)) %>%
    filterMz(120, 135)
ms[precursorIntensity(ms) > 11e6, ]
```

As shown on Figure 2 (c), this lazy mechanism is significantly faster than its application on in-memory data. The advantageous reading and execution times and memory footprint of the on-disk backend are possible by retrieving only spectra data from the selected subset hence avoiding access to the full raw data. Once access to the spectra m/z and intensity values become mandatory (for example for plotting), then the in-memory backend becomes more efficient, as illustrated on Figure 2 (d). This gain is maximal when the whole dataset is accessed (i.e. all spectra are already in memory) and negligible when only subsets of the data are requested.

Prototyping

The MSnExp data structure and its interface constitute an efficient prototyping environment for computational method development. We illustrate this by demonstrating how to implement the BoxCar¹⁴ acquisition method. In a nutshell, BoxCar acquisition aims at improving the detection of intact precursor ions by distributing the charge capacity over multiple nar-

row m/z segments and thus limiting the proportion of highly abundant precursors in each segment. A full scan is reconstructed by combining the respective adjacent segments of the BoxCar acquisitions. The MSnbaseBoxCar package ¹⁵ is a small package that demonstrates this. The simple method is composed of three steps is described below and illustrated with code from MSnbaseBoxCar in the following code chunk.

- 1. Identify and filter the groups of spectra that represent adjacent BoxCar acquisitions (Figure 3 (b)). This can be done using the *filterString* metadata variable that identifies BoxCar spectra by their adjacent m/z segments with the bc_groups() function and filtering relevant spectra with the filterBoxCar().
- 2. Remove any signal outside the BoxCar segments using the bc_zero_out_box() function from MSnbaseBoxCar (Figures 3 (c) and (d)).
- 3. Using the combineSpectra function from the MSnbase, combine the cleaned BoxCar spectra into a new, full spectrum (Figure 3 (e)).

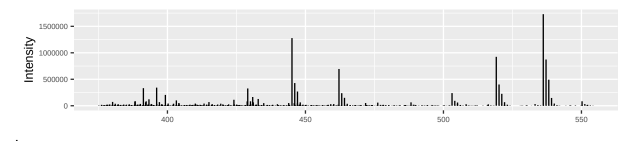
After processing of the BoxCar data, the final object can either be further analysed in R using MSnbase or written back to disk as an mzML file using writeMSData() for processing with other tools.

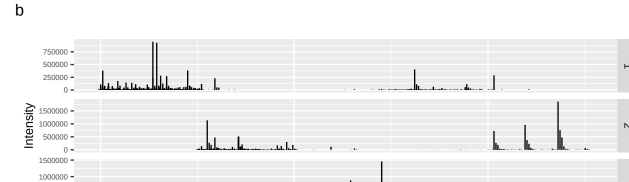
All the functions for the processing of BoxCar spectra and segments in MSnbaseBoxCar were developed using existing functionality implemented in MSnbase, illustrating the flexibil-



500000 -

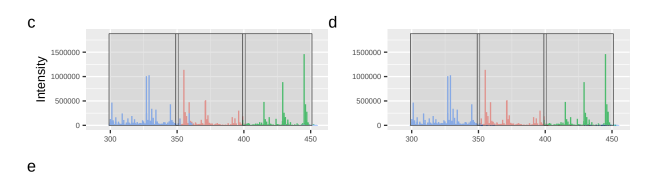
300





ω

500



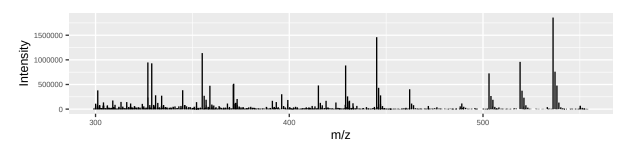


Figure 3: BoxCar processing with MSnbase. (a) Standard full scan with (b) three corresponding BoxCar scans showing the adjacent segments. Figure (c) shows the overlapping intact BoxCar segments and (d) the same segments after cleaning, i.e. where peaks outside of the segments were removed. The reconstructed full scan is shown on panel (e). Spectra visualisation, as shown here, rely on the ggplot2¹⁶ package.

