

## Chapter 11

# Mass Spectrometry Analysis Using MALDIquant<sup>1</sup>

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**Abstract** MALDIquant and associated R packages provide a versatile and completely free open-source platform for analyzing 2D mass spectrometry data as generated for instance by MALDI and SELDI instruments. We first describe the various methods and algorithms available in MALDIquant. Subsequently, we illustrate a typical analysis workflow using MALDIquant by investigating an experimental cancer data set, starting from raw mass spectrometry measurements and ending at multivariate classification.

## 11.1 Introduction

Mass Spectrometry (MS), a high-throughput technology commonly used in proteomics, enables the measurement of the abundance of proteins, metabolites, peptides and amino acids in biological samples. The study of changes in protein expression across subgroups of samples and through time provides valuable insights into cellular mechanisms and offers a means to identify relevant biomarkers, e.g. to distinguish among tissue types, or for predicting health status. In practice, however, there still remain many analytic and computational challenges to be addressed, especially in clinical diagnostics (Leichtle et al., 2013). Among these challenges the availability of open and easy-to-extend processing and analysis software is highly important (Aebersold and Mann, 2003; Lilley et al., 2011).

Here, we present MALDIquant (Gibb and Strimmer, 2012), a complete open-source analysis pipeline for the R platform (R Core Team, 2015). In the first half of this chapter we describe the methodology implemented and available in MALDIquant. In the second half we illustrate the versatility of MALDIquant by application to an experimental data set, showing, how raw intensity measurements are preprocessed and how peaks relevant for a specific outcome can be identified.

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Current documentation of specific version of MALDIquant can be found on its homepage at <http://strimmerlab.org/software/malDIquant/> where we also provide instructions for installing the software. In addition, we provide a number of example R scripts on the MALDIquant homepage. Direct download of the MALDIquant software is also possible from the CRAN server at <http://cran.r-project.org/package=MALDIquant>.

## 11.2 Methodology Available in MALDIquant

### 11.2.1 General Workflow

The purpose of MALDIquant is to provide a complete workflow to facilitate the complex preprocessing tasks needed to convert raw two-dimensional MS data, as generated for example by MALDI or SELDI instruments, into a matrix of feature intensities required for high-level analysis. A typical workflow is depicted in figure 11.1.

Each analysis with MALDIquant consists of all or some of the following steps (see also Norris et al. (2007); Morris et al. (2010) for related analysis pipelines): First, the raw data is imported into the *R* environment. Subsequently, the data are smoothed to remove noise and also transformed for variance stabilization. Next, to remove chemical background noise a baseline correction is applied. This is followed by a calibration step to allow comparison of intensity values across different baseline-corrected spectra. As a next step a peak detection algorithm is employed to identify potential features and also to reduce the dimensionality of the data. After peaks have been identified a peak alignment procedure is applied as the *mass-to-charge ratios* ( $m/z$ s) typically differ across different measurements and need to be adjusted accordingly. Finally, after feature binning an intensity matrix is produced that can be used as starting point for further statistical analysis, for example for variable selection or classification.

In the following subsections we discuss each of these steps in more detail.

### 11.2.2 Import of Raw Data

A prerequisite of any analysis is to import the raw data into the *R* environment. Unfortunately, nearly every vendor of mass spectrometry machinery has its own native and often proprietary data format. This complicates the exchange of experimental data between laboratories, the use of analysis software and the comparison of results. Fortunately, there is now much effort to create generic and open formats, such as *mzXML* (Pedrioli et al., 2004) and its successor *mzML* (Martens et al., 2011) or *imzML* (Schramm et al., 2012) for Mass Spectrometry Imaging (MSI) data. Nevertheless, the support of these formats is still limited and often conversion is needed to get the data into a suitable format for subsequent analysis (Chambers et al., 2012).

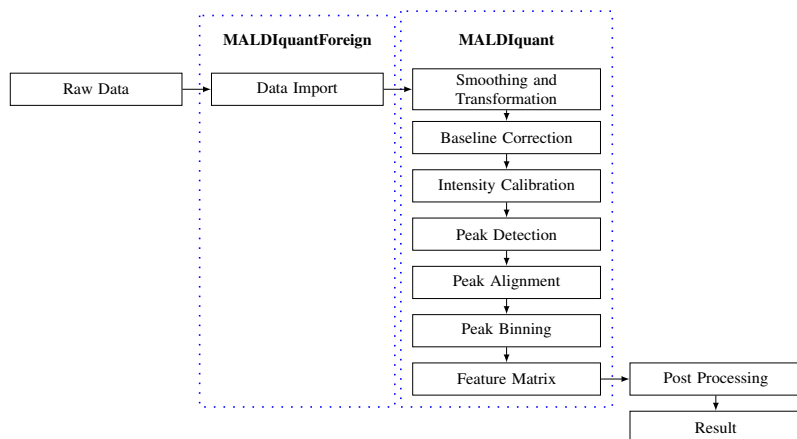


Figure 11.1: Preprocessing workflow for MS data using MALDIquantForeign and MALDIquant.

Importing of raw data in MALDIquant is performed by its sister R package MALDIquantForeign. It offers import routines for numerous native and public data formats. In addition to the open XML formats (*mzXML*, *mzML*) it supports *Ciphergen XML*, *ASCII*, *CSV*, *NetCDF*, and *Bruker DatIonics \*flex Series* files. It can also read MSI formats like *imzML* and *ANALYZE 7.5* (Robb et al., 1989).

A very useful feature of MALDIquantForeign is that it reads and traverses whole directory trees containing supported file formats so that simultaneous import of many spectra is straightforward. Furthermore, MALDIquantForeign allows to import data from remote resources so the spectral data can be read over an Internet connection from a website or database.

After importing the raw spectra an important step is quality control. This includes checking the mass range, the length of each spectra and also visual exploration of spectra to find and remove potentially defective measurements. MALDIquant provides functions to facilitate this often neglected task.

