# **IPPD** package vignette

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#### Abstract

This is the vignette of the Bioconductor add-on package IPPD which implements automatic isotopic pattern extraction from a raw protein mass spectrum. Basically, the user only has to provide mass/charge channels and corresponding intensities, which are automatically decomposed into a list of monoisotopic peaks. IPPD can handle several charge states as well as overlaps of peak patterns.

### 1 Aims and scope of IPPD

A crucial challenge in the analysis of protein mass spectrometry data is to automatically process the raw spectrum to a list of peptide masses. IPPD is tailored to spectra where peptides emerge in the form of isotope patterns, i.e. one observes several peaks for each peptide mass at a given charge state due to the natural abundance of heavy isotopes. Datasets with a size of up to 100,000 mass/charge channels and the presence of isotope patterns at multiple charge states frequently exhibiting overlap make the manual annotation of a raw spectrum a tedious task. IPPD provides functionality to perform this task in a fully automatic, transparent and user-customizable way. Basically, one feeds the raw spectrum into one single function to obtain a list of monoisotopic peaks described by a mass/charge channel, a charge and an intensity. What makes our approach particularly user-friendly is its dependence on only a small set of easily interpretable parameters. We also offer a method to display the decomposition of the spectrum graphically, thereby facilitating a manual validation of the output.

### 2 Methodology

#### 2.1 Template model

In the context of this package, a protein mass spectrum is understood as a sequence of pairs  $\{x_i, y_i\}_{i=1}^n$ , where  $x_i = m_i/z_i$  is a mass  $(m_i)$  per charge  $(z_i)$  value (measured in Thomson) and  $y_i$  is the intensity, i.e. the abundance of a particular mass (modulo charge state), observed at  $x_i$ , i = 1, ..., n, which are assumed to be in an increasing order. The  $y_i$  are modeled as a linear combination of template functions representing prior knowledge about



Figure 1: Illustration of the template construction as described in the text. The left panel depicts different templates of different charge states (1 to 4). The right panel zooms at the charge two template  $\varphi_2$ .

peak shapes and the composition of isotopic patterns. If our model were exact, we could write

$$\boldsymbol{y} = \boldsymbol{\Phi}\boldsymbol{\beta}^*, \quad \boldsymbol{y} = (y_1, \dots, y_n)^\top,$$
 (1)

where  $\Phi$  is a matrix template functions and  $\beta^*$  a vector of weights for each template. Only a small fraction of all templates are needed to fit the signal, i.e.  $\beta^*$  is highly sparse. Since  $y \ge 0$ , where ' $\ge$ ' is understood componentwise, all template functions are nonnegative and accordingly  $\beta^* \ge 0$ . Model (1) can equivalently be written as

$$\boldsymbol{y} = \begin{bmatrix} \boldsymbol{\Phi}_1 & \dots & \boldsymbol{\Phi}_C \end{bmatrix} \begin{bmatrix} \boldsymbol{\beta}_1^* \\ \vdots \\ \boldsymbol{\beta}_C^* \end{bmatrix} = \sum_{c=1}^C \boldsymbol{\Phi}_c \boldsymbol{\beta}_c^*, \qquad (2)$$

where  $\Phi_c, \beta_c^*$  denote the matrix of template functions and weight vector to fit isotopic patterns of a particular charge state  $c, c = 1, \ldots, C$ . Each submatrix  $\Phi_c$  can in turn be divided into columns  $\varphi_{c,1}, \ldots, \varphi_{c,p_c}$ , where the entries of each column vector store the evaluations of a template  $\varphi_{c_j}, j = 1, \ldots, p_c$ , at the  $x_i, i = 1, \ldots, n$ . Each template  $\varphi_{c,j}$ depends on parameter  $m_{c,j}$  describing the m/z position at which  $\varphi_{c,j}$  is placed. A template  $\varphi_{c,j}$  is used to fit an isotopic pattern of peaks composed of several single peaks, which is modeled as

$$\varphi_{c,j} = \sum_{k \in Z_{c,j}} a_{c,j,k} \,\psi_{c,j,k,\boldsymbol{\theta}_{c,j}}, \quad Z_{c,j} \subset \mathbb{Z}$$
(3)

