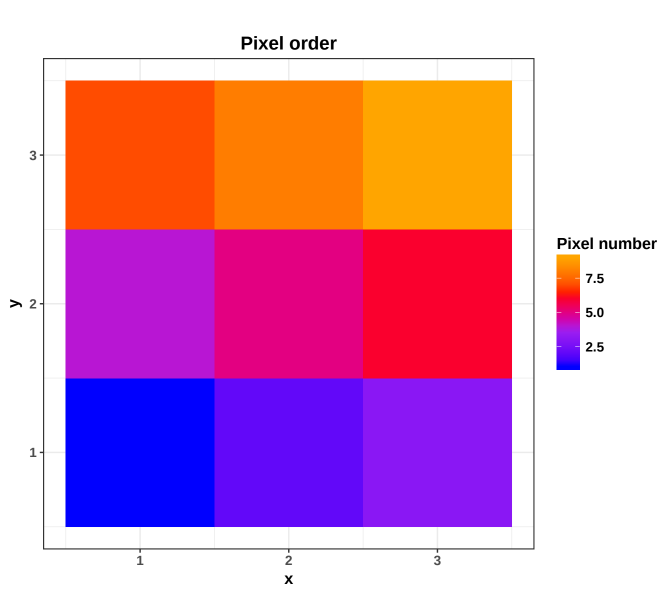
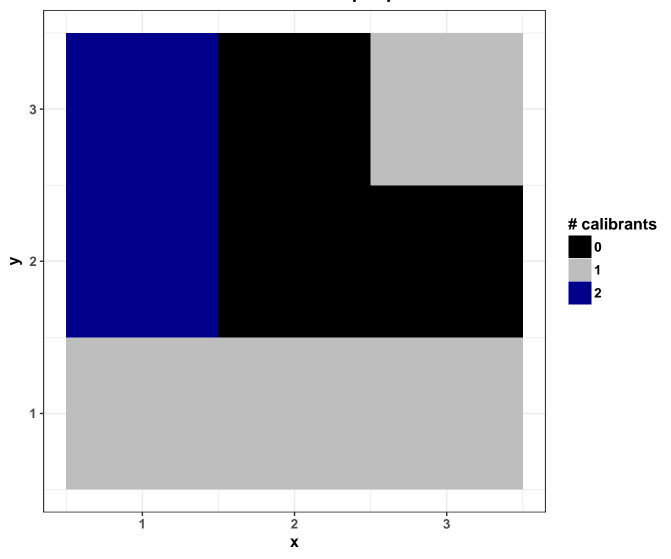
Testfile_imzml

properties	values
Number of m/z features	5199
Range of m/z values	100 – 799.81
Number of pixels	9
Range of x coordinates	1 – 3
Range of y coordinates	1 – 3
Range of intensities	0 – 9.24
Median of intensities	0
Intensities > 0	35.16 %
Number of zero TICs	0
Median TIC	161.81
Median # peaks per spectrum	1961
Normalization	FALSE
Smoothing	FALSE
Baseline reduction	FALSE
Peak picking	FALSE
Centroided	FALSE
# peptides in inputpeptides.txt	3/6
# calibrants in inputcalibrantfile1.txt	3/3

