

HowTo Use the Bioconductor PROcess package

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1 Introduction

The `PROcess` package contains a collection of functions for processing spectra to remove baseline drifts if any, detect peaks and align them to a set of protobiomarkers. This document serves as a quick tutorial for using the `PROcess` package.

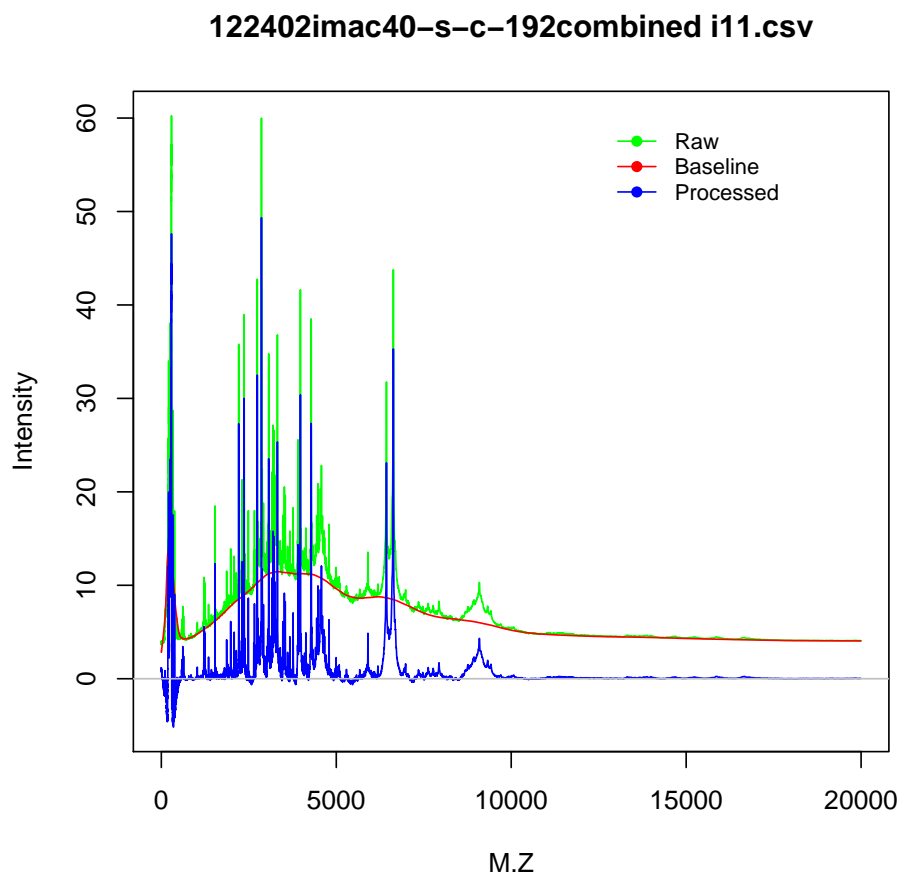
2 Baseline subtraction

Our first observation of a raw spectrum is that it exhibits elevated baseline, more so at smaller m/z values than at larger values. This elevated baseline is mostly caused by the chemical noises in the EAM and ion overload. Ideally a spectrum should rest more or less on the zero horizontal line. This baseline needs to be subtracted from each raw spectrum. The following example shows the result of a spectrum with its baseline removed.

```
> library(PROcess)
```

Loading required package: Icms

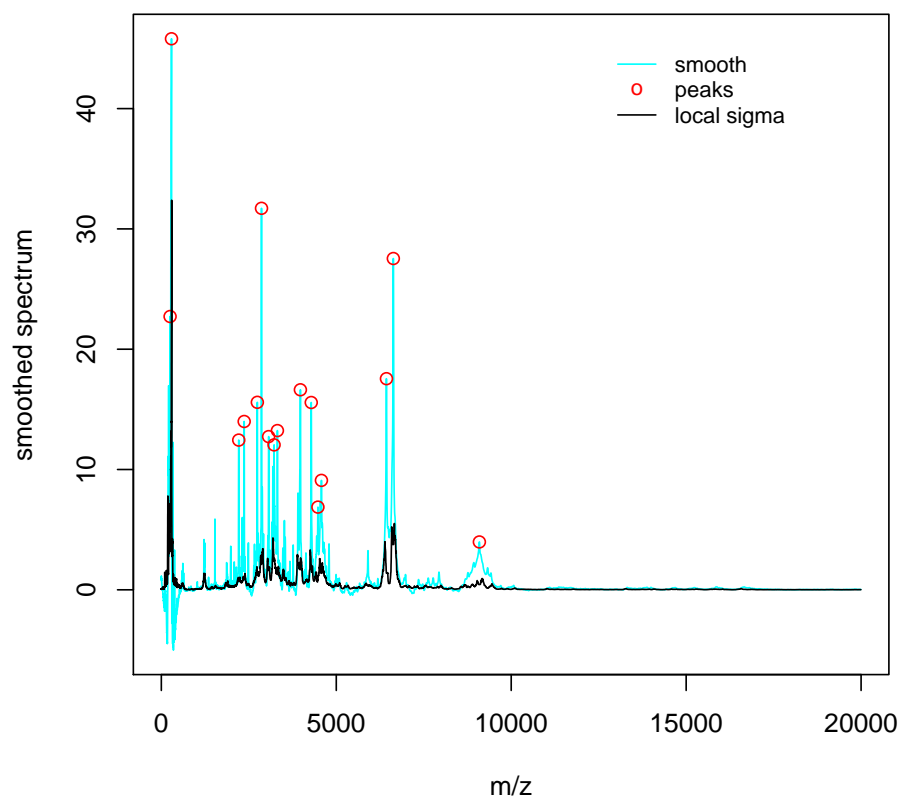
```
> fdat <- system.file("Test", package = "PROcess")
> fs <- list.files(fdat, pattern = "*.csv", full.names = TRUE)
> f1 <- read.files(fs[1])
> fcut <- f1[f1[, 1] > 0, ]
> bseoff <- bslnoff(fcut, method = "loess", plot = TRUE, bw = 0.1)
> title(basename(fs[1]))
```