where the  $\psi_{c,j,k}$  are functions representing a peak of a single isotope within an isotopic pattern. They depend on  $m_{c,j}$  and a parameter vector  $\boldsymbol{\theta}_{c,j}$ . The nonnegative weights  $a_{c,j,k}$  reflect the relative abundance of the isotope indexed by k. The  $a_{c,j,k}$  are computed according to the averagine model (Senko et al. [1995]) and hence are fixed in advance. Each  $\psi_{c,j,k}$  is linked to a location  $m_{c,j,k}$  at which it attains its maximum. The  $m_{c,j,k}$  are calculated from  $m_{c,j}$  as  $m_{c,j,k} = m_{c,j} + \kappa \frac{k}{c}$ , where  $\kappa$  equals 1 Dalton ( $\approx 1.003$ ). The rationale behind Eq. (3) and the definitions that follow is the fact that the location of the most intense isotope is taken as characteristic location of the template, i.e. we set  $m_{c,j,0} = m_{c,j}$  so that the remaining  $m_{c,j,k}$ ,  $k \neq 0$ , are computed by shifting  $m_{c,j}$  in both directions on the m/zaxis. By 'most intense isotope', we mean that  $a_{c,j,0} = \max_k a_{c,j,k} = 1$ . The set  $Z_{c,j}$  is a subset of the integers which depends on the averagine model and a pre-specified tolerance, i.e. we truncate summation in Eq. (3) if the weights drop below that tolerance. Figure 1 illustrates the construction scheme and visualizes our notation.

#### 2.2 Peak shape

In an idealized setting, the  $\psi_{c,j,k}$  are delta functions at specific locations. In practice, however, the shape of a peak equals that of a bump which may exhibit some skewness. In the case of no to moderate skewness, we model peaks by Gaussian functions:

$$\psi_{c,j,k}(x) = \exp\left(-\frac{(x - m_{c,j,k})^2}{\sigma_{c,j}}\right).$$
(4)

The parameter to be determined is  $\theta_{c,j} = \sigma_{c,j} > 0$ . In the case of considerable skewness, peaks are modeled by exponentially modified Gaussian (EMG) functions, see for instance Grushka [1972], Marco and Bombi [2001], and Schulz-Trieglaff et al. [2007] in the context of protein mass spectrometry:

$$\psi_{c,j,k}(x) = \frac{1}{\alpha_{c,j}} \exp\left(\frac{\sigma_{c,j}^2}{2\alpha_{c,j}^2} + \frac{\mu_{c,j} - (x - m_{c,j,k})}{\alpha_{c,j}}\right) \left(1 - F\left(\frac{\sigma_{c,j}}{\alpha_{c,j}} + \frac{\mu_{c,j} - (x - m_{c,j,k})}{\sigma_{c,j}}\right)\right),$$

$$F(t) = \int_{-\infty}^t \frac{1}{\sqrt{2\pi}} \exp\left(-\frac{u^2}{2}\right) \, du.$$
(5)

The EMG function involves a vector of three parameters  $\boldsymbol{\theta}_{c,j} = (\alpha_{c,j}, \sigma_{c,j}, \mu_{c,j})^{\top} \in \mathbb{R}^+ \times \mathbb{R}^+ \times \mathbb{R}$ . The parameter  $\alpha_{c,j}$  controls the additional length of the right tail as compared to a Gaussian. For  $\alpha_{c,j} \downarrow 0$ , the EMG function becomes a Gaussian. For our fitting approach as outlined in Section 2.3, it is crucial to estimate the  $\boldsymbol{\theta}_{c,j}$ , which are usually unknown, from the data as good as possible. To this end, we model each component  $\theta_l$  of  $\boldsymbol{\theta}$  as a linear combination of known functions  $g_{l,m}$  of x = m/z and an error component  $\varepsilon_l$ , i.e.

$$\theta_l(x) = \sum_{m=1}^{M_l} \nu_{l,m} g_{l,m}(x) + \varepsilon_l(x).$$
(6)

In the case of no prior knowledge about the  $g_{l,m}$ , we model  $\theta_l$  as a constant independent of x. In most cases, it is sensible to assume a linear trend, i.e.  $\theta_l(x) = \nu_{l,1} + \nu_{l,2}x$ . In order to fit a model of the form (6), we have to collect information from the data  $\{x_i, y_i\}_{i=1}^n$ . To be precise, we proceed according to the following steps.

