Raman Spectra of Chondrocytes in Cartilage: hyperSpec's chondro data set

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1 Introduction

This vignette describes the chondro data set. It shows a complete data analysis work flow on a Raman map demonstrating frequently needed preprocessing methods

- baseline correction
- normalization
- ullet smoothing / interpolating spectra
- plotting spectra
- ullet plotting false color maps

and other basic work techniques

- cutting the spectral range,
- selecting (extracting) or deleting spectra, and
- aggregating spectra (e.g. calculating cluster mean spectra).

The chemometric methods used are

- Principal Component Analysis (PCA) and
- hierarchical cluster analysis,

showing how to use data analysis procedures provided by R and other packages.

2 The Data Set

Raman spectra of a cartilage section were measured on each point of a grid, resulting in a so-called *Raman map*. Figure 1 shows a microscope picture of the measured area and its surroundings.

The measurement parameters were:

Excitation wavelength: 633 nm
Exposure time: 10 s per spectrum

Objective: $100 \times$, NA 0.85

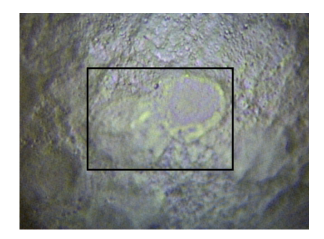


Figure 1: Microscope view of the cartilage section. The frame indicates the measurement area (35 $\times 21 \mu m$).

Measurement grid: $35 \times 21 \mu m$, $1 \mu m$ step size

Spectrometer: Renishaw InVia

All data to reproduce this Vignette is accessible at *hyperSpec*'s homepage, http://r-forge.r-project.org/projects/hyperspec/, as the original file is far too large to be included in the package.

3 Data Import

Renishaw provides a converter to export their proprietary data in a so-called long format ASCII file. Raman maps are exported having four columns, y, x, raman shift, and intensity. hyperSpec comes with a function to import such files, scan.txt.Renishaw. The function assumes a map as default, but can also handle single spectra (data = "spc"), time series (data = "ts"), and depth profiles (data = "depth"). In addition, large files may be processed in chunks. In order to speed up the reading scan.txt.Renishaw does not allow missing values, but it does work with NA.

```
> library (hyperSpec)
> filename <- "chondro.txt"

> chondro <- scan.txt.Renishaw (filename, data = "xyspc")
> chondro

hyperSpec object
   875 spectra
   3 data columns
   1272 data points / spectrum
wavelength: tilde(nu)/cm^-1 [numeric 1272] 601.622 602.664 ... 1802.15
data: (875 rows x 3 columns)
   (1) y: y/(mu * m) [numeric 875] range -4.77 -3.77 ... 19.23
   (2) x: x/(mu * m) [numeric 875] range -11.55 -10.55 ... 22.45
   (3) spc: I / a.u. [AsIs matrix 875 x 1272] range 52.2573 52.5012 ... 1884.25 + NA
```

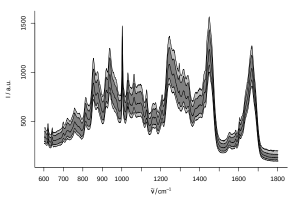
To get an overview of the spectra,:

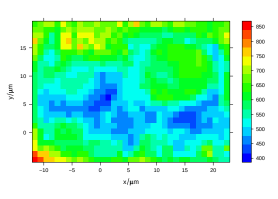
```
> plot (chondro, "spcprct15")
```

A sum intensity map is produced by:

```
> print (plotmap (chondro, na.rm = TRUE))
```

Figure 2 shows the results.





- (a) The raw spectra: median, $16^{\rm th}$ and $84^{\rm th}$, and $5^{\rm th}$ and $95^{\rm th}$ percentile spectra.
- (b) The sum intensity of the raw spectra.

Figure 2: The raw data.

4 Preprocessing

As usual in Raman spectroscopy of biological tissues, the spectra need some preprocessing.