ity and adaptability of the MSnbase package for computational mass spectrometry method development.

Visualisation

The R environment is well known for the quality of its visualisation capacity. This also holds true for mass spectrometry. ^{17–20} Here, we conclude the overview of version 2 of the MSnbase package by highlighting the flexibility of the software to visualise and assess the efficiency of raw data processing. Figure 4 compares the raw MS profile data imported from an mzML file for the serine and the same data after smoothing, centroiding and m/z refinement, as illustrated in the code chunk below. Detailed execution and description of these operations can be found in the MSnbase: centroiding of profile-mode MS data MSnbase vignette.

```
serine_mz <- 106.049871
serine_proc <- ms %>%
    smooth(method = "SavitzkyGolay", halfWindowSize = 4L) %>%
    pickPeaks(refineMz = "descendPeak") %>%
    filterMz(c(serine_mz - 0.01, serine_mz + 0.01)) %>%
    filterRt(c(175, 187))
```

Package maintenance and governance

The first public commit to the MSnbase GitHub repository was in October 2010. Since then, the package benefited from 12 contributors²¹ that added various features, some particularly significant ones such as the on-disk backend described herein. Contributions to the package are explicitly encouraged, rewarded by an official contributor status and coverned by a code of conduct.

According to MSnbase's Bioconductor page, there are 36 Bioconductor packages that depend, import or suggest it. Among these are pRoloc²² to analyse mass spectrometry-

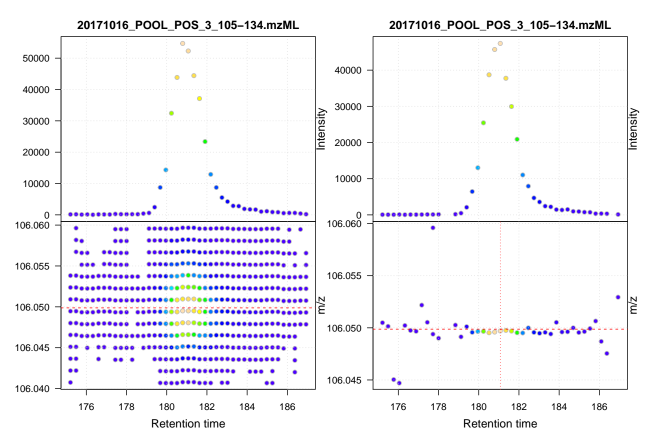


Figure 4: Visualisation of data smoothing and m/z refinement using MSnbase. (a) Raw MS profile data for serine. Upper panel shows the base peak chromatogram (BPC), lower panel the individual signals in the retention time -m/z space. The horizontal dashed red line indicates the theoretical m/z of the [M+H]+ adduct of serine. (b) Smoothed and centroided data for serine with m/z refinement. The horizontal red dashed line indicates the theoretical m/z for the [M+H]+ ion and the vertical red dotted line the position of the maximum signal.

based spatial proteomics data, msmsTests, ²³ DEP, ²⁴ DAPAR and ProStaR ²⁵ for the statistical analysis for of quantitative proteomics data, RMassBank ²⁶ to process metabolomics tandem MS files and build MassBank records, MSstatsQC ²⁷ for longitudinal system suitability monitoring and quality control of targeted proteomic experiments and the widely used xcms ²⁸ package for the processing and analysis of metabolomics data. MSnbase is also used in non-R/Bioconductor software, such as for example IsoProt, ²⁹ that provides a reproducible workflow for iTRAQ/TMT experiments. MSnbase currently ranks 101 out of 1823 packages based on the monthly downloads from unique IP addresses, tallying over 1000 downloads from unique IP addresses each months.