- 1. We apply a simple peak detection algorithm to the spectrum to identify disjoint regions  $\mathcal{R}_r \subset \{1, \ldots, n\}, r = 1, \ldots, R$ , of well-resolved peaks.
- 2. For each region r, we fit the chosen peak shape to the data  $\{x_i, y_i\}_{i \in \mathcal{R}_r}$  using nonlinear least squares:

$$\min_{\boldsymbol{\theta}} \sum_{i \in \mathcal{R}_r} (y_i - \psi_{\boldsymbol{\theta}}(x_i))^2, \tag{7}$$

yielding an estimate  $\widehat{\theta}_r(\widehat{x}_r)$ , where  $\widehat{x}_r$  denotes an estimation for the mode of the peak in region  $\mathcal{R}_r$ .

3. The sequence  $\{\widehat{x}_r, \widehat{\theta}_r\}_{r=1}^R$  is then used as input for the estimation of the parameters  $\nu_{l,m}$  in model (6).

Step 2. is easily solved by the general purpose nonnegative least squares routine nls in R:::stats for a Gaussian peak shape. For the EMG, we have to perform a grid search over all three parameters to find a suitable starting value, which is then passed to the general purpose optimization routine optim in R:::stats with the option method = "BFGS" and a specification of a closed form expression of the gradient via the argument gr. For step 3., we use least absolute deviation regression because of the presence of outliers arising from less well-resolved, wiggly or overlapping peaks. The whole procedure is performed by the function fitModelParameters as demonstrated below. After loading the package, we access the real world dataset myo500 and extract m/z channels (x) and the corresponding intensities (y). For computational convenience and since they contain very few relevant information, we discard all channels above 2500.

```
R> library(IPPD)
R> data(myo500)
R> x <- myo500[,"mz"]
R> y <- myo500[,"intensities"]
R> y <- y[x <= 2500]
R> x <- x[x <= 2500]</pre>
```

To have a look at the data, we plot the first 1000 (x,y) pairs:



In the plot, one identifies a prominent peak pattern beginning at about 804, which is zoomed at in the right panel.

We now apply fitModelParameters to fit model (6) for the width parameter  $\sigma$  of a Gaussian function (4). For simplicity, we take  $g_1(x) = 1$ ,  $g_2(x) = x$ . The model is specified by using an R formula interface.

An analogous command for the EMG (5) with the model formulae  $\alpha(x) = \nu_{1,1} + \nu_{1,2}x$ ,  $\sigma(x) = \nu_{2,1} + \nu_{2,2}x$ ,  $\mu(x) = \nu_{3,1}$  is given by

Inspecting the results, we find that R = 55 peak regions are used to fit an EMG parameter model. Moreover, it turns out that the EMG model is a more appropriate peak model for the data when visually comparing the list of mean residual sums of squares of the EMG fits and the Gauss fits extracted from slot(fitEMG, "peakfitresults") and

slot(fitGauss, "peakfitresults"), respectively. The figure shows an example where the EMG shape comes relatively close to the observed data. A long right tail indicates that a Gaussian would yield a rather poor fit here.

```
R> show(fitEMG)
```



R> visualize(fitEMG, type = "peak", cex.lab = 1.5, cex.axis = 1.25)



To assess the fit of the two linear models for the EMG parameters  $\alpha$  and  $\sigma$ , we use again the function visualize as follows:

R>



While the fit for  $\sigma$  seems to be reasonable except for some extreme outliers, the fit for  $\alpha$  is not fully convincing. Nevertheless, in the absence of further knowledge, the fit produces good results in the template matching step detailed in the next section.

#### 2.3 Template fitting

Once all necessary parameters have been determined, the positions at which the templates are placed have to be fixed. In general, one has to choose positions from the interval  $[x_1, x_n]$ . We instead restrict us to a suitable subset of the finite set  $\{x_i\}_{i=1}^n$ . The deviations from the true positions is then at least in the order of the sampling rate, but this can be improved by means of a postprocessing step described in 2.4. Using the whole set  $\{x_i\}_{i=1}^n$  may be computationally infeasible if n is large. Such an approach would be at least computationally wasteful, since 'genuine' peaks patterns occur very sparsely in the spectrum. Therefore, we apply a pre-selection step on the basis of what we term 'local noise level' (LNL). The LNL is defined as a quantile (typically the median) of the intensities  $y_i$ falling into a sliding window of fixed width around a specific position. Given the LNL, we place templates on an  $x_i$  (one for each charge state) if and only if the corresponding  $y_i$  exceeds the LNL at  $x_i$  by a factor factor.place, which typically equals three or four and has to be specified by the user. Given the positions of the templates, we compute the matrix  $\Phi$  according to Eqs. (1) and (3). It then remains to estimate the coefficient vector  $\beta^*$  on the basis of two structural assumptions, sparsity and nonnegativity of all quantities involved. Related approaches in the literature (Du and Angeletti [2006], Renard et al. [2008]) account for sparsity of  $\beta^*$  by using  $\ell_1$ -regularized regression (Tibshirani [1996]). We here argue empirically that  $\ell_1$  regularization is not the best to do, since it entails the selection of a tuning parameter which is difficult to choose in our setting, and secondly the structural constraints concerning nonnegativity turn out to be so strong that sparsity is more conveniently achieved by fitting followed by hard thresholding. We first determine

