1 Aims

So the point of this is multi-faceted:

- To provide a tutorial/introduction and access to the use of some of the crappy R code I wrote in the process of analysing peaklist MALDI imaging data during my thesis.
- To provide some example code on how (not) to use R, knitr, and LATEX, including referencing using bibtex.

2 Disclaimer

This code is a steaming pile of crap. Use it with extreme skeptisism – as you should use any software, and do any analysis, skeptisism is the lifeblood of a scientist, embrace it. When I get some free time I plan on re-writing all the plotting functions from scratch to make them nicer. Which should make this steaming pile of crap slightly more bearable.

Also, I really ought to have wrapped my code up in a package, but I am shit and haven't, so for the time being all the relevant functions can be loaded into the workspace by calling

```
source('localFunctions.R')
```

the fact that this is here may prompt me to getting around to cleaning it up and wrapping it up in package form at some point in the future, but probably not in the immediate future.

3 Setup and Reading Peaklists

I set the current dataset_name to a variable,

```
and then call

pl_all <- readPeaklists(dataset_name)

This reads in peaklist files from</pre>
```

<parent_folder_name>/<dataset_name>/<peaklist_folder_name>

and returns the number of empty spectra after writing the three files:

```
<dataset_name>_comprehensive_peaklist.txt
<dataset_name>_fExists.txt
<dataset_name>_LXY.txt
```

```
./<data_folder>
```

dataset_name is required, but the other arguments are optional and default to:

```
• peaklist_folder_name = "peaklists"
```

- parent_folder_name = "."
- data_folder = "./data"

Once created the files written to data_folder can be read in easily by calling

```
pl_all <- load_peaklist(dataset_name)

LXY <- load_LXY(dataset_name)

fExists <- load_fExists(dataset_name)</pre>
```

respectively. These load_* functions also accept an optional data_folder argument if an alternative location is used to store these files.

4 Peak Grouping and Peaklist Subsets

There are three functions included for assigning 'peakgroup' labels to peaks:

4.1 Mass Matching

Peaks can be matched to known masses by mass-error. In these data for example, there are some internal calibrants of known mass, as described by [2].

The function mzMatch extracts peaks from the first argument about the m/z values in the second argument.

```
pl_cal <- mzMatch(pl_all,cal_df$m.z)</pre>
```

Optional arguments binMargin (mass error to be allowed) and use_ppm (mass error measured in ppm or Da) can also be specified, but otherwise default to:

- binMargin = 0.3, and
- use_ppm = FALSE.

The function mzMatch returns the subset of the input peaklist with an added column, PeakGroup, specifying the theoretical mass that peak is matched too. Not sure how this would react to overlapping mass windows but it was not intended for that.

4.2 Tolerance Clustering

When the masses of interest are not known, peakgroups can be formed via a onedimensional clustering of the m/z values of the peaks. Tolerance Clustering is one of the simplest ways of doing this, and boils down to finding the equivalence classes of the relation defined on two peaks as 'being within some tolerance tol of each other'.

```
pl_tol <- groupPeaks(pl_all)</pre>
```

The optional tolerance argument tol defaults to a value of 0.1Da, and an additional optional argument minGroupSize can be specified to label any equivalence classes with less than that many peaks in them zero. By default all peaks will be labeled. The function groupPeaks returns the the input peaklist with an added column, PeakGroup, specifying a peakgroup label.

4.3 DBSCAN

A more sophisticated clustering approach is to use a density based clustering such as DBSCAN, or more precisely its deterministic version DBSCAN*, as described in [1].

```
pl_dbs <- dbscan_lw(pl_all,pp=FALSE)</pre>
```

The function dbscan_lw works similarly to the function groupPeaks, in that it takes a peaklist and returns the same peaklist with an added column, PeakGroup, containing a peakgroup label. The function dbscan_lw also takes optional arguments eps (similar to the tol of the tolerance clustering, specifying the width of the rectangular kernal used), mnpts (the minimum number of points within a eps-neighbourhood considered significant – adjusting mnpts can fix the problem in large datasets of different masses being combined), cvar (specifying the variable in the input peaklist to be clustered) and pp (print progress to console logical). These optional arguments default to:

```
• eps = 0.05,
```