As is custom with Bioconductor packages, MSnbase comes with ample documentation. Every user-accessible function is documented in a dedicated manual page. In addition, the package offers 5 vignettes, including one aimed at developers. The package is checked nightly on the Bioconductor servers: it implements unit tests covering 72% of the code base and, through its vignettes, also provides integration testing. Questions from users and developers are answered on the Bioconductor support forum as well as on the package GitHub page. The package provides several sample and benchmarking datasets, and relies on other dedicated experiment packages such as msdata³⁰ for raw data or pRolocdata²² for quantitative data. MSnbase is available on Windows, Mac OS and Linux under the open source Artistic 2.0 license and easily installable using standard installation procedures.

The growth of MSnbase and the user support provided over the years attest to the core maintainers commitment to long-term development, and the quality and maintainability of the code base.

Discussion

We have presented here some important functionality of MSnbase version 2. The new on-disk infrastructure enables large scale data analyses, 31 either using MSnbase directly or through

packages that rely on it, such as xcms. We have also illustrated how MSnbase can be used for standard data analysis and visualisation, and how it can be used for method development and prototyping.

The version of MSnbase used in this manuscript is 2.14.2. The main features presented here were available since version 2.0. The code to reproduce the analyses and figures in this article is available at https://github.com/lgatto/2020-msnbase-v2/.

Acknowledgement

The authors thank the various contributors and users who have provided constructive input and feedback that have helped, over the years, the improvement of the package. The authors declare no conflict of interest.

References

- (1) Côté Richard G, M. L., Reisinger Florian jmzML, an open-source Java API for mzML, the PSI standard for MS data. *Proteomics* **2010**, *10*, 1332–1335.
- (2) Chambers, M. C. et al. A cross-platform toolkit for mass spectrometry and proteomics.

 Nat Biotechnol 2012, 30, 918–20.
- (3) Cox, J.; Mann, M. MaxQuant enables high peptide identification rates, individualized p.p.b.-range mass accuracies and proteome-wide protein quantification. *Nat Biotechnol* **2008**, *26*, 1367–1372.
- (4) Vaudel, M.; Burkhart, J. M.; Zahedi, R. P.; Oveland, E.; Berven, F. S.; Sickmann, A.; Martens, L.; Barsnes, H. PeptideShaker enables reanalysis of MS-derived proteomics data sets. *Nat Biotechnol* 2015, 33, 22–24.
- (5) Gatto, L.; Lilley, K. S. MSnbase an R/Bioconductor package for isobaric tagged mass

- spectrometry data visualization, processing and quantitation. *Bioinformatics* **2012**, *28*, 288–9.
- (6) Huber, W. et al. Orchestrating high-throughput genomic analysis with Bioconductor.

 Nat Methods 2015, 12, 115–21.
- (7) Rost, H. L.; Schmitt, U.; Aebersold, R.; Lars, M. pyOpenMS: a Python-based interface to the OpenMS mass-spectrometry algorithm library. *Proteomics* **2014**, *14*, 74–77.
- (8) Bittremieux, W. spectrum_utils: A Python package for mass spectrometry data processing and visualization. *Analytical Chemistry* **2020**, *92*, 659–661.
- (9) Goloborodko, A. A.; Levitsky, L. I.; Ivanov, M. V.; Gorshkov, M. V. Pyteomics a Python Framework for Exploratory Data Analysis and Rapid Software Prototyping in Proteomics. J. Am. Soc. Mass Spectrom. 2013, 24, 301–304.
- (10) R Core Team, R: A Language and Environment for Statistical Computing. R Foundation for Statistical Computing: Vienna, Austria, 2020.
- (11) Stanstrup, J. et al. The metaRbolomics Toolbox in Bioconductor and Beyond. *Metabolites* **2019**, *9*, 200.
- (12) Gatto, L. MSnbase Base Functions and Classes For Mass Spectrometry and Proteomics. 2020; R package version 2.14.2.
- (13) Rainer, J. Metabolomics data pre-processing using xcms. 2020.