3 Peak detection

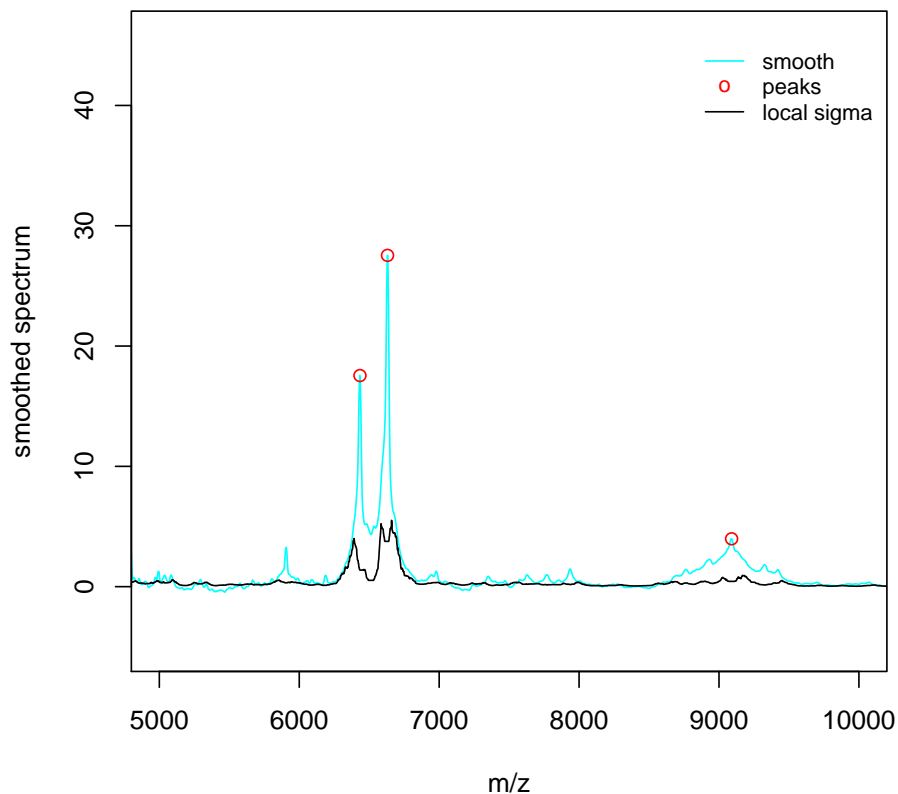
After baseline is removed, peaks can be located by using `isPeak`. A spectrum is smoothed first using moving average of the k nearest neighbours. Smoothing helps to enhance peaks and get rid of spurious peaks. However, we do not recommend large amount of smoothing (controlled by parameter `sm.span`) in this step because we do not wish to smooth away too many short and wide peaks and also we need the precision in peak locations. As a first step we do not mind getting more potential features.

```
> pkgobj <- isPeak(bseoff, span = 81, sm.span = 11, plot = TRUE)
```



We can also zoom in to inspect peaks in a particular range of m/z values.

```
> specZoom(pkgobj, xlim = c(5000, 10000))
```



4 Batch operation

We demonstrate the batch functionality of this package using a set of 2 spectra.