$$\widehat{\boldsymbol{\beta}} \in \underset{\boldsymbol{\beta}}{\operatorname{argmin}} \| \boldsymbol{y} - \boldsymbol{\Phi} \boldsymbol{\beta} \|_{q}^{q}, \ q = 1 \text{ or } q = 2,$$
subject to  $\boldsymbol{\beta} \ge 0.$ 
(8)

The optimization problem (8) is a quadratic (q = 2) or linear (q = 1) program and is solved using standard techniques (Boyd and Vandenberghe [2004]); we omit further details here. We remark that in the presence of high noise, it is helpful to subtract the LNL from  $\boldsymbol{y}$ . Concerning the choice of q, we point out that q = 1 can cope better with deviations from model assumptions, i.e. deviations from the averagine model or from the peak model and thus may lead to a reduction of the number of false positives.

#### 2.4 Postprocessing

Given an estimate  $\hat{\beta}$ , we define  $\mathcal{M}_c = \{m_{c,j} : \hat{\beta}_{c,j} > 0\} \subset \{x_i\}_{i=1}^n, c = 1, \ldots, C$ , as the set of all template locations where the corresponding coefficient exceeds 0, separately for each charge. Due to a limited sampling rate, different sources of noise and model misfit, the locations in the sets  $\{\mathcal{M}_c\}_{c=1}^C$  may still deviate considerably from the set of true peak pattern locations. Specifically, the sets  $\{\mathcal{M}_c\}_{c=1}^C$  tend to be too large, mainly caused by what we term 'peak splitting': for the reasons just mentioned, it frequently occurs that several templates are used to fit the same peak. This can at least partially be corrected by means of the following merging procedure.

- 1. Separately for each c, divide the sets  $\mathcal{M}_c$  into groups  $\mathcal{G}_{c,1}, \ldots, \mathcal{G}_{c,G_c}$  of 'adjacent' positions. Positions are said to be adjacent if their distance on the m/z scale is below a certain tolerance as specified via a parts per million (ppm) value.
- 2. For each c = 1, ..., C and each group  $g_c = 1, ..., G_c$ , we solve the following optimization problem.

$$(\widetilde{m}_{c,g},\widetilde{\beta}_{c,g}) = \min_{m_{c,g},\beta_{c,g}} \left\| \sum_{m_{c,j}\in\mathcal{G}_{c,g}} \widehat{\beta}_{c,j}\psi_{m_{c,j}} - \beta_{c,g}\psi_{m_{c,g}} \right\|_{L^2}^2$$
(9)

In plain words, we take the fitted function resulting from the functions  $\{\psi_{m_{c,j}}\}$  representing the most intense peak of each peak pattern in the same group and then determine a function  $\psi_{\widetilde{m}_{c,g}}$  placed at location  $\widetilde{m}_{c,g}$  and weighted by  $\widetilde{\beta}_{c,g}$  such that  $\widetilde{\beta}_{c,g}\psi_{\widetilde{m}_{c,g}}$  approximates the fit of multiple functions  $\{\psi_{m_{c,j}}\}$  best (in a least squares sense).

3. One ends up with sets  $\widetilde{\mathcal{M}}_c = \{\widetilde{m}_{c,g}\}_{g=1}^{G_c}$  and coefficients  $\{\widetilde{\beta}_{c,g}\}_{g=1}^{G_c}, c = 1, \ldots, C$ .

The additional benefit of step 2. as compared to the selection of the function with the largest coefficient as proposed in Renard et al. [2008] is that, in the optimal case, we are able to determine the peak pattern location even more accurate as predetermined by a limited sampling rate. The integral in (9) can be solved analytically for a Gaussian function, and we resort to numeric approximations for the EMG function.

The sets  $\{\mathcal{M}_c\}$  tend to be too large in the sense that they still contain noise peak patterns.

Therefore, we apply hard thresholding to the  $\{\tilde{\beta}_{c,g}\}_{g=1}^{G_c}$ ,  $c = 1, \ldots, C$ , discarding all positions where the corresponding coefficients is less than a significance level times the LNL, where the significance level has to be specified by the user.