### 11.2.3 Intensity Transformation and Smoothing

The raw data obtained from mass spectrometry experiments are counts of ionized molecules, with intensity values approximately following a *Poisson* distribution (Sköld et al., 2007; Du et al., 2008). Consequently, the variance depends on the mean, as mean and variance are identical for a *Poisson* distribution. However, by applying a square root transformation ( $f(x) = \sqrt{x}$ ) we can convert the Poisson distributed data to approximately normal data, with constant variance independent of mean, which is an important requirement for many statistical tests (Purohit and Rocke, 2003). In the preprocessing noise models other than the Poisson may be also assumed, which

lead to different variance-stabilizing functions such as the logarithmic transformation (Tibshirani et al., 2004; Coombes et al., 2005). These can be easily applied in MALDIquant as well.

Subsequently, the transformed spectral data is smoothed to reduce small and high-frequent variations and noise. For this purpose MALDIquant offers the moving average smoother and the *Savitzky-Golay*-filter (Savitzky and Golay, 1964). The latter is based on polynomial regressions in a moving window. In contrast to the moving average, the Savitzky-Golay filter preserves the shape of the local maxima.

Note that both algorithms require the specification of window size, which according to Bromba and Ziegler (1981) should be chosen to be smaller than twice the Full Width at Half Maximum (FWHM) of the peaks.

### 11.2.4 Baseline Correction

The elevation of the intensity values in a typical Matrix-Assisted Laser Desorption/Ionization – Time-Of-Flight Mass Spectrometer (MALDI-TOF) spectrum is called *baseline* and is caused by chemical noise such as matrix-effects and pollution. It is recommended to remove these background effects to reduce their influence in quantification of the peak intensities.

In the last few years many algorithms to adjust for the baseline have been developed, ranging from simple methods like the subtraction of the absolute minimum (Gammerman et al., 2008) or the moving minimum or median (Liu et al., 2010) to more elaborate methods such as fitting a LOWESS curve, a spline or an exponential function against the moving minima respectively median values (Tibshirani et al., 2004; Williams et al., 2005; Li, 2005; Liu et al., 2009; He et al., 2011; House et al., 2011). Other authors prefer morphological filters such as *TopHat* (Sauve and Speed, 2004), iterative methods as the Statistics-sensitive Non-linear Iterative Peak-clipping algorithm (SNIP) (Ryan et al., 1988) or the convex hull approach (Liu et al., 2003).

Unfortunately, there is no automatic way to select among the available procedures to find the baseline correction method that is most suitable for a given spectrum at hand. Instead, it is recommended to investigate multiple baseline estimations by visual inspection (Williams et al., 2005). As shown in Fig. 11.2 the algorithms can indeed differ substantially.

MALDIquant provides three complex baseline correction algorithms that have been selected for inclusion in MALDIquant because of their favorable properties, such as respecting peak form and non-negativity of intensity values:

1. The *convex hull* algorithm (Andrew, 1979) doesn't need a tuning parameter and is often very effective to find the baseline. Unfortunately, for concave matrix effects as common in MALDI-TOF spectra this algorithm cannot be applied — see Fig. 11.2B ( $m/z \approx 1500Da$ ).
2. *TopHat* (van Herk, 1992; Gil and Kimmel, 2002) is a morphological filter combining a moving minimum (erosion filter) followed by a moving maximum (dilation filter). In contrast to the convex hull approach it has an additional

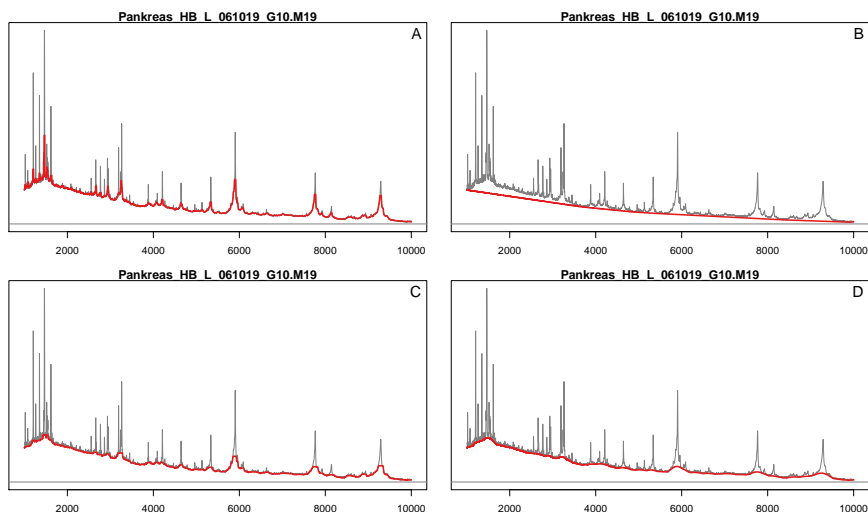


Figure 11.2: Estimated baselines for an raw MALDI-TOF spectrum from Fiedler et al. (2009). The following algorithms were applied: (A) moving median, (B) convex hull, (C) *TopHat*, (D) SNIP.

tuning parameter, the window size of the moving window, that controls the smoothness of the estimated baseline. The narrower the window the more of the baseline is removed but also of the peak heights. A wider window will preserve the peak intensities and produce a smoother baseline but will also cause some local background variation to remain (Fig. 11.2C).

3. The default baseline correction algorithm in MALDIquant is SNIP (Ryan et al., 1988). Essentially, this is a local window-based algorithm in which a baseline is reconstructed by replacing the intensities in a window by the mean of the surrounding points, if the mean is smaller than the local intensity, with window size decreasing iteratively starting from a specified upper limit (Morhác, 2009).

In addition to the above MALDIquant also supports the moving-median algorithm (Fig. 11.2A) which is commonly used in the literature but may lead to negative intensity values after baseline subtraction.

### 11.2.5 Intensity Calibration

The intensity values in mass spectrometry data represent the relative amount of analytes, such as peptides. The measured intensity strongly depends on preanalytical and environmental factors like sample collection, sample storing, room temperature, air humidity, crystallization etc. (Baggerly et al., 2004; Leek et al., 2010; Leichtle et al., 2013).