4.1 Apply baseline subtraction to a set of spectra

```
> testdir <- system.file("Test", package = "PROcess")
> testM <- rmBaseline(testdir)
```

4.2 Renormalize spectra

Suppose we want to normalize a set of spectra to their median AUC (Area Under the Curve), where an AUC is calculated for m/z values greater than a cutoff point, 1500.

```
> rtM <- renorm(testM, cutoff = 1500)
```

4.3 Identify peaks of spectra

```
> peakfile <- paste(tempdir(), "testpeakinfo.csv", sep = "/")
> getPeaks(testM, peakfile)
```

4.4 Quality assessment

Quality assessment is necessary because some spectra are very noisy and have hardly any peaks. Function `quality` computes three parameters `Quality`, `Retain` and `peak` for assessing a set of spectra.

```
> qualRes <- quality(testM, peakfile, cutoff = 1500)
> print(qualRes)
```

	Quality	Retain	peak
122402imac40-s-c-192combined i11.csv	0.5419809	0.3825606	1.28
122402imac40-s-c-192combined i12.csv	0.6234699	0.3548255	0.72

A spectrum is deemed of poor quality and should be removed from subsequent analyses if it meets the following 3 conditions simultaneously:

1. `Quality < 0.4`;
2. `Retain < 0.1`;
3. `peak < 1/2` of the mean peak number in the chip.

4.5 Get protobiomarkers

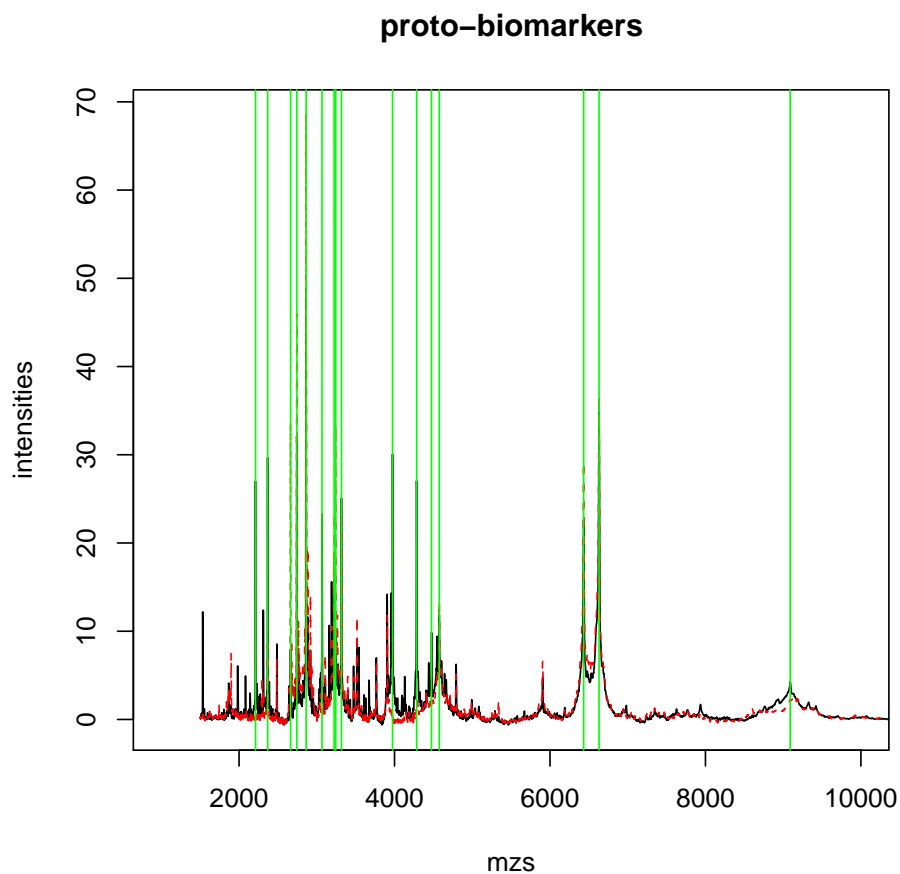
One challenge in MS data is that not only they exhibit variation vertically but also do they horizontally. This horizontal variation is not simply a constant shift but associated with value of m/z . Currently the accuracy in the m/z position is believed to be within 0.3% of the m/z value. Once the peaks are detected, we align the peaks by first generating an interval of size 0.3% of the m/z value which centers at m/z for each m/z where a peak is detected. We treat those m/z intervals as interval censored data. Maximum likelihood estimate of the distribution of survival times conditional upon the observed intervals can be computed by the method of Gentleman and Geyer (1994). Protobiomarkers are computed as the center of the MLE intervals. For a spectrum whose peaks are in the clique that define a protobiomarker, its value at this protobiomarker is set to be the maximum of the peaks in the clique, otherwise its value is set to be the value of its nearest neighbour.

```
> bmfile <- paste(tempdir(), "testbiomarker.csv", sep = "/")
> testBio <- pk2bmkr(peakfile, testM, bmfile)
> mzs <- as.numeric(rownames(rtM))
```

```

> matplot(mzs, rtM, type = "l", xlim = c(1000, 10000), ylab = "intensities",
+         main = "proto-biomarkers")
> bks <- getMzs(testBio)
> abline(v = bks, col = "green")

```



References

Robert Gentleman and Charles J. Geyer. Maximum likelihood for interval censored data: Consistency and computation. *Biometrika*, 81:618–623, 1994.