- (14) Meier, F.; Geyer, P.; Virreira Winter, S.; Juergen, C.; Matthias, M. BoxCar acquisition method enables single-shot proteomics at a depth of 10,000 proteins in 100 minutes. Nature Methods 2018, 15, 440-448.
- (15) Gatto, L. MSnbaseBoxCar: BoxCar Data Processing with MSnbase. 2020; R package version 0.2.0.

- (16) Wickham, H. ggplot2: Elegant Graphics for Data Analysis; Springer-Verlag New York, 2016.
- (17) Gatto, L.; Breckels, L. M.; Naake, T.; Gibb, S. Visualization of proteomics data using R and Bioconductor. *PROTEOMICS* **2015**, *15*, 1375–1389.
- (18) Panse, C.; Grossmann, J. protViz: Visualizing and Analyzing Mass Spectrometry Related Data in Proteomics. 2020; R package version 0.6.
- (19) Sauteraud, R.; Jiang, M.; Gottardo, R. Pviz: Peptide Annotation and Data Visualization using Gviz. 2019; R package version 1.21.0.
- (20) Bemis, K. D.; Harry, A.; Eberlin, L. S.; Ferreira, C.; van de Ven, S. M.; Mallick, P.; Stolowitz, M.; Vitek, O. Cardinal: an R package for statistical analysis of mass spectrometry-based imaging experiments. *Bioinformatics* **2015**,
- (21) Gatto, L. MSnbase contributors 2010 2020. 2020; https://lgatto.github.io/msnbase-contribs-2/.
- (22) Gatto, L.; Breckels, L. M.; Wieczorek, S.; Burger, T.; Lilley, K. S. Mass-spectrometry-based spatial proteomics data analysis using pRoloc and pRolocdata. *Bioinformatics* **2014**, 30, 1322–4.
- (23) Gregori, J.; Sanchez, A.; Villanueva, J. msmsTests: LC-MS/MS Differential Expression Tests. 2019; R package version 1.25.0.
- (24) Zhang, X.; Smits, A. H.; van Tilburg, G. B.; Ovaa, H.; Huber, W.; Vermeulen, M. Proteome-wide identification of ubiquitin interactions using UbIA-MS. *Nature Protocols* **2018**, *13*, 530–550.
- (25) Wieczorek, S.; Combes, F.; Lazar, C.; Giai Gianetto, Q.; Gatto, L.; Dorffer, A.; Hesse, A. M.; Couté, Y.; Ferro, M.; Bruley, C.; Burger, T. DAPAR & ProStaR: software to

- perform statistical analyses in quantitative discovery proteomics. *Bioinformatics* **2017**, 33, 135–136.
- (26) Stravs, M. A.; Schymanski, E. L.; Singer, H. P.; Hollender, J. Automatic Recalibration and Processing of Tandem Mass Spectra using Formula Annotation. *Journal of Mass Spectrometry* 2013, 48, 89–99.
- (27) Dogu, E.; Mohammad-Taheri, S.; Abbatiello, S. E.; Bereman, M. S.; MacLean, B.; Schilling, B.; Vitek, O. MSstatsQC: Longitudinal System Suitability Monitoring and Quality Control for Targeted Proteomic Experiments. Mol Cell Proteomics 2017, 16, 1335–1347.
- (28) Smith, C. A.; Want, E. J.; O'Maille, G.; Abagyan, R.; Siuzdak, G. XCMS: processing mass spectrometry data for metabolite profiling using nonlinear peak alignment, matching, and identification. *Anal Chem* **2006**, *78*, 779–87.
- (29) Griss, J.; Vinterhalter, G.; Schwämmle, V. IsoProt: A Complete and Reproducible Workflow To Analyze iTRAQ/TMT Experiments. J Proteome Res 2019, 18, 1751– 1759.
- (30) Neumann, S.; Gatto, L.; Rainer, J. msdata: Various Mass Spectrometry raw data example files. 2019; R package version 0.27.0.
- (31) Nothias, L. F. et al. Feature-based Molecular Networking in the GNPS Analysis Environment. bioRxiv 2019,