Further confounders are introduced by so-called batch effects. These are systematic differences that hide the true biological effect and that are caused by different experimental conditions, for instance a different preanalytical processing, measurements on different days by different operators in different laboratories on different devices (Hu et al., 2005; Leek et al., 2010; Gregori et al., 2012).

The systematic errors can be stronger than the real biological effect, and are best minimized already at the stage of data acquisition by strictly adhering to a standardized preanalytical and experimental protocol (Baggerly et al., 2004).

Note that unlike in other omics data, such as gene expression data, batch effects and other systematic errors can be the source of shifts both on the  $x$ -axis ( $m/z$  values) and on the  $y$ -axis (intensity values). Hence, to ensure the validity of any subsequent statistical analysis, great care must be taken to address both of these shift in preprocessing, by *intensity calibration* (often called normalization) and by *peak alignment/warping* (see also 11.2.7).

Methods to calibrate peak intensities can be divided into *local* and *global* approaches (Meuleman et al., 2008). In a *local* calibration each single spectrum is calibrated on its own, by matching a specified characteristic such as the median, the mean or the Total Ion Current (TIC) (Callister et al., 2006; Meuleman et al., 2008; Borgaonkar et al., 2010). In contrast, *global* approaches use information across multiple spectra, e.g. employing linear regression normalization (Callister et al., 2006), quantile normalization (Bolstad et al., 2003), or Probabilistic Quotient Normalization (PQN) (Dieterle et al., 2006).

MALDIquant supports two *local* and one *global* method. Specifically, it implements the TIC and median calibration as well as PQN. In PQN all spectra are calibrated using the TIC calibration first. Subsequently, a median reference spectrum is created and the intensities in all spectra are standardized using the reference spectrum and a spectrum-specific median is calculated for each spectrum. Finally, each spectrum is rescaled by the median of the ratios of its intensity values and that of the reference spectrum (Dieterle et al., 2006).

It has been shown that applying intensity calibration is an essential step in preprocessing (Meuleman et al., 2008). Despite its simplicity TIC is often the best choice, especially to account for effects between technical replicates (Shin and Markey, 2006; Meuleman et al., 2008).

### 11.2.6 Peak Detection

Peak detection is a further step in processing mass spectrometry data, serving both to identify potential relevant features as well as to reduce the dimensionality of the data.

MALDIquant provides the most commonly used peak detection method based on finding local maxima (Yasui et al., 2003a; Tibshirani et al., 2004; Li, 2005; Morris et al., 2005; Smith et al., 2006; Tracy et al., 2008). First a window is moved across the spectra and local maxima are detected. Subsequently these local maxima are compared against a noise baseline which is estimated by the Median Absolute Deviation (MAD) or alternatively Friedman's SuperSmoother (Friedman, 1984). If

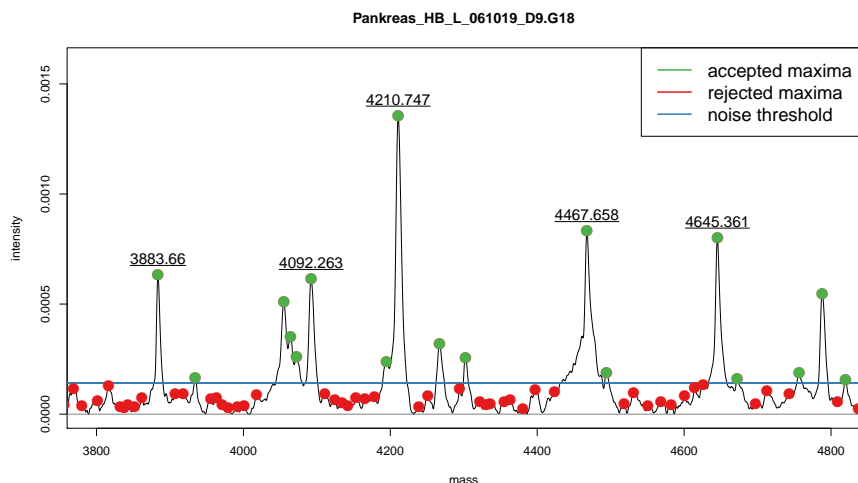


Figure 11.3: Detail view of a MALDI-TOF spectrum from Fiedler et al. (2009). Local maxima are marked with points (red: rejected maxima, green: peaks). The blue line represents the estimated noise baseline as estimated by MAD.

a local maximum is above a given Signal-to-Noise Ratio (SNR) it is considered a peak, whereas local maxima below the SNR threshold are discarded (Fig 11.3).

Some authors advocate peak detection methods based on *wavelets* (Du et al., 2006; Lange et al., 2006). These methods are implemented in the Bioconductor R packages *MassSpecWavelet* and *xcms* (Smith et al., 2006) and thus are readily available if needed.

### 11.2.7 Peak Alignment

As already noted above in Section 11.2.5 not only the intensities but also the  $m/z$  values differ across spectra, as result of the many possible sources of variation in the acquisition of mass spectrometry data. Methods to recalibrate the  $m/z$  values of the spectra are referred to as *peak alignment* or *warping*.

A simple approach is the Correlation Optimized Warping (COW) algorithm (Veselkov et al., 2009; Morris et al., 2010; Wang et al., 2010). COW is based on pairwise comparisons of spectra and maximizes the correlation to find an optimal shift. The advantage of this approach is that correlation is fast to compute and with the use of a reference spectrum the method is also applicable to simultaneous alignment of multiple spectra. However, in actual data the location shifts are typically of a nonlinear nature (He et al., 2011), thus methods based on global linear shifts will often be ineffective in achieving an optimal alignment. A possible workaround is to divide the spectrum into several parts and perform local linear alignment instead.