## 3 Case study

We continue the data analysis starting in Section 2.2. The methodology of the Sections 2.3 and 2.4 is implemented in the function getPeaklist. For the computation of the template functions, we recycle the object fitEMG obtained in Section 2.2.

```
R> EMGlist <- getPeaklist(mz = x, intensities = y, model = "EMG",
model.parameters = fitEMG,
loss = "L2", trace = FALSE,
control.localnoise = list(factor.place = 2),
control.basis = list(charges = c(1, 2)),
control.postprocessing = list(ppm = 200))
R> show(EMGlist)
An object of class 'peaklist'(with postprocessing)
Loss function used: L2
Peak model used: EMG
number of peaks: 1222
charge states used: 1,2
```

R>

The argument list can be summarized as follows: we compute EMG templates for charges 1 and 2; templates are placed on all m/z-positions in the spectrum where the intensity is at least two times the LNL; the fit is least squares (loss = L2); postprocessing is performed by merging peaks within a tolerance of 200 ppm. Subsequently, only the patterns with signal-to-noise ratio bigger than three are maintained. The result is of the following form.

R> threshold(EMGlist, threshold = 3, refit = TRUE, trace = FALSE)

	loc_init	loc_most_intense	charge	quant	amplitude	localnoise	ratio
[1,]	800.4642	800.4642	1	79.44145	50.273955	9.411770	5.341605
[2,]	808.2414	808.2414	1	80.36507	50.663658	10.196100	4.968925
[3,]	829.2292	829.2292	1	93.77381	58.509383	9.411770	6.216618
[4,]	842.4935	842.4935	1	65.95672	40.883691	8.366010	4.886880
[5,]	864.4065	864.4065	1	142.82358	87.563490	13.333300	6.567278
[6,]	877.0373	877.0373	1	115.96709	70.647034	8.888890	7.947790
[7,]	908.4247	908.4247	1	77.95258	46.743175	8.104580	5.767501
[8,]	908.9339	908.9339	1	57.31024	34.357187	9.411770	3.650449
[9,]	923.4380	923.4380	1	71.87806	42.784829	7.581700	5.643171
[10,]	924.4543	924.4543	1	72.74984	43.281494	8.366010	5.173493
[11,]	927.4746	927.4746	1	109.40194	64.990843	8.888890	7.311469
[12,]	927.9689	927.9689	1	60.24656	35.781209	9.934640	3.601661
[13,]	941.4672	941.4672	1	6891.18891	4065.615602	12.549000	323.979249