- mnpts = 100,
- cvar ="m.z", and
- pp =TRUE

4.4 Evaluating Parameter Choices for Peak Grouping

Both the tolerance clustering of Section 4.2 and the DBSCAN approach of Section 4.3 involve the choice of a number of tuning parameters, in the former only in grouping tolerance tol and in the latter both a grouping tolerance eps and a minimum density mnpts. In this section I'll demonstrate how to make a heuristic diagnostic plot I use as a sanity check that my parameter selection has been somewhat reasonable. First I summarise the peak grouping by calculating a number of statistics for each peak group:

```
pgs_tol <- ddply(pl_tol,</pre>
                 "PeakGroup",
                 summarise.
                 AWM = weighted.mean(m.z,log1p(intensity)),
                 Range = max(m.z) - min(m.z),
                 nPeaks = length(m.z),
                 nDupPeaks = (length(m.z)
                 - length(unique(Acquisition))))
pgs_dbs <- ddply(subset(pl_dbs, PeakGroup!=0),</pre>
                 "PeakGroup",
                 summarise,
                 AWM = weighted.mean(m.z,log1p(intensity)),
                 Range = \max(m.z) - \min(m.z),
                 nPeaks = length(m.z),
                 nDupPeaks = (length(m.z)
                 - length(unique(Acquisition))))
```

In particular:

- The AWM: the abundance weighted mean m/z of peaks in the peakgroup this is mostly useful for reference when matching particular peakgroups of interest to LC-MS/MS identifications, or some such, for example.
- The Range: the difference between the minimum and maximum m/z in the peak group.
- The nPeaks: the number of peaks in the peak group.
- The nDupPeaks: the number of duplicate peaks, or the number of occurrances of more than one peak from the same spectrum being included in the same peakgroup

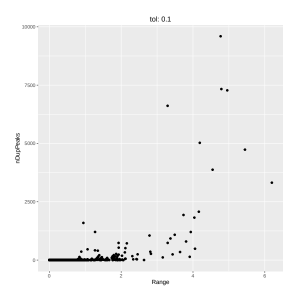


Figure 1: Plot for heuristic evaluation of peak grouping efficacy: tolerance clustering.

Ideally, the Range should be small, and the nDupPeaks should be zero. In reality, compromises must be made. When using the simplistic tolerance clustering approach, optimising parameter choice is correspondingly simple. You could for example consider a plot such as Figure 1:

```
p = (ggplot(pgs_tol,aes(x=Range,y=nDupPeaks))
    + geom_point()
    + ggtitle(paste('tol:',toString(0.1))))
print(p)
```

The nPeaks is also useful as peakgroups with nPeaks too small will typically be uninteresting, and so you could consider the number of peak groups with at least some threshold value of nPeaks, you could apply a similar filter when making a plot such as Figure 1 in order to get rid of some of the junk lying around the origin. For peptide data, typically you shouldnt really have any peakgroups with a Range above 1 Da, and any half-decent quality peakpicking should allow for you do do peak peaking that keeps the nDupPeaks in the single digits, generally speaking. So Figure 1 shows that this is really quite terrible, and we should try a smaller value of tol. Hopefully, reproducing this analysis with a smaller tol will improve these things. Generally, you want to choose a tol as large as possible so that the Range and nDupPeaks are somewhat reasonable. Sometimes this is not possible – there simply is no happy medium, no value of tol for which simple tolerance clustering produces nice peak groups. This is usually when using a more sophisticated approach, such as DBSCAN, is a

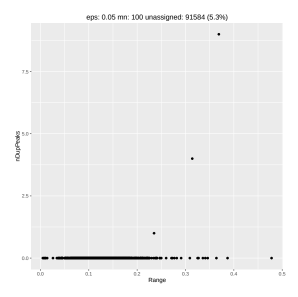


Figure 2: Plot for heuristic evaluation of peak grouping efficacy: DBSCAN clustering.

good idea. Being a more sophisticated approach, it also involves more tuning paramters, more factors to consider. We can have a look at Figure 2, analogous to Figure 1:

```
p = (ggplot(pgs_dbs,aes(x=Range,y=nDupPeaks))
+ geom_point()
+ ggtitle(paste(
   'eps: ',toString(0.05),
   'mn: ',toString(100),
   'unassigned: ',
   toString(sum(pl_dbs$PeakGroup==0)),' (',
   toString(round(100*sum(pl_dbs$PeakGroup==0)/nrow(pl_dbs),1)),
   '%)',sep="")))
print(p)
```

The interpretation of Figure 2 itself is similar to that of Figure 1, the main differences are two-fold:

- There is an extra tuning parameter (density, mnpts) which can be thought
 of as the extra degree of freedom allowing us to find nice solutions which
 cannot be acheived with tolerance clustering alone.
- DBSCAN will not allocate some number of peaks to any peakgroup dbscan_lw() will assign these peaks a PeakGroup of 0. As this is essentially throwing away the information in the peaks, you kinda want to minimise how much

data you throw away in order to acheive a nice peak grouping, so this adds an additional factor into the equation when trying to tune your tuning parameters.