An alternative, and much more flexible approach, is Dynamic Time Warping (DTW) which is based on dynamic programming (Torgrip et al., 2003; Toppoo et al., 2008; Clifford et al., 2009; Kim et al., 2011). DTW is a pairwise alignment approach that is guaranteed to find the optimal alignment by comparing each point in the first spectrum to every other point in the second spectrum, and optimizing a distance score. Dynamic programming techniques are used to substantially shortcut computational time by means of an underlying decision tree (e.g. Sakoe and Chiba, 1978). Still, DTW is a computationally very expensive algorithm that also requires substantial computer memory, especially for multiple alignment.

As a compromise, recently the Parametric Time Warping (PTW) approach has been suggested (Jeffries, 2005; Lin et al., 2005; Bloemberg et al., 2010; He et al., 2011; Wehrens et al., 2015) where a polynomial functions is used to stretch or shrink a spectrum to increase the similarity between them. PTW is a very fast method that is also able to correct for non-linear shifts. As in all previously mentioned method a reference spectrum is necessary for multiple alignment. Note that the use of a reference spectrum requires prior calibration of the intensity values (Smith et al., 2013).

Finally, another simple strategy to align peaks is based on clustering respectively creation of bins of similar  $m/z$  values (Yasui et al., 2003b; Tibshirani et al., 2004; Tracy et al., 2008). This is a fast and easy to implement approach, and in contrast to DTW, COW and PTW it offers the possibility to align all spectra simultaneously. However, the clustering approach is valid only when there are relatively small shifts around the true peak position, hence this approach is only applicable if there are only mild distortions in the  $m/z$  values.

In MALDIquant we use nonlinear warping of peaks (He et al., 2011; Wehrens et al., 2015). First we align the  $m/z$  values of the peaks using PTW and subsequently we employ binning to identify common peak positions across spectra. Note that in contrast to the standard version of PTW we work on peak level rather than on the whole spectrum.

Our peak alignment algorithm in MALDIquant starts by looking for stable peaks, which are defined as high peaks in defined, coarse  $m/z$  ranges that are present in most spectra. The  $m/z$  of the peaks is averaged and used as reference peak list (also known as anchor or landmark peaks — see Wang et al. (2010)). Next, MALDIquant computes a LOcally WEighted Scatterplot Smoothing (LOWESS) curve or polynomial-based function to warp the peaks of each spectra against the reference peaks (Fig. 11.4). As not all reference peaks are found in each spectrum, the number of matched peaks out of all reference peaks is reported by MALDIquant for information.

Due to using the peaks instead of the whole spectral data the alignment approach implemented in MALDIquant is much faster than traditional PTW, still the results are comparable (Fig. 11.5). Another important advantage of our approach is that only  $m/z$  values are used for calibration, which implies that our approach does not require perfectly calibrated intensity values as is the case for full spectrum-based alignment methods.

After performing alignment, peak positions of identical features across spectra



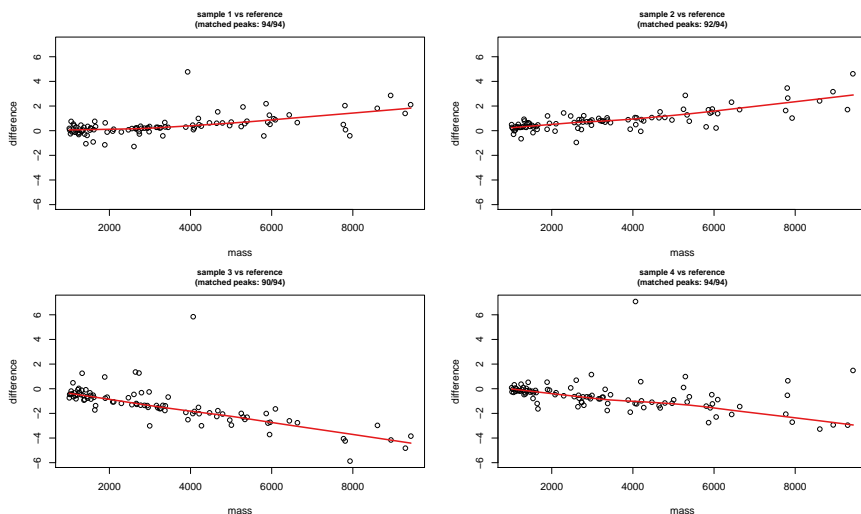


Figure 11.4: Example warping function for four different peak lists. The x-axis represents the peak position and the y-axis the difference from the reference peak list. The red line shows the calculated warping function. The number of matched peaks out of all reference peaks is also shown for each spectrum.

will become very similar but in general not numerically identical. Thus, as final step grouping of  $m/z$  values into bins is needed. For this purpose MALDIquant uses the following simple clustering algorithm: The  $m/z$  values are sorted in ascending order and split recursively at the largest gap until all  $m/z$  values in the resulting bins are from different samples and their individual  $m/z$  values are in a small user-defined tolerance range around their mean. The latter becomes the new  $m/z$  value for all corresponding peaks in the associated bin.

## 11.2.8 Subsequent Statistical Analysis

With peak alignment the task of MALDIquant to transform raw mass spectrometry data into a matrix containing intensity measurements of potentially useful  $m/z$  values is complete.

Subsequently, the resulting feature intensity matrix can be used with any preferred univariate or multivariate analysis technique, e.g. to identify peaks that are useful for predicting a desired outcome, or simply to rank features with regard to group separation (e.g. Gibb and Strimmer, 2015).

In the following section we will describe in detail how such an analysis may be conducted. For more examples please see the MALDIquant homepage.