Γ1/ <b>Ι</b>	0/1 6761	041 6761	1	125 00052	90 010510	12 504000	E 0007E0
[14,]	941.0701	941.0701	1	135.99053	160,219510	16 722000	5.900750
[10,]	942.4249	942.4249	1	2/1.00230	100.310307	7 200060	9.501419
[10,]	949.4364	949.4304	1	76 11004	42.198106	7.320260	5.764564
[10]	952.5318	952.5318	1	/6.11884	44.659496	7.843140	5.694084
[18,]	954.4820	954.4820	1	106.27402	62.290498	8.104580	7.685839
[19,]	963.4553	963.4553	1	148.93764	86.903255	6.535950	13.296193
[20,]	967.4826	967.4826	1	145.35796	84.644336	8.104580	10.444013
[21,]	969.4694	969.4694	1	262.52317	152.715484	10.457500	14.603441
[22,]	985.4424	985.4424	1	125.56428	72.456118	7.320260	9.898025
[23,]	991.5025	991.5025	1	143.70662	82.668836	9.411770	8.783559
[24,]	992.0244	992.0244	1	126.06443	72.501354	10.196100	7.110695
[25,]	999.4665	999.4665	1	153.63582	88.022072	7.320260	12.024446
[26,]	1023.4509	1023.4509	1	72.97884	41.200637	6.535950	6.303695
[27,]	1057.4478	1057.4478	1	61.59659	34.040137	7.320260	4.650127
[28,]	1086.5573	1086.5573	1	163.01456	88.424740	6.013070	14.705423
[29,]	1125.5187	1125.5187	1	82.19542	43.609009	6.797390	6.415552
[30,]	1151.4789	1151.4789	1	97.86679	51.210334	5.751630	8.903621
[31,]	1168.6235	1168.6235	1	88.00277	45.625087	6.274510	7.271498
[32,]	1192.7011	1192.7011	1	73.30092	37.507669	6.535950	5.738671
[33,]	1271.6609	1271.6609	1	3373.47413	1646.576999	10.457500	157.454172
[34,]	1297.6800	1297.6800	1	143.23324	68.796163	7.581700	9.073976
[35,]	1360.7611	1360.7611	1	3816.67186	1720.040463	13.071900	131.583049
[36,]	1361.7379	1361.7379	1	1393.29669	627.255229	20.653600	30.370261
[37,]	1378.8379	1378.8379	1	2998.21544	1325.426782	15.686300	84.495820
[38,]	1394.8413	1394.8413	1	133.38764	57.958356	9.150330	6.334018
[39,]	1474.6387	1474.6387	1	154.35911	63.760694	9.934640	6.418018
[40,]	1484.6606	1484.6606	1	554.63328	227.677756	11.764700	19.352619
[41,]	1500.6588	1500.6588	1	186.99548	76.008359	14.902000	5.100548
[42,]	1501.6659	1501.6659	1	237.89772	96.648852	20.130700	4.801068
[43,]	1502.6668	1502.6668	1	6512.20399	2644.083819	27.973900	94.519671
[44,]	1506.9383	1506.9383	1	1362.30919	551.937920	18.039200	30.596585
[45,]	1518.6637	1518.6637	1	3192.21932	1285.242860	14.902000	86.246333
[46,]	1519.6113	1519.6113	1	167.63225	67.460094	15.424800	4.373483
[47,]	1524.6527	1524.6527	1	142.44825	57.174412	9.411770	6.074778
[48.]	1534.6603	1534.6603	1	271.30974	108.319909	12.287600	8.815384
[49.]	1546.6550	1546.6550	1	180.73996	71.699011	10.196100	7.032003
[50.]	1588.8538	1588.8538	1	206.83078	80.197111	12.287600	6.526670
[51.]	1589.8317	1589.8317	1	205.68076	79.705151	14.379100	5.543125
[52.]	1606.8601	1606.8601	1	47656.89747	18288.544972	27.451000	666.225091
[53.]	1622.8482	1622.8482	- 1	282.47887	107.338422	8,104580	13.244168
[54.]	1628.8462	1628.8462	- 1	197,47210	74,757363	7.843140	9.531560
[55]	1632 8754	1632 8754	1	231 61098	87 460710	9 673200	9 041549
[56]	1643 8448	1643 8448	1	201.01000	110 769208	10 980400	10 087903
[57]	1650 83/8	1650 83/8	1	200.00001	81 883060	9 /11770	8 700070
[58 ]	1660 8520	1660 2520	1	346 30821	128 /03100	15 40/800	8 3300010
[50,]	1661 9502	1661 05020	1	7021 60500	120.430103	10.424000	108 751005
[60]	1675 9106	1675 0102	1	1/0 11/67	51 /00000	7 050000	7 2017715
[00,]	1693 0003	1602 0002	1	115 10006	01.492290 10 111001	6 525050	1.234140 6 112001
	1607 0674	1607.0293	1	120 47405	42.111204	7 501700	0.443024
[02,]	1710 2272	100/.00/4	1	138.4/195	50.494593	1.581/00	0.000062
Lp3,]	1/12.00/0	1/12.66/6	1	110.1/816	41.//1601	o.2/4510	0.05/349