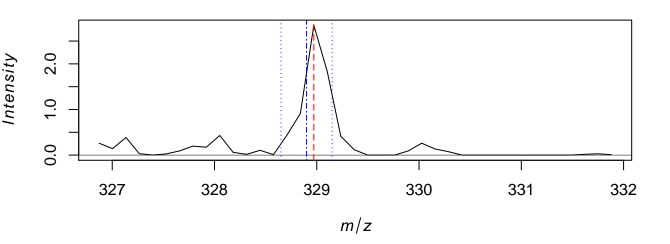


Number of calibrants per pixel

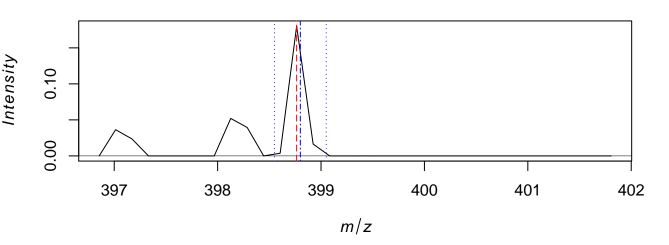


Control of fold change plot

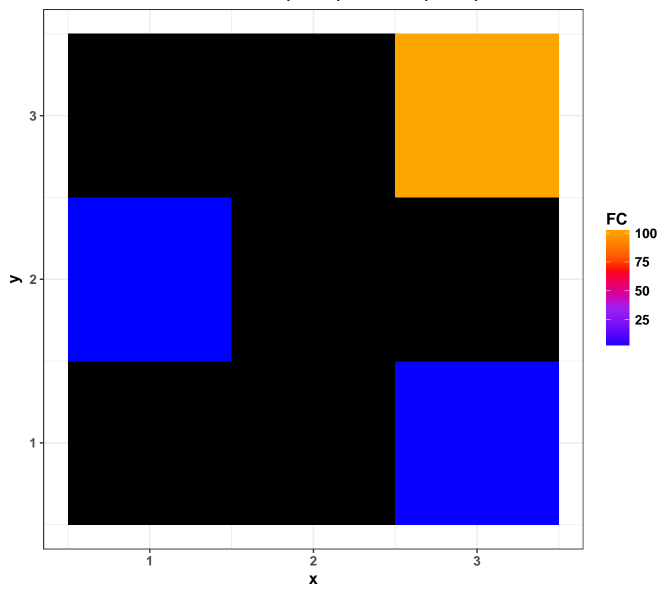
average spectrum 328.9 Da



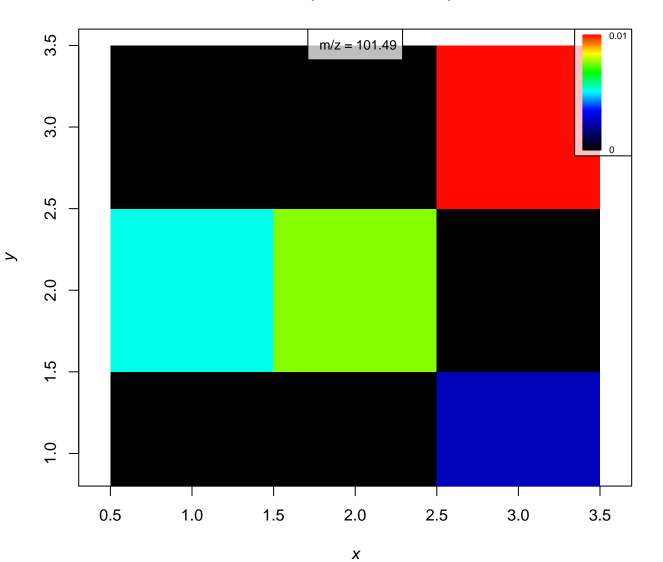
average spectrum 398.8 Da



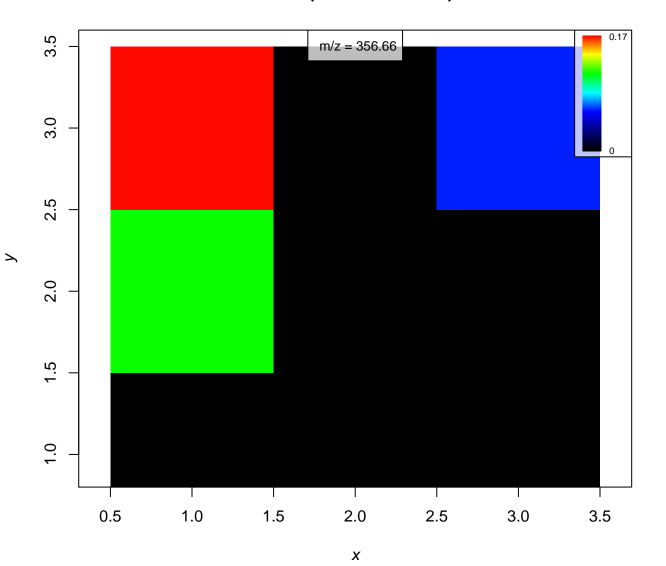
Ratio of mass1 (328.9) / mass2 (398.8)