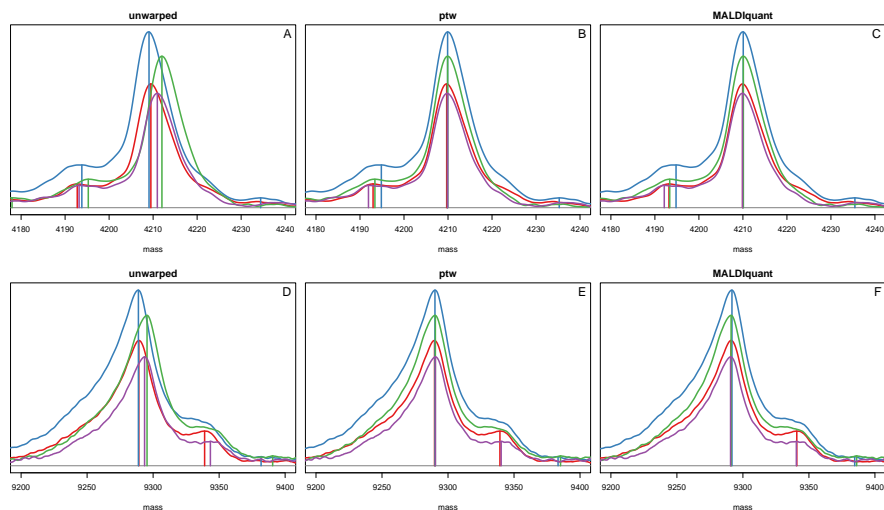


Figure 11.5: Comparison of two peaks (top row and bottom row) present in four MALDI-TOF spectra from Fiedler et al. (2009). (A, D) unaligned; (B, E) warped using the PTW algorithm; (C, F) warped using MALDIquant’s peak based PTW.

## 11.3 Case Study

### 11.3.1 Dataset

For illustration how to use MALDIquant in practical data analysis we now show in detail how to use the software by application to the mass spectrometry data published in Fiedler et al. (2009). The aim of this study was to determine proteomic biomarkers to discriminate patients with pancreas cancer from healthy persons. As part of their study the authors collected serum samples of 40 patients with diagnosed pancreas cancer as well as 40 healthy controls as training dataset. For each sample 4 technical replicates were obtained. These 320 samples were processed following a standardized protocol for serum peptidomics and subsequently analyzed in a linear MALDI-TOF mass spectrometer. For details on the experimental setup we refer to the original study.

Half of the patients and controls were recruited at the University Hospital Heidelberg and the University Hospital Leipzig. Due to the presence of strong batch effects we restrict ourselves to the samples from Heidelberg, leading to a raw data set containing 160 spectra for 40 probands, of which 20 were diagnosed with pancreatic cancer and 20 are healthy controls. Fiedler et al. (2009) found marker peaks at  $m/z$  3884 (double charged) and 7767 (single charged) and correspondingly suggested Platelet Factor 4 (PF4) as potential marker, arguing that PF4 is down-regulated in blood serum of patients with pancreatic cancer.

### 11.3.2 Preparations

Prior to preprocessing the data we first need to set up our *R* environment by install the necessary packages, namely MALDIquant (Gibb and Strimmer, 2012), MALDIquantForeign, sda and crossval, and also download the data set:

```
install.packages(c("MALDIquant", "MALDIquantForeign",  
                  "sda", "crossval"))
```

```
## load packages  
library("MALDIquant")  
  
Loading required package: methods  
This is MALDIquant version 1.13  
Quantitative Analysis of Mass Spectrometry Data  
See '?MALDIquant' for more information about this package.  
  
library("MALDIquantForeign")  
  
## download the raw spectra data (approx. 90 MB)  
githubUrl <- paste0("https://raw.githubusercontent.com/sgibb/",  
                    "MALDIquantExamples/master/inst/extdata/",  
                    "fiedler2009/")  
downloader::download(paste0(githubUrl, "spectra.tar.gz"),  
                     "fiedler2009spectra.tar.gz")  
  
## download metadata  
downloader::download(paste0(githubUrl, "spectra_info.csv"),  
                     "fiedler2009info.csv")
```

### 11.3.3 Import Raw Data and Quality Control

The first step in the analysis comprises importing the raw data into the *R* environment. As the raw data set contains both the samples from Heidelberg and Leipzig we filter out the samples from Leipzig, so that our final data set only contains the Heidelberg patients and controls:

```
## import the spectra  
spectra <- import("fiedler2009spectra.tar.gz", verbose=FALSE)  
  
## import metadata  
spectra.info <- read.csv("fiedler2009info.csv")  
  
## keep data from Heidelberg
```

```
isHeidelberg <- spectra.info$location == "heidelberg"

spectra <- spectra[isHeidelberg]
spectra.info <- spectra.info[isHeidelberg,]
```

After importing the raw data it is recommend to perform basic sanity checks for quality control. Below, we test whether all spectra contain the same number of data points, are not empty and are regular, i.e. whether the differences between subsequent  $m/z$  values are constant:

```
table(lengths(spectra))

42388
 160

any(sapply(spectra, isEmpty))

[1] FALSE

all(sapply(spectra, isRegular))

[1] TRUE
```

Next, we ensure that all spectra cover the same  $m/z$  range. The ‘trim’ function automatically determines a suitable common  $m/z$  range if it is called without any additional arguments:

```
spectra <- trim(spectra)
```

Finally, it is advised to inspect the spectra visually to discover any obviously distorted measurements. Here, for reasons of space we only plot a single spectrum:

```
plot(spectra[[47]], sub="")
```

### 11.3.4 Transformation and Smoothing

Next, we perform variance stabilization by applying the square root transformation to the raw data, and subsequently use a 41 point *Savitzky-Golay*-Filter (Savitzky and Golay, 1964) to smooth the spectra:

```
spectra <- transformIntensity(spectra, method="sqrt")

spectra <- smoothIntensity(spectra, method="SavitzkyGolay",
                           halfWindowSize=20)
```

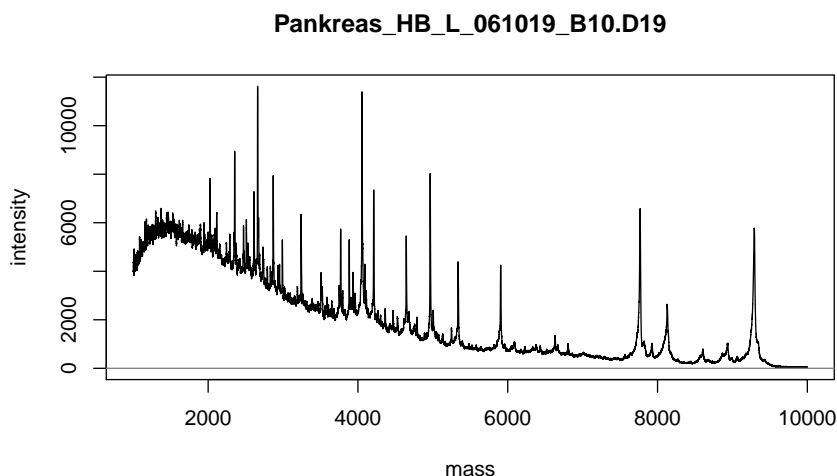


Figure 11.6: Example of a raw, uncalibrated mass spectrum.