	1717 0000	1717 0000		100 70000	00 450007	0 074540	E 201710
[64,]	1/1/.8060	1717.8060	1	100.79866	36.152097	6.274510	5.761740
[65,]	1/18.8/42	1718.8742	1	78.34258	28.083333	6.535950	4.296748
[66,]	1753.7102	1753.7102	1	89.35124	31.479318	5.490200	5.733729
[67,]	1777.8665	1777.8665	1	106.10237	36.930658	6.013070	6.141731
[68,]	1798.8819	1798.8819	1	175.36134	60.386534	7.581700	7.964775
[69,]	1815.9052	1815.9052	1	9222.12252	3149.052692	24.052300	130.925221
[70,]	1831.9004	1831.9004	1	327.93016	111.105627	5.751630	19.317242
[71,]	1837.8935	1837.8935	1	368.32466	124.419634	6.274510	19.829379
[72,]	1847.8954	1848.9034	1	373.93783	125.768310	6.274510	20.044324
[73,]	1852.9575	1853.9655	1	162.82743	54.759110	10.980400	4.986987
[74,]	1853.9657	1854.9737	1	2843.31754	956.193683	12.549000	76.196803
[75,]	1869.9595	1870.9675	1	428.17363	143.949136	6.013070	23.939375
[76,]	1885.0257	1886.0337	1	2133.58694	717.090626	9.150330	78.367734
[77,]	1885.9545	1886.9625	1	157.60503	52.969463	9.673200	5.475899
[78,]	1897.9419	1898.9499	1	73.78262	24.791973	4.44440	5.578200
[79,]	1901.0127	1902.0207	1	88.40222	29.699601	4.705880	6.311168
[80,]	1919.0078	1920.0158	1	146.81349	49.262051	6.013070	8.192496
[81,]	1937.0230	1938.0310	1	10570.05666	3542.225521	27.712400	127.820958
[82,]	1953.0164	1954.0244	1	116.41572	38.970328	3.660130	10.647252
[83,]	1958.9972	1960.0052	1	146.47000	49.010723	3.921570	12.497730
[84,]	1963.0361	1964.0441	1	219.18803	73.322477	6.013070	12.193851
[85,]	1969.9485	1970.9565	1	300.77516	100.566547	4.444440	22.627496
[86,]	1981.0615	1982.0695	1	244.81985	81.793238	13.594800	6.016509
[87,]	1982.0622	1983.0702	1	5026.45793	1679.199820	15.686300	107.048815
[88,]	1994.0448	1995.0528	1	131.82473	44.002939	3.921570	11.220746
[89,]	1995.0238	1996.0318	1	80.08518	26.730494	3.660130	7.303154
[90,]	1998.0483	1999.0563	1	86.94326	29.013517	3.398690	8.536676
[91,]	2004.0431	2005.0511	1	76.49469	25.516107	3.660130	6.971366
[92,]	2008.0804	2009.0884	1	135.36525	45.140515	3.660130	12.333036
[93,]	2039.0826	2040.0906	1	56.21758	18.706444	2.875820	6.504734
[94,]	2052.9999	2054.0079	1	68.79131	22.867988	2.352940	9.718900
[95,]	2092.1287	2093.1367	1	81.25866	26.938525	3.660130	7.359991
[96.]	2098.0541	2099.0621	1	76.68584	25.411940	3.137260	8.100043
[97.]	2105.0082	2106.0162	1	172.44634	57.105965	3.398690	16.802346
[98.]	2109.1722	2110.1802	1	245.67708	81.321211	11.764700	6.912306
[99.]	2110, 1592	2111.1672	1	5284,76546	1749,119243	14,117600	123,896359
[100]	2126 1453	2127 1533	1	79 79170	26 366710	2 352940	11 205857
[101]	2132 1398	2133 1478	1	59 99236	19 812067	2 352940	8 420133
[102]	2136 1764	2137 1844	1	73 48754	24 258854	2 614380	9 279008
[103]	2154 1284	2155 1364	1	41 38814	13 637610	2 091500	6 520493
[104]	2157 9980	2159 0060	1	41 70488	13 736534	1 830070	7 506015
[105]	2166 1065	2167 11/5	1	73 99906	24 353297	2 3529/0	10 350156
[106]	2100.1000	2107.1140	1	33 10578	10 087074	1 830070	6 003745
[100,]	2172.1009	2173.1103	1 1	26 07294	10.307274	1 622097	7 420025
[107,]	2100.0000	2109.0740	1	50.97204 69 16075	12.140300	2 001500	10 700004
[100,]	2211.1144	2212.1224	1	100.07004	22.421494	2.091500	10.720294
LIU9,j	2220.1425	2221.1505	1	102.27084	33.43U38l	1 622007	20.409386
LIIV,j	2203.0991	2234.10/1	1	33.8356/	11.000000	1 622007	0./02944
	2201.0012	2200.0102	Ţ	100 54400	23.802/04	1.03398/	14.004006
[112,]	2283.2155	2284.2235	1	182.54499	59.241203	2.091500	28.324745
L113,∫	2284.1744	2285.1824	1	24.07025	1.810563	2.091500	3.734431

[114,] 2427.2620 2428.2700 1	50.99399	16.212221	1.633987	9.921876
------------------------------	----------	-----------	----------	----------

R>

The results can be examined in detail graphically. We finally present some selected regions to demonstrate that our method performs well. The pre-defined method **visualize** can be used display the template fitting at several stages for regions within selected m/z intervals as specified by the arguments lower and upper.

```
R> visualize(EMGlist, x, y, lower= 963, upper = 973,
    fit = FALSE, fittedfunction = TRUE, fittedfunction.cut = TRUE,
    localnoise = TRUE, quantile = 0.5,
    cutoff.functions = 3)
```