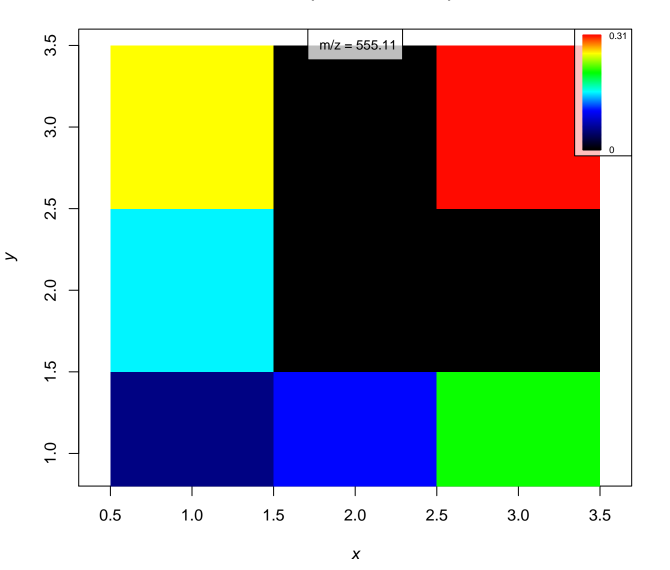
101.5 (101.5 ± 0.25 Da)



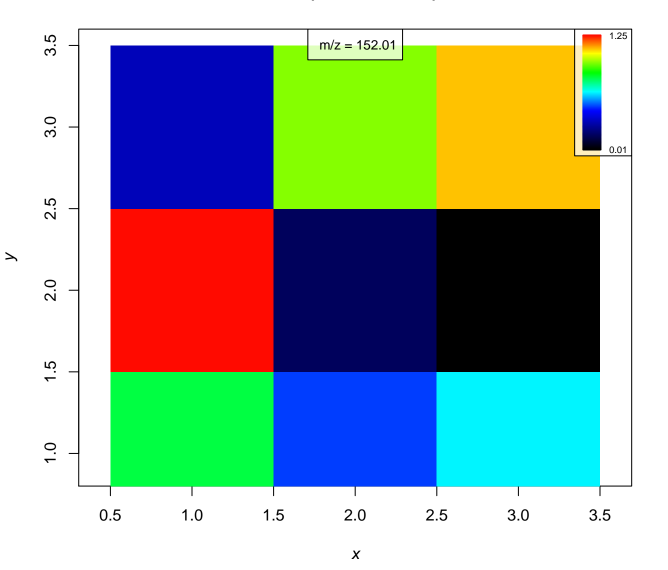
356.7 (356.7 ± 0.25 Da)



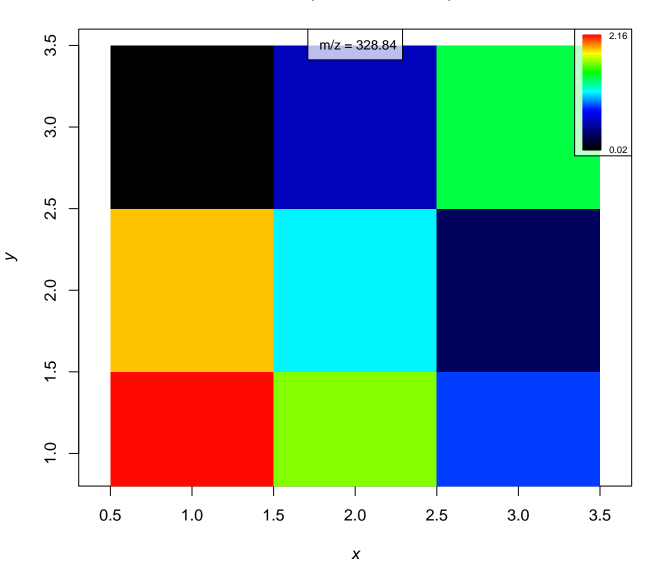
555.1 (555.1 ± 0.25 Da)



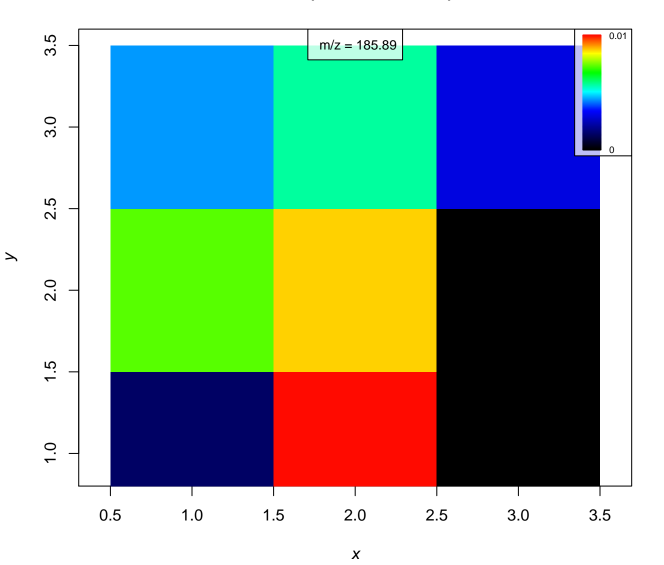
152 (152 ± 0.25 Da)



