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1 Introduction

This document describes handling of mass spectrometry data from large experiments using the MSnbase package and more specifically its *on-disk* backend. For demonstration purposes, the MassIVE data set MSV000080030 is used. This consists of over 1,000 mzXML files from swab-samples collected from hands and various personal objects of 80 volunteers.

2 Data handling and analysis with MSnbase

In this section we demonstrate data handling and access by MSnbase on a large experiment consisting of more than 1,000 data files.

To reproduce the analysis in this document, download the MSV000080030 folder from ftp://massive.ucsd.edu/MSV000080030/ and place it into the same folder than this document.

Below we load the required libraries and define the files to be analyzed.

The data set consists of in total 1182 mzXML files. We next load the data using the two different MSnbase backends "inMemory" and "onDisk". For the in-memory backend, due to the larger memory requirements, we import the data only from a subset of the files.

```
ms_mem <- readMSData(fls[grep("Hand", fls)], mode = "inMemory")</pre>
```

Next we load data from all mzXML files as an on-disk MSnExp object.

```
ms_dsk <- readMSData(fls, mode = "onDisk")</pre>
```

Below we count the number of spectra per MS level of the whole experiment.

```
table(msLevel(ms_dsk))
##
## 1 2
## 1173678 4599786
```

Note that the in-memory MSnExp object contains only MS2 spectra (in total 2140520) from a subset of data files, still, data import was much slower (over \sim 12 hours for the in-memory backend while creating the on-disk object from the full data data set took \sim 3 hours).

Next we subset the on-disk object to contain the same set of spectra than the in-memory MSnExp and compare their memory footprint.

```
ms_dsk_hands <- ms_dsk %>%
    filterFile(grep("Hand", fls)) %>%
    filterMsLevel(2L)

object_size(ms_mem)
```

```
## 21.8 GB
object_size(ms_dsk_hands)
## 617 MB
```

Since the on-disk object stores only spectra metadata in memory it occupies also much less system memory. As a comparison, the on-disk MSnExp for the full experiment was still much smaller than the in-memory object:

```
object_size(ms_dsk)
## 1.66 GB
```

2.1 Performance of the on-disk backend on large scale data sets

To demonstrate MSnbase's efficiency in processing large scale experiments we perform some standard subsetting, data access and manipulation operations.

We first compare the performance of the on-disk and in-memory backend on accessing m/z values with the mz function on a set of 100 randomly selected spectra. The performance is assessed with the microbenchmark function.

```
set.seed(123)
idx <- sample(seq_along(ms_mem), 100)</pre>
library(microbenchmark)
microbenchmark(mz(ms_mem[idx]),
               mz(ms_dsk_hands[idx]),
               times = 5)
## Unit: seconds
##
                     expr
                                min
                                            lq
                                                  mean
                                                           median
                                                                         uq
         mz(ms_mem[idx]) 51.514084 55.542641 59.32276 55.586931 60.34142 73.62871
##
## mz(ms_dsk_hands[idx]) 3.892099 3.893295 13.41512 3.976054 26.77394 28.54019
    neval
##
       5
        5
##
```

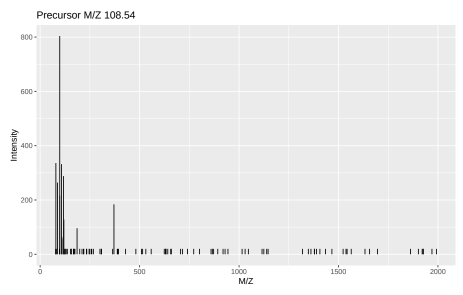
Thus, for this combined subsetting and data access operation the on-disk backend performed better than the in-memory MSnExp, while even requiring much less memory.

Next we extract all MS2 spectra with a retention time between 50 and 60 seconds and a precursor m/z of 108.5362 (+/- 5ppm). This subsetting operation is performed on the on-disk MSnExp object representing the full experiment with the 1182 data files/samples. To assess the performance of the following operations we use system.time calls that record elapsed time in seconds.

```
system.time(
    ms_sub <- ms_dsk %>%
        filterMsLevel(2L) %>%
        filterRt(c(50, 60)) %>%
        filterPrecursorMz(mz = 108.5362, ppm = 5)
)["elapsed"]
## elapsed
## 6.698
```

In total <code>length(ms_sub)</code> spectra were selected from in total 928 data files/samples. The plot below shows the data for the first spectrum.

```
system.time(
    plot(ms_sub[[1]])
)["elapsed"]
```