4.1 Spectral Smoothing

As the overview print shows that the spectra contain NAs (from cosmic spike removal that was done previously), the first step is to remove these. Another issue that can be solved at the same time is that the wavelength axis is not evenly spaced (the data points are between 0.85 and 1 cm⁻¹ apart from each other). Furthermore, it would be good to trade some spectral resolution for higher signal to noise ratio. All three of these issues are tackled by interpolating and smoothing of the wavelength axis by spc.loess. The resolution is to be reduced to 8 cm⁻¹, or 4 cm⁻¹ data point spacing.

```
> chondro <- spc.loess (chondro, seq (602, 1800, 4))
> chondro

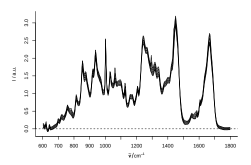
hyperSpec object
   875 spectra
   3 data columns
   300 data points / spectrum

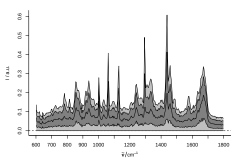
wavelength: tilde(nu)/cm^-1 [numeric 300] 602 606 ... 1798
data: (875 rows x 3 columns)
   (1) y: y/(mu * m) [numeric 875] range -4.77 -3.77 ... 19.23
   (2) x: x/(mu * m) [numeric 875] range -11.55 -10.55 ... 22.45
   (3) spc: I / a.u. [matrix 875 x 300] range 80.04420 81.75761 ... 1858.881
```

4.2 Baseline Correction

Next, we do a linear baseline correction. spc.fit.poly.below tries to automatically find appropriate support points for polynomial baselines. The default is a linear baseline, which is appropriate in our case:

```
> baselines <- spc.fit.poly.below (chondro)
Fitting with npts.min = 15
> chondro <- chondro - baselines</pre>
```





- (a) The spectra after smoothing, baseline correction, and normalization.
- (b) The spectra after subtracting the $5^{\rm th}$ percentile spectrum.

Figure 3: The preprocessed spectra.

4.3 Normalization

As the spectra are quite similar, area normalization should work well:.

```
> chondro <- sweep (chondro, 1, apply (chondro, 1, mean), "/")
> plot (chondro, "spcprct15")
```

For the results of these preprocessing steps, see figure 3a.

4.4 Subtracting the Overall Composition

The spectra are very homogeneous, but I'm interested in the differences between the different regions of the sample. Subtracting the minimum spectrum cancels out the matrix composition that is common to all spectra. But the minimum spectrum also picks up a lot of noise. So instead, the $5^{\rm th}$ percentile spectrum is subtracted:

```
> chondro <- sweep (chondro, 2, apply (chondro, 2, quantile, 0.05), "-")
> plot (chondro, "spcprctl5")
```

The resulting data set is shown in figure 3b. Some interesting differences start to show up: there are distinct lipid bands in some but not all of the spectra.

4.5 Outlier Removal by Principal Component Analysis (PCA)

PCA is a technique that decomposes the data into scores and loadings (virtual spectra). It is known to be quite sensitive to outliers. Thus, I use it for outlier detection. The resulting scores and loadings are put again into *hyperSpec* objects by decomposition:

```
> pca <- prcomp (~ spc, data = chondro$., center = TRUE)
> scores <- decomposition (chondro, pca$x, label.wavelength = "PC", label.spc = "score / a.u.")
> loadings <- decomposition (chondro, t(pca$rotation), scores = FALSE, label.spc = "loading I / a.u.")</pre>
```

Plotting the scores of each PC against all other gives a good idea where to look for outliers.

```
> pairs (scores [[,,1:20]], pch = 19, cex = 0.5)
```

> chondro <- chondro [- out]

Now the spectra can be found either by plotting two scores against each other (by plot) and identifying with identify, or they can be identified in the score map by map.identify. There is also a function to identify spectra in a spectra plot, spc.identify, but this is not helpful here.

```
> out <- map.identify (scores [,,5])</pre>
> out <- c (out, map.identify (scores [,,6]))</pre>
> out <- c (out, map.identify (scores [,,7]))</pre>
> out
[1] 105 140 216 289 75 69
> outcols <- c ("red", "blue", "#800080", "orange", "magenta", "brown")</pre>
> cols <- rep ("black", nrow(chondro))</pre>
> cols [out] <- outcols
We can check our findings by comparing the spectra to the bulk of spectra (figure):
> plot(chondro[1], plot.args = list (ylim = c (1, length (out) + .7)), lines.args = list( type = "n"))
> for (i in seq (along = out)){
     plot(chondro, "spcprct15", yoffset = i, add = TRUE, col = "gray")
     plot (chondro [out[i]], yoffset = i, col = outcols[i] , add = TRUE, lines.args = list (lwd = 2))
     text (600, i + .33, out [i]) }
and also by looking where these spectra appear in the scores pairs plot (figure):
> png ("chondro-pca-pairs2.png", width = 1000, height = 1000)
> pairs (scores [[,,1:7]], pch = 19, cex = 1, col = cols)
> dev.off ()
Finally, the outliers are removed:
```