### 11.3.5 Baseline Correction

In the next step we address the problem of matrix-effects and chemical noise that result in an elevated baseline. In our analysis we use the *SNIP* algorithm (Ryan et al., 1988) to estimate the baseline for each spectrum. Subsequently, the estimated baseline is subtracted to yield baseline-adjusted spectra:

```
baseline <- estimateBaseline(spectra[[1]], method="SNIP",  
                             iterations=150)  
plot(spectra[[1]], sub="")  
lines(baseline, col="red", lwd=2)
```

```
spectra <- removeBaseline(spectra, method="SNIP",  
                           iterations=150)  
plot(spectra[[1]], sub="")
```

### 11.3.6 Intensity Calibration and Alignment

After baseline correction we calibrate each spectrum by equalizing the TIC across spectra. After normalizing the intensities we also need to adjust the mass values. This is done by the peak based warping algorithm implemented in *MALDIquant*. In the example code the function `alignSpectra` acts as a simple wrapper around more

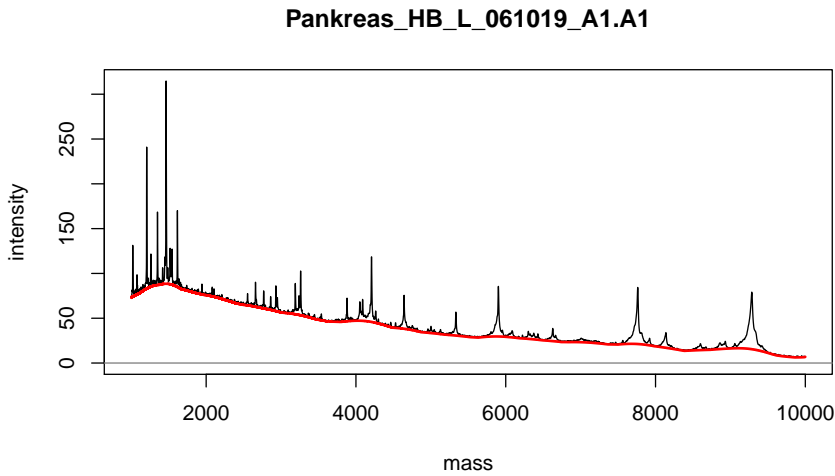


Figure 11.7: Baseline estimated using the SNIP method (red line).

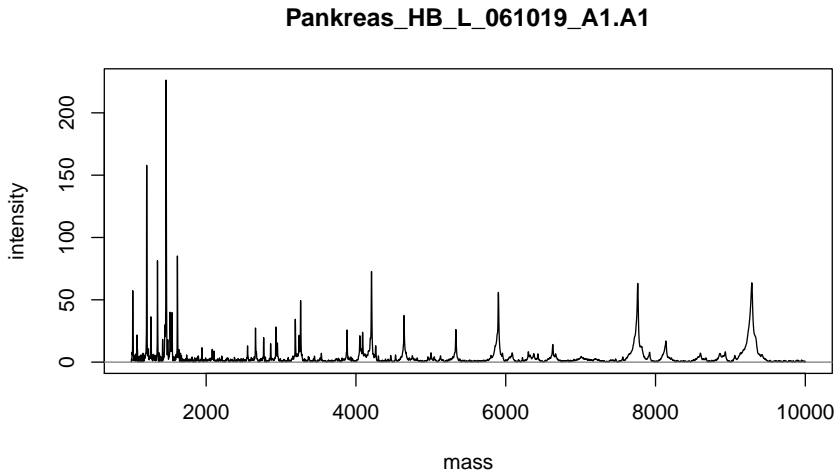


Figure 11.8: Mass spectrum after baseline correction.

complicated procedures. For a finer control of the underlying procedures the function `determineWarpingFunctions` may be used alternatively:

```
spectra <- calibrateIntensity(spectra, method="TIC")
spectra <- alignSpectra(spectra)
```

Next, we average the technical replicates before we search for peaks and update our meta information accordingly:

```
avgSpectra <-
  averageMassSpectra(spectra, labels=spectra.info$patientID)
avgSpectra.info <-
  spectra.info[!duplicated(spectra.info$patientID), ]
```

### 11.3.7 Peak Detection and Computation of Intensity Matrix

Peak detection is the crucial step to identify features and to reduce the dimensionality of the data. Before performing peak detection we first estimate the noise of selected spectra to investigate suitable settings for the *signal-to-noise ratio* (SNR):

```
noise <- estimateNoise(avgSpectra[[1]])
plot(avgSpectra[[1]], xlim=c(4000, 5000), ylim=c(0, 0.002))
lines(noise, col="red") # SNR == 1
lines(noise[, 1], 2*noise[, 2], col="blue") # SNR == 2
```

In this case we decide to set a SNR of 2 (blue line) and then run the peak detection algorithm:

```
peaks <- detectPeaks(avgSpectra, SNR=2, halfWindowSize=20)
```

```
plot(avgSpectra[[1]], xlim=c(4000, 5000), ylim=c(0, 0.002))
points(peaks[[1]], col="red", pch=4)
```

After the alignment the peak positions (mass) are very similar but not numerically identical. Consequently, binning is required to achieve identity:

```
peaks <- binPeaks(peaks)
```