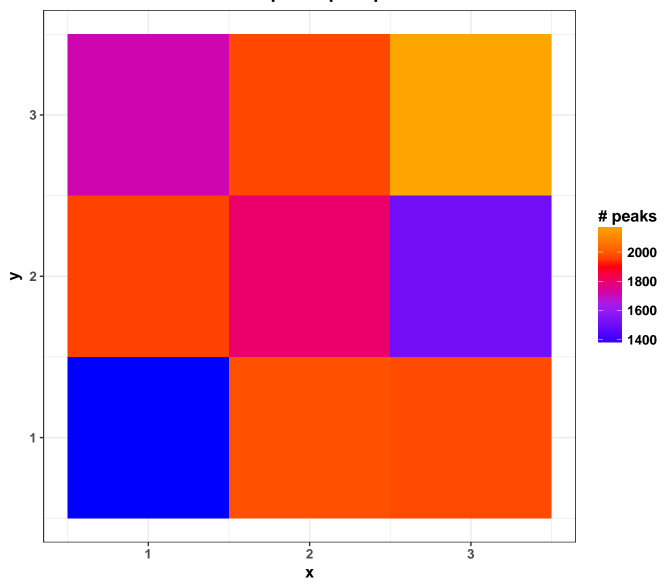
328.9 (328.9 ± 0.25 Da)



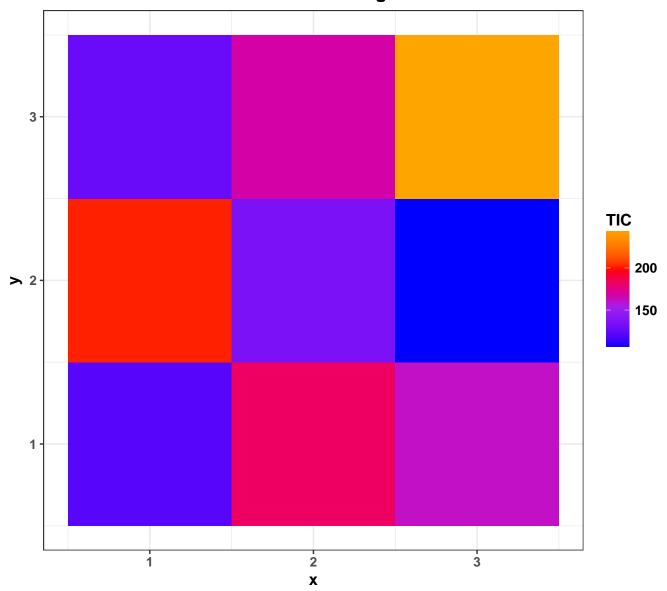
185.9 (185.9 ± 0.25 Da)