R>





R>





```
R> visualize(EMGlist, x, y, lower= 1360, upper = 1364,
    fit = FALSE, fittedfunction = TRUE, fittedfunction.cut = TRUE,
    localnoise = TRUE, quantile = 0.5,
    cutoff.functions = 2)
```

R>



# 4 Extension to process LC-MS runs

In the preceding sections, it has been demonstrated how IPPD can be used to process single spectrums. For LC-MS, multiple spectra, one for a sequence of retention times  $\{t_l\}_{l=1}^L$ , have to be processed. In this context, a single spectrum is referred to as scan. The resulting data can be displayed as in Figure 2 by plotting intensities over the plane defined by retention times and m/z-values. IPPD offers basic functionality to process this kind of data. Support for mzXML format as well as an implementation of the sweep line scheme as suggested in Schulz-Trieglaff et al. [2008] is provided, which is briefly demonstrated in the sequel.

The sweep line scheme aggregates the peaklists of multiple scans by looking for blocks of consecutive retention times at which there is signal at nearby m/z-positions. The output is a quadruple consisting of a retention time interval, a m/z-position, a charge state and



Figure 2: Graphical display of the sample mzXML file used in the code.

a quantification of the intensity. The intervals are found by sequentially processing the results of getPeaklist, where the results of each peaklist will lead to extensions of existing interval of preceding lists or to the creation of new intervals; intervals are closed once they have not been extended after processing more than gap additional peaklists, where gap is a parameter to be specified by the user. For more details, we refer to Schulz-Trieglaff et al. [2008]. The function analyzeLCMS runs getPeaklist for each scan and then calls the function sweepline, which can as well be run independently from analyzeLCMS to aggregate the results. While there is a default setting, parameters can be changed by passing appropriate arguments.

The output can be displayed as follows. The retention time intervals are given by the two columns  $rt_begin$  and  $rt_end$ , the corresponding m/z-positions are given by the column loc. Quantitive information is contained in the column quant. The output is visualized by means of a contour plot, where the contour lines depict intensities over the plane defined by m/z-positions and retention times. The intervals of the output are drawn as red lines.

R> print(boxes)

	loc	charge	quant	rt_begin	$rt_end$	npeaks	gapcount
[1,]	503.3474	1	4804929	2093.26	2113.46	26	6
[2,]	505.8353	2	58628590	2067.62	2097.79	34	13
[3,]	520.3157	1	4060317	2097.79	2118.62	22	11
[4,]	521.6749	3	174362565	1929.97	1971.22	57	6

```
[5,] 534.3278
                    2 82024006 2068.95 2099.10
                                                      35
                                                                12
 [6,] 535.3383
                    1 242848614 2099.10 2150.15
                                                      70
                                                                 9
 [7,] 546.8268
                    2 43092586 2112.83 2137.57
                                                                 2
                                                      37
 [8,] 549.3673
                    1 10241899 1882.23 1925.06
                                                      56
                                                                 8
                    2 358822314 2090.02 2155.51
 [9,] 584.8515
                                                      91
                                                                10
[10,] 585.3650
                    2 17016750 2132.31 2155.51
                                                      31
                                                                 5
[11,] 585.3691
                    2 13830263 2175.63 2193.95
                                                                 3
                                                      25
[12,] 597.3784
                    2 10873799 1882.96 1906.53
                                                      35
                                                                 1
R> rtlist <- lapply(data$scan, function(x)</pre>
                  as.numeric(sub("([^0-9]*)([0-9|.]+)([^0-9]*)", "\\2", x$scanAttr)))
R> rt <- unlist(rtlist)</pre>
R> nscans <- length(rt)
R> npoints <- length(data$scan[[1]]$mass)</pre>
R> Y <- matrix(unlist(lapply(data$scan, function(x) x$peaks)),</pre>
             nrow = nscans,
             ncol = npoints,
             byrow = TRUE)
R> contour(rt, data$scan[[1]]$mass, Y, xlab = "t", ylab = "mz",
         levels = 10^{(seq(from = 5, to = 6.75, by = 0.25))},
         drawlabels = FALSE)
R> for(i in 1:nrow(boxes))
   lines(c(boxes[i,"rt_begin"], boxes[i,"rt_end"]), rep(boxes[i,"loc"], 2), col = "red")
R>
R>
```



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