In peak detection we choose a very low signal-to-noise ratio to keep as many features as possible. Using the information about class labels we can now filter out false positive peaks, by removing peaks that appear in less than 50 % of all spectra in each group:

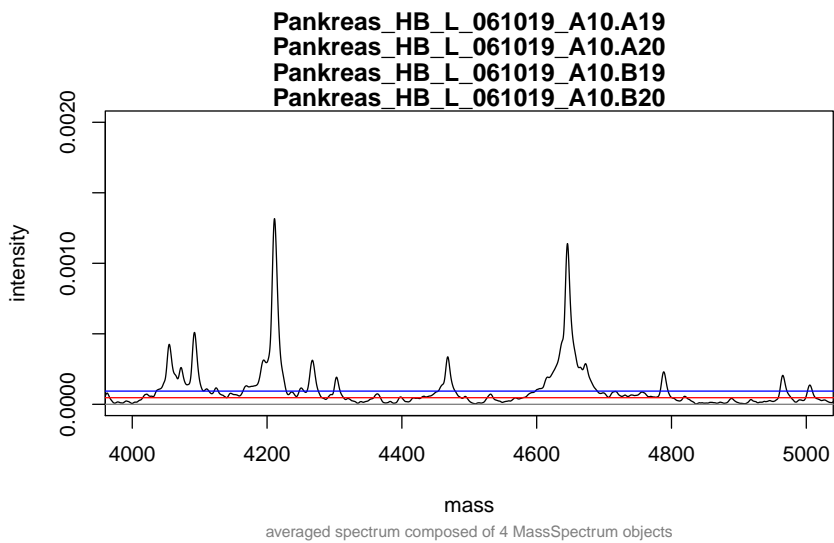


Figure 11.9: Thresholds based on signal to noise ratio (SNR): SNR=1 (red line) and SNR=2 (blue line).

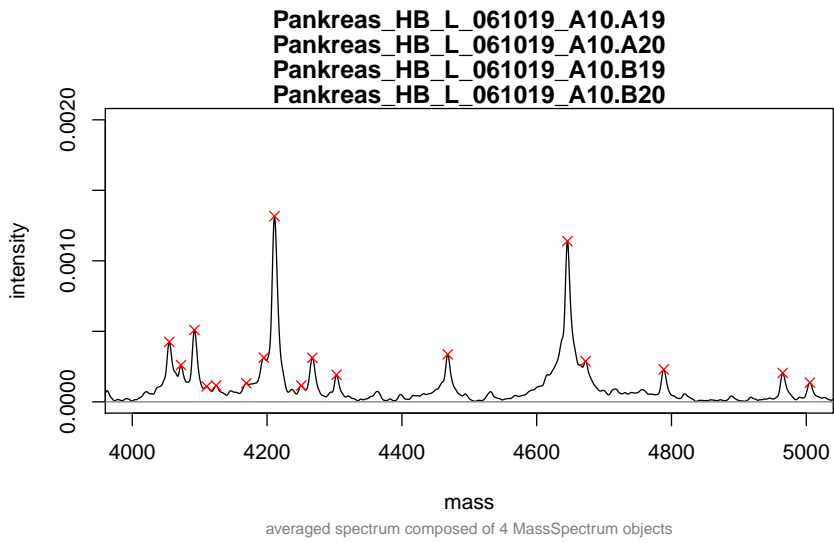


Figure 11.10: Peaks indetified above SNR=2 threshold.



```
peaks <- filterPeaks(peaks, minFrequency=c(0.5, 0.5),
                    labels=avgSpectra.info$health,
                    mergeWhitelists=TRUE)
```

As final step in MALDIquant we create the feature intensity matrix, and for convenience label the rows with the corresponding patient ID:

```
featureMatrix <- intensityMatrix(peaks, avgSpectra)
rownames(featureMatrix) <- avgSpectra.info$patientID
dim(featureMatrix)
```

```
[1] 40 166
```

This matrix is the final output of MALDIquant and contains the calibrated intensity values for identified features across all spectra. It forms the basis for higher-level statistical analysis.

### 11.3.8 Feature Ranking and Classification

The Fiedler et al. (2009) data set contains class labels for each spectrum (healthy versus cancer), hence it is natural to perform a standard classification and feature ranking analysis. A commonly used approach is Fisher's linear discriminant analysis (LDA), see Mertens et al. (2006) for details and applications to mass spectrometry analysis. Many other classification approaches may also be applied, such as based on Random Forests (Breiman, 2001) or peak discretization (Gibb and Strimmer, 2015).

Here, we use a variant of LDA implemented in the R package *sda* (Ahdesmäki and Strimmer, 2010). In particular, we use diagonal discriminant analysis (DDA), a special case of LDA with the assumption that the correlation among features (peaks) is negligible. Despite this simplification this approach to classification is very effective, especially in high dimensions (Tibshirani et al., 2003). In order to identify the most important class discriminating peaks we use standard *t*-scores, which are the natural variable importance measure in DDA.

As a first step in our analysis we therefore compute the ranking of features by *t*-scores, and list the 10 top-ranking features in Table 11.1:

```
library("sda")
colnames(featureMatrix) <-
  round(as.double(colnames(featureMatrix)), 2)
Ytrain <- avgSpectra.info$health
ddar <- sda.ranking(Xtrain=featureMatrix, L=Ytrain, fdr=FALSE,
                  diagonal=TRUE, verbose=FALSE)
```

To illustrate that feature selection based on the above feature ranking is indeed beneficial for subsequent analysis we apply hierarchical cluster analysis based on the euclidean distance first to the data set containing all features:

	idx	score	t.cancer	t.control
	8936.97	158.00	90.69	9.52
	4468.07	116.00	80.80	8.99
	8868.27	157.00	80.06	8.95
	4494.8	117.00	67.00	8.19
	8989.2	159.00	66.19	8.14
	5864.49	135.00	37.56	-6.13
	5906.17	136.00	34.43	-5.87
	2022.94	49.00	33.30	5.77
	5945.57	137.00	32.66	-5.71
	1866.17	44.00	32.12	5.67