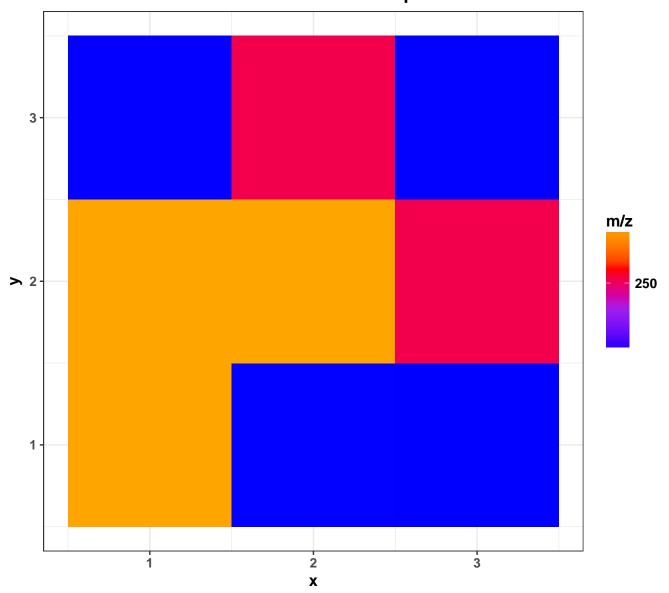
Number of peaks per spectrum



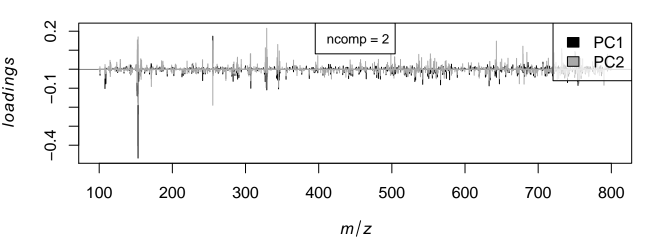
Total Ion Chromatogram

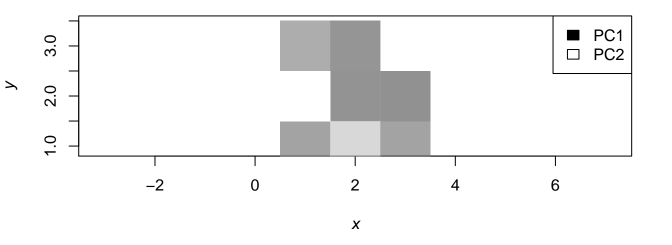


Most abundant m/z in each spectrum

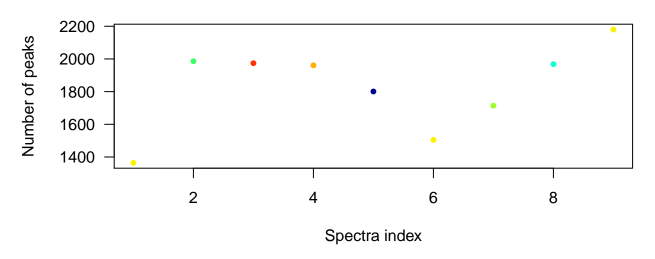


PCA for two components

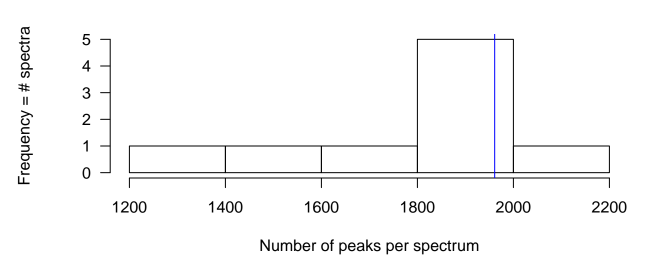


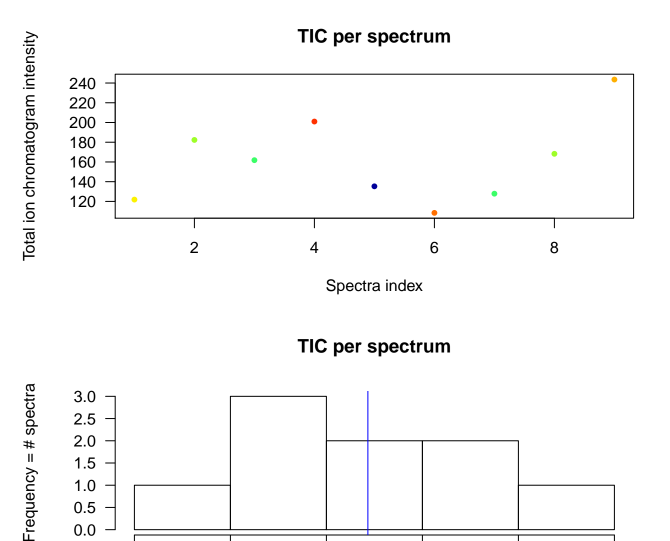


Number of peaks per spectrum



Number of peaks per spectrum





0.5 0.0

4.6

4.8

log(TIC per spectrum)