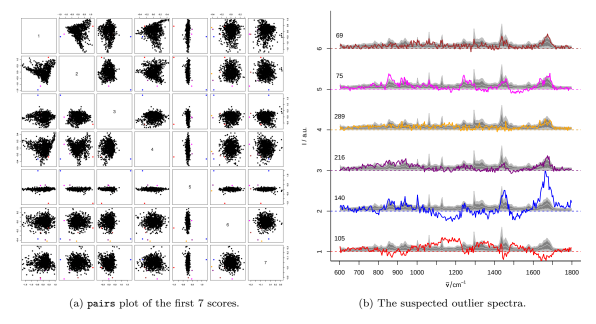


Figure 4: Outlier removal by PCA

5 Hierarchical Cluster Analysis (HCA)

HCA fuses objects according to their (dis)similarity. The result is a dendrogram, a graph stating at which level two objects are similar and thus grouped together.

The first step in HCA is the choice of the distance. The R function dist offers a variety of distance measures to be computed. The so-called Pearson distance $D_{Pearson}^2 = \frac{1 - COR(X)}{2}$ is popular in data analysis of vibrational spectra and is provided by hyperSpec.

Also for computing the dendrogram, a number of choices are available. I choose WARD's method, and, as it uses Euclidean distance for calculating the dendrogram, Euclidean distance also for the distance matrix:

```
> dist <- dist (chondro [[]])
> dendrogram <- hclust (dist, method = "ward")</pre>
```

> plot (dendrogram)

In order to get clusters, the dendrogram is cut at a level specified either by height or by the number of clusters. The result for k=3 clusters is plot as a map. If plotmap's z is a factor, the legend bar does not show intermediate colors.

```
> clusters <- cutree (dendrogram, k = 3)
> print (plotmap (chondro, z = as.factor (clusters)))
```

The cluster membership can also be marked in the dendrogram:

```
> plot (dendrogram, labels = FALSE, hang = 0)
> col.clust <- matlab.palette(3)
> points (seq_along (dendrogram$order), rep (-3, length (dendrogram$order)),
+ col = col.clust [clusters [dendrogram$order]], pch = "|")
```

Figure 5a shows the dendrogram and 5b the resulting cluster map. The three clusters correspond to the cartilage matrix, the lacuna and the cells. The left cell is destroyed and its contents are leaking into the matrix, while the right cells looks intact.

We can calculate the cluster mean spectra using aggregate. However, we can do even better and plot the cluster mean spectra \pm 1 standard deviation (see figure 6a):

```
> cluster.means <- aggregate (chondro, clusters, mean_pm_sd)
> plot(cluster.means, stacked = ".aggregate", fill = ".aggregate",
+ col = matlab.palette (3))
```

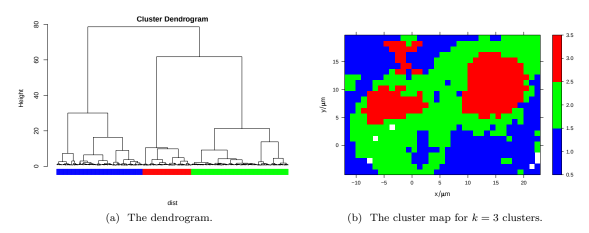


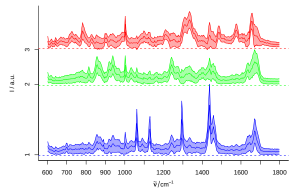
Figure 5: Hierarchical cluster analysis.

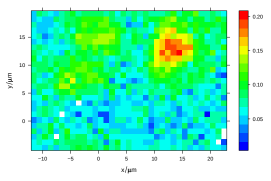
6 Plotting a False-Colour Map of Certain Spectral Regions

hyperSpec comes with a sophisticated interface for specifying spectral ranges. Expressing things like $1000 \text{ cm}^{-1} \pm 1$ data points are easily possible. Thus, we can have a fast look at the nucleic acid distribution, using the DNA bands at 728, 782, 1098, 1240, 1482, and 1577 cm⁻¹:

```
> print (plotmap (chondro[, , c( 728, 782, 1098, 1240, 1482, 1577)]))
```

The result is shown in figure 6b. While the nucleus of the right cell shows up nicely, nothing is detected in the remainders of the left cell.





- (a) The cluster mean \pm 1 standard deviation spectra. The blue cluster shows distinct lipid bands, the green cluster collagen, and the red cluster proteins and nucleic acids.
- (b) False colour map of the DNA band intensities.

Figure 6