Table 11.1: The ten top-ranking peaks as identified in the analysis.

```
distanceMatrix <- dist(featureMatrix, method="euclidean")
hClust <- hclust(distanceMatrix, method="complete")
plot(hClust, hang=-1)
```

Next, we repeat the above clustering on the data set containing only the best two top-ranking peaks:

```
top <- ddar[1:2, "idx"]
distanceMatrixTop <- dist(featureMatrix[, top],
                          method="euclidean")
hClustTop <- hclust(distanceMatrixTop, method="complete")
plot(hClustTop, hang=-1)
```

As can be seen by comparison of the two trees, as a result of the feature selection we obtain a nearly perfect split between the Heidelberg pancreas cancer samples (labeled "HP") and the Heidelberg control group (labeled "HC").

The strong predictive capabilities of the first two discovered peaks can be further quantified by conducting a cross-validation analysis to estimate the prediction error. We use the `crossval` (Strimmer, 2014) package to perform a 10-fold cross validation using the predictor containing only the two selected peaks:

```
library("crossval")
# create a prediction function for the cross validation
predfun.dda <- function(Xtrain, Ytrain, Xtest, Ytest,
```

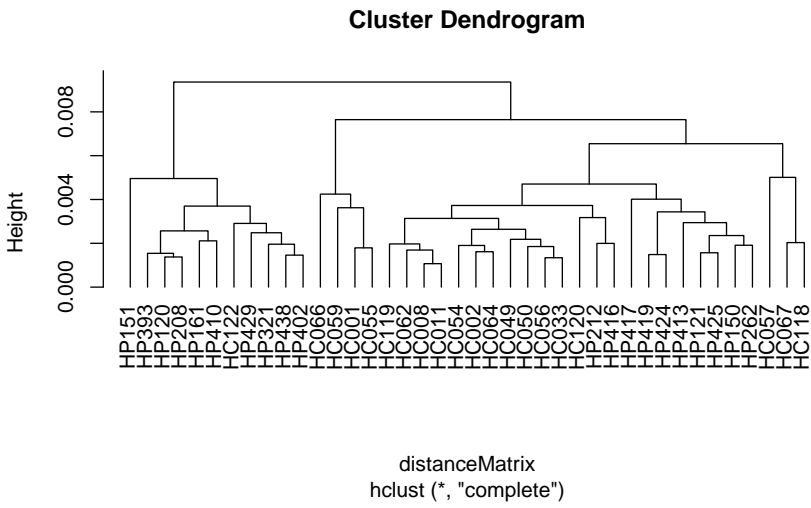


Figure 11.11: Hierarchical clustering of patient samples using all features.

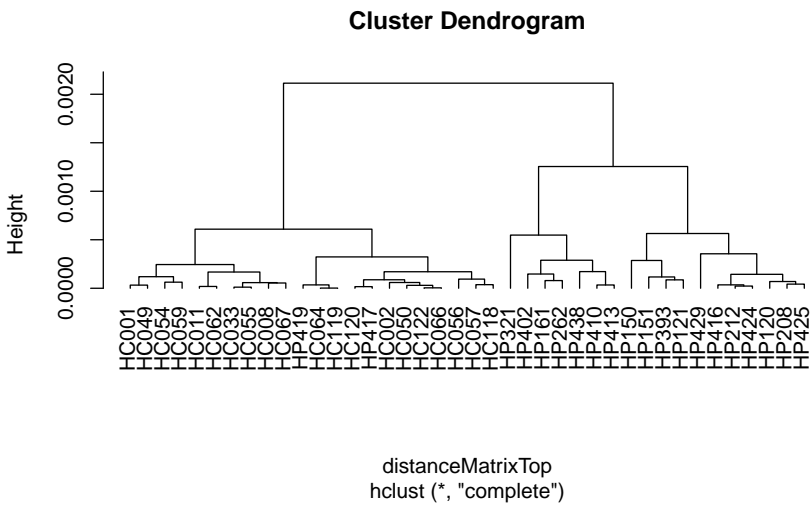


Figure 11.12: Hierarchical clustering of patient samples using only the best two top-ranking peaks.

```

                                negative) {
  dda.fit <- sda(Xtrain, Ytrain, diagonal=TRUE, verbose=FALSE)
  ynew <- predict(dda.fit, Xtest, verbose=FALSE)$class
  return(confusionMatrix(Ytest, ynew, negative=negative))
}

# set seed to get reproducible results
set.seed(1234)

cv.out <- crossval(predfun.dda,
                  X=featureMatrix[, top],
                  Y=avgSpectra.info$health,
                  K=10, B=20,
                  negative="control",
                  verbose=FALSE)
diagnosticErrors(cv.out$stat)

```

acc	sens	spec	ppv	npv	lor
0.9500000	0.9000000	1.0000000	1.0000000	0.9090909	Inf

As a result of the above analysis, we conclude that the identified peaks with mass  $m/z$  8937 and 4467 allow for the construction of a very low-dimensional predictor function that is highly effective in separating cancer and control group with both high accuracy as well as high sensitivity.

## 11.4 Conclusion

The large-scale acquisition of mass spectrometry data is becoming routine in many experimental settings. In MALDIquant we have put together a robust *R* pipeline for preprocessing these data to allow subsequent high-level statistical analysis. All methods implemented in MALDIquant have been selected both for computational efficiency and for biological validity. In this chapter we have given an overview over the most commonly used procedures of MALDIquant as well as demonstrated their application in detail.

A topic that has not been covered here is Mass Spectrometry Imaging (MSI), which combines spectral measurements with spatial information (Cornett et al., 2007). MALDIquant also enables some simple MSI analysis, for practical examples in *R* we refer to the homepage of MALDIquant at <http://strimmerlab.org/software/malDIquant/> as well as the associated web page <https://github.com/sgibb/MALDIquantExamples/>.

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