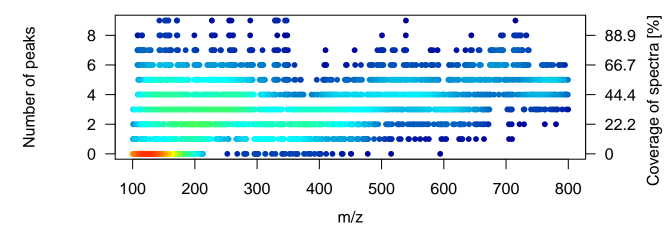
5.2

5.4

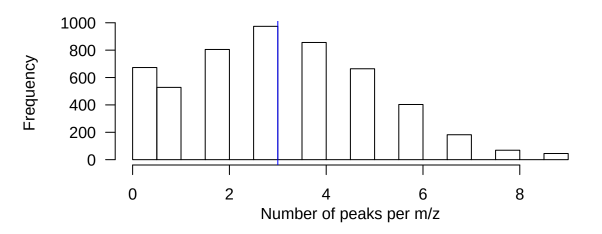
5.6

5.0

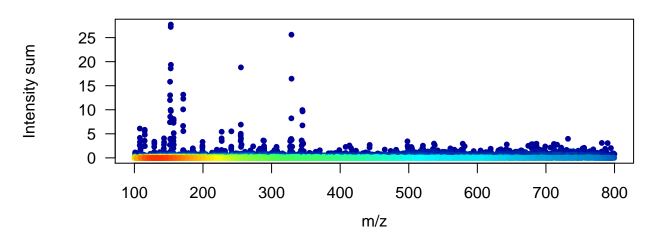
Number of peaks per m/z



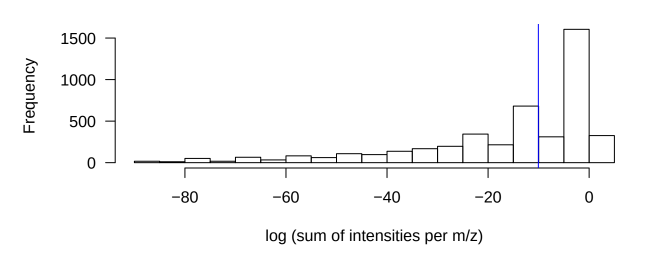
Number of peaks per m/z



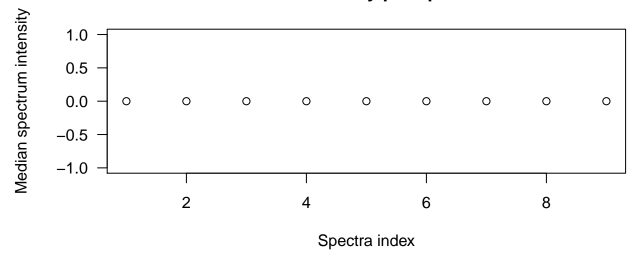
Sum of intensities per m/z



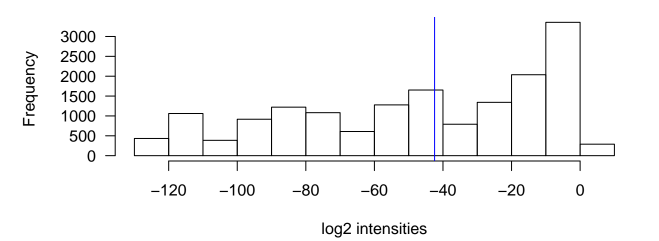
Sum of intensities per m/z



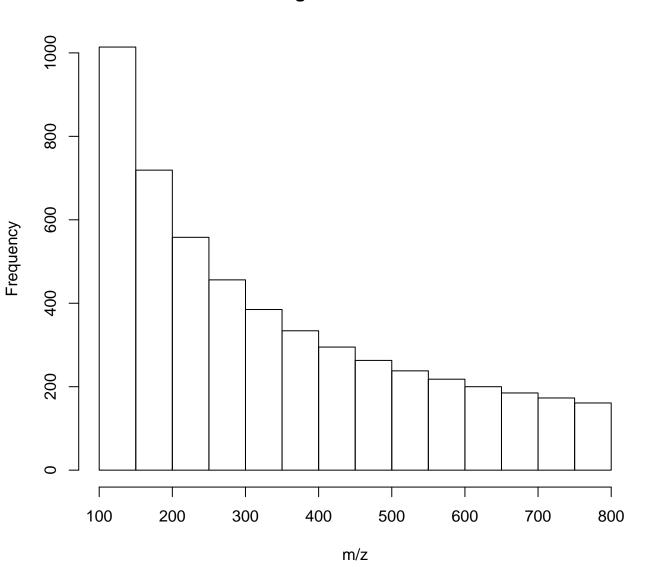
Median intensity per spectrum

