# Processing Tandem-MS and $MS^n$ data with xcms

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## Introduction

This document describes how to use xcms for processing of Tandem-MS and MS<sup>n</sup> data from e.g. triple-quad, QTOF, ion trap or orbitrap mass spectrometers.

> library(xcms)

> library(msdata)

### 1 Raw Data File Preparation

The *xcms* package reads full-scan LC/MS data and associated Tandem-MS and  $MS^n$  spectra from mzData and mzXML files (CDF does not support  $MS^n$ ).

For the purposes of demonstration, we will use a some sample files included in the *msdata* package. The (truncated) raw data files are contained in the **data** directory. To access the mzData files, we first locate the *data* directory in the *msdata* package.

```
> mzdatapath <- system.file("iontrap", package = "msdata")
> list.files(mzdatapath, recursive = TRUE)
```

[1] "extracted.mzData"

## 2 Accessing, combining and Visualising $MS^n$ spectra

The  $MS^n$  spectra are read into xcms just like plain LC-MS files. Inclusion of the additional scans has to be explicitly enabled. The xcmsRaw summary shows the additional content:

```
> library(xcms)
> mzdatafiles <- list.files(mzdatapath, pattern="extracted.mzData", recursive = TRUE,
> xraw <- xcmsRaw(mzdatafiles[1], includeMSn=TRUE)
> xraw
```

There is also a new peak picker, which actually returns all parent ions as annotated in the mzData / mzXML files. A warning will be issued if this is not present, either because of a poor converter or if xraw is read from a CDF file:

```
> peaks <- findPeaks(xraw, method="MS1")</pre>
```

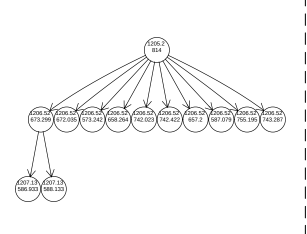
By giving larger windows, subsets of spectra can be returned, and these can be combined to obtain so called compound spectra containing e.g.  $MS^2$  to  $MS^5$ .

#### 3 xcmsFragments

A xcmsFragments-Object contains a Table with informations about all Peaks from MS1 to  $MS^n$  of one Experiment, including mz-value, retentiontime, MSlevel, and parentpeak. The data can be used for visualization or (later) computing alignments between different Experiments. The object is created as empty and can be filled with collect(). This method requires a peaktable or a xcmsSet for the MS1peaks and the original xcmsRaw for the msn-Data.

```
> xs <- xcmsSet(mzdatafiles, method="MS1")</pre>
```

> xfrag <- xcmsFragments(xs)
394 Peaks picked, 0 MSn-Specs ignored.
> xfrag



|##6\_M814T1205.2F814 | ##75\_M814T1206.52F673.2994 | ##85\_M814T1207.13F586.9333 | ##86\_M814T1207.13F588.1333 | ##76\_M814T1206.52F672.0354 | ##77\_M814T1206.52F573.2417 | ##78\_M814T1206.52F658.2642 | ##79\_M814T1206.52F742.0226 | ##80\_M814T1206.52F742.4218 | ##81\_M814T1206.52F657.1997 | ##81\_M814T1206.52F587.0795 | ##83\_M814T1206.52F755.1951 | ##84\_M814T1206.52F743.2867

Figure 1: The  $MS^n$  fragmentation tree for the MS1 peak 6.

An "xcmsFragments" object with 394 peaks in 77 Spectra From Level 1 to 3 Number of Samples: 1 .

Sample 1 : 37 Peaks in Level 1 351 Peaks in Level 2 6 Peaks in Level 3

Memory usage: 0.0289 MB

plotTree prints out a simplyfied tree of the peaks in a xcmsFragments. The user can define borders for the retentionTime and for the MassRange, he can also print the tree for a single MS1-Peak. PlotTree uses the package RgraphViz and plots the tree text-based if requested or if the package cannot be found.

> plotTree(xfrag,xcmsFragmentPeakID=6)