When trying to pick some tuning parameters, I will typically produce a bunch of plots like Figure 2 with various (reasonable) choices for tuning parameters and then try to choose a healthy medium between all the different factors I'm trying to optimise.

5 Simple Plots

I also have some functions that make plots specially for peaklist data. This part is the most hacked up, as I have repeatedly modified these functions to perform various different tasks, while trying to maintain backwards compatability and its all ended in a giant mess. These are such a mess I am not even going to bother trying to explain them, and instead just recommend you write your own plotting functions, as then you can be sure your plotting the thing you want to plot. Your also welcome to look at the code under spatialPlot and acquisitionPlot and canabilise code to your hearts content. I provide an example of one use of the function spatialPlot below when I produce a DIPPS map using it, although it can do alot more than this – I plan on coming back and re-writing the plotting functions at some point so they make more sense. But good luck on that ever happening, hahaha.

6 DIPPS

Now say you have produced some peakgroups one way or another, and now you have two regions you want to compare using DIPPS. For example, here I have annotation of the center of cancer tumours stored in an xml 'ROI' file. So I'll read the annotations into R using the XML package and merge them onto the LXY variable as a 'ROI' column with value 'None' for spectra not in any annotated region.

We can take a quick look at these annotations by plotting them. Figure 3 demonstrates this, as well as providing a simple (less confusing?) example of a straightforward way to make spatial plots without using my gargantuan spatialPlot function (although you could equally make this plot using spatialPlot if you really wanted to.

Now I create a simplified peaklist variable pl_uni which intially has only two variables, Acquisition (indexing the originating spectrum) and PeakGroup (indexing the peakgroup). I also make sure the rows of pl_uni are unique – this is important, as having multiple peaks from the same peakgroup in the same spectrum will otherwise affect your results, although in the senario that this occurs more than a couple of times I would suggest perhaps revisiting whatever decisions you made at your peakgrouping step. I add a third variable, Group to the peaklist pl_uni, identifying each peak as originating from either an annotated region (2), or not (1). Now that we have cleaned up pl_uni and ensured it has the three neccessary columns, and that they are correctly formatted we can plug this right into the DIPPS function.

```
pl_uni = unique(pl_tol[,c("Acquisition","PeakGroup")])
pl_uni = subset(pl_uni,PeakGroup!=0)
temp = match(pl_uni$Acquisition,LXY$Acquisition)
pl_uni$Group = 1+(LXY[temp,]$ROI != "None")
dipsum = DIPPS(pl_uni)
nStar = dippsHeur(pl_uni,dipsum)
```

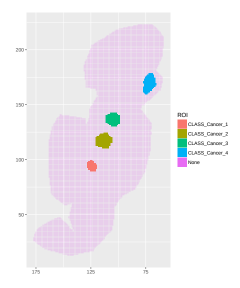


Figure 3: Annotation Regions

Note that in this case I used pl_tol to make pl_uni which I used in the DIPPS analyses – this was the peaklist with the PeakGroup column created by tolerance clustering. I could easily have used pl_dbs or even p_cal (if I was only interested in the calibrants) instead of pl_tol. Another option would be to bin the peaks in a data-independent manner using the R base function cut – a potentially useful option when interested in prediction, because of its independence on the data.

Also note how I generate nStar, which is the number of variables suggested to be optimal by the heuristic. One way to visualise these top nStar 'DIPPS features' is in a 'DIPPS map', so I might as well demonstrate how to do that using the hugely dodgey spatialPlot function:

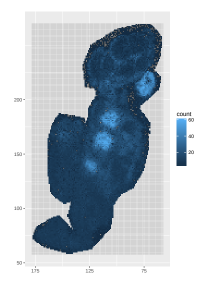


Figure 4: DIPPS Map

```
+ xlab("")
)
# DIPPS Map
print(p)
```

One could also produce individual intensity plots of particular peakgroups, for example for the peakgroup with highest DIPPS statistic value:

and you could for example have a look at the AWM or various other statistics for these peakgroups if you wanted. For example:

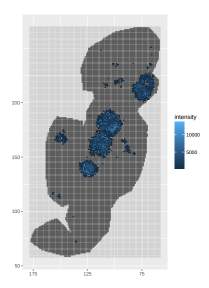


Figure 5: Intensity Map for peakgroup most highly ranked by DIPPS

```
pgs_tol = merge(pgs_tol,dipsum)
pgs_tol = pgs_tol[rev(order(pgs_tol$d)),]
print(head(pgs_tol), digits=3)
##
        PeakGroup AWM Range nPeaks nDupPeaks
                                                  p.d
## 818
              818 1407 0.430
                              1523
                                            0 0.0767 0.963 0.886
##
  2948
             2948 2854 0.203
                               1304
                                            0 0.0611 0.944 0.883
## 2504
             2504 2484 0.150
                               709
                                            0 0.0190 0.882 0.863
## 782
              782 1391 0.413
                               2363
                                            0 0.1382 0.994 0.856
## 767
              767 1384 0.242
                               1090
                                            0 0.0475 0.880 0.832
## 1743
             1743 1998 0.256
                                            0 0.0141 0.821 0.807
                                612
print(pgs_tol[57:61,], digits=3)
##
        PeakGroup AWM Range nPeaks nDupPeaks
                                                   p.d
                                                         p.u
## 692
              692 1347 0.221
                                221
                                            0 0.00806 0.219 0.211
## 293
              293 1155 4.163
                              12379
                                         2074 0.73300 0.942 0.209
## 1817
             1817 2054 0.265
                                741
                                            0 0.04561 0.252 0.207
## 1416
             1416 1776 2.789
                              12067
                                         1053 0.78525 0.959 0.174
## 727
              727 1362 0.185
                                212
                                             0 0.00888 0.181 0.172
```

In addition to calculating the AWM here I have also calculated the m/z range over which each peakgroup spans, and the number of duplicated peaks (a number above zero indicated there are individual spectra with more than one peak in the indicated peakgroup). Notice that the most highly ranked variables by DIPPS

seem fine, but that there are some (AWM = 1155 and AWM = 1776) for which it seems the peakgrouping has failed pretty badly, producing peakgroups that span 4.16Da and 2.789Da respectively. Note that you can look at such things without having done the DIPPS step, and it is worth doing so as a quality-control/sanity-check step. You could for example try using the dbscan peakgroups instead – I'll leave that as an exercise.

References

- [1] Ricardo JGB Campello, Davoud Moulavi, and Joerg Sander. Density-based clustering based on hierarchical density estimates. In *Advances in Knowledge Discovery and Data Mining*, pages 160–172. Springer, 2013.
- [2] Johan OR Gustafsson, James S Eddes, Stephan Meding, Tomas Koudelka, Martin K Oehler, Shaun R McColl, and Peter Hoffmann. Internal calibrants allow high accuracy peptide matching between MALDI imaging MS and LC-MS/MS. *Journal of Proteomics*, 75(16):5093–5105, 2012.

Session Info and pdflatex Version

```
sessionInfo()
## R version 3.1.0 (2014-04-10)
## Platform: x86_64-w64-mingw32/x64 (64-bit)
##
## locale:
## [1] LC_COLLATE=English_United States.1252
## [2] LC_CTYPE=English_United States.1252
## [3] LC_MONETARY=English_United States.1252
## [4] LC_NUMERIC=C
## [5] LC_TIME=English_United States.1252
##
## attached base packages:
## [1] grid
                stats graphics grDevices utils
                                                              datasets methods
## [8] base
##
## other attached packages:
## [1] plyr_1.8.3 ggplot2_2.1.0
## [5] stringr_1.0.0 knitr_1.12.3
                                            reshape2_1.4.1 data.table_1.9.6
## loaded via a namespace (and not attached):
## [1] chron_2.3-47 colorspace_1.2-6 digest_0.6.9
## [5] formatR_1.3 gtable_0.2.0 highr_0.5.1
                                                              magrittr_1.5
                           gtable_0.2.0 highr_0.5.1 Rcpp_0.12.4 scales_0.4.0
## [9] munsell_0.4.3 Rcpp_0.12.4
                                                                stringi_1.0-1
## [13] tools_3.1.0
```

```
system('pdflatex --version',intern=TRUE)

## [1] "MikTeX-pdfTeX 2.9.5840 (1.40.16) (MikTeX 2.9 64-bit)"

## [2] "Copyright (C) 1982 D. E. Knuth, (C) 1996-2014 Han The Thanh"

## [3] "TeX is a trademark of the American Mathematical Society."

## [4] "compiled with zlib version 1.2.8; using 1.2.8"

## [5] "compiled with libpng version 1.6.19; using 1.6.19"

## [6] "compiled with poppler version 0.32.0"

## [7] "compiled with jpeg version 8.4"
```