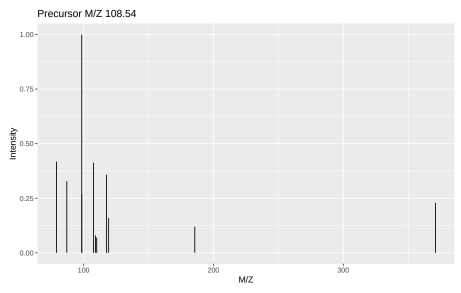
```
## elapsed
## 0.354
```

Since there seems to be quite some background noise in the MS2 spectrum we next remove peaks with an intensity below 50 by first replacing their intensities with 0 (with the remove Peaks call) and subsequently removing all 0-intensity peaks from each spectrum with the clean call. In addition we *normalize* each spectrum by dividing the maximum intensity per spectrum from the spectrum's intensities.

```
system.time(
    ms_sub <- ms_sub %>%
    removePeaks(t = 50) %>%
    clean(all = TRUE) %>%
    normalize(method = "max")
)["elapsed"]
## elapsed
## 0.043
```

The result on the first spectrum is shown below.

```
system.time(
    plot(ms_sub[[1]])
)["elapsed"]
```



```
## elapsed
## 0.316
```

Note that any of the data manipulations above are not directly applied to the data but *cached* in the object's internal *lazy processing queue* (explaining the very short running time of the normalization call). The operations are only effectively applied to the data when m/z or intensity values are extracted from the object as it is the case in the plot call above.

For additional workflows employing MSnbase see also metabolomics2018¹ that explains filtering, plotting and centroiding of profile-mode MS data with MSnbase and subsequent pre-processing of the (label free/untargeted) LC-MS data with the xcms package (that builds upon MSnbase for MS data representation and access).

¹https://github. com/jorainer/ metabolomics2018

2.2 System information

The present analysis was run on a MacBook Pro 16,1 with 2.3 GHz 8-Core Intel Core i9 CPU and 64 GB 2667 MHz DDR4 memory running macOS version 10.15.5. The R version and the version of the used packages are listed below.

```
sessionInfo()
## R version 4.0.0 (2020-04-24)
## Platform: x86_64-apple-darwin17.0 (64-bit)
## Running under: macOS Catalina 10.15.5
##
## Matrix products: default
## BLAS: /Library/Frameworks/R.framework/Versions/4.0/Resources/lib/libRblas.dylib
## LAPACK: /Library/Frameworks/R.framework/Versions/4.0/Resources/lib/libRlapack.dylib
##
## locale:
## [1] en_US.UTF-8/en_US.UTF-8/en_US.UTF-8/c/en_US.UTF-8/en_US.UTF-8
##
## attached base packages:
## [1] stats4 parallel stats graphics grDevices utils datasets
```

```
## [8] methods
                base
##
## other attached packages:
## [1] microbenchmark_1.4-7 BiocParallel_1.22.0 pryr_0.1.4
## [4] magrittr_1.5
                           MSnbase_2.14.2
                                                ProtGenerics_1.20.0
## [7] S4Vectors_0.26.1
                            mzR_2.22.0
                                                Rcpp_1.0.4.6
## [10] Biobase_2.48.0
                            BiocGenerics_0.34.0 BiocStyle_2.16.0
## [13] rmarkdown_2.3
##
## loaded via a namespace (and not attached):
## [1] tinytex_0.24 tidyselect_1.1.0
                                                  xfun_0.15
## [4] purrr_0.3.4
                           lattice_0.20-41
                                                  colorspace_1.4-1
## [7] vctrs_0.3.1
                             generics_0.0.2
                                                  htmltools_0.5.0
## [10] yaml_2.2.1
                             vsn_3.56.0
                                                  XML_3.99-0.3
## [13] rlang_0.4.6
                             pillar_1.4.4
                                                  glue_1.4.1
## [16] affy_1.66.0
                             foreach_1.5.0
                                                  affyio_1.58.0
## [19] lifecycle_0.2.0
                             plyr_1.8.6
                                                  mzID_1.26.0
## [22] stringr_1.4.0
                             zlibbioc_1.34.0
                                                  munsell_0.5.0
## [25] pcaMethods_1.80.0
                             gtable_0.3.0
                                                  codetools\_0.2-16
## [28] evaluate_0.14
                             labeling_0.3
                                                  knitr_1.29
## [31] IRanges_2.22.2
                             doParallel_1.0.15
                                                  preprocessCore_1.50.0
## [34] scales_1.1.1
                             BiocManager_1.30.10
                                                  limma_3.44.3
## [37] farver_2.0.3
                             impute_1.62.0
                                                  ggplot2_3.3.2
## [40] digest_0.6.25
                             stringi_1.4.6
                                                  bookdown_0.20
## [43] dplyr_1.0.0
                             ncdf4_1.17
                                                  grid_4.0.0
## [46] tools_4.0.0
                             tibble_3.0.1
                                                  crayon_1.3.4
## [49] pkgconfig_2.0.3
                             ellipsis_0.3.1
                                                  MASS_7.3-51.6
## [52] iterators_1.0.12
                             R6_2.4.1
                                                  MALDIquant_1.19.3
## [55] compiler_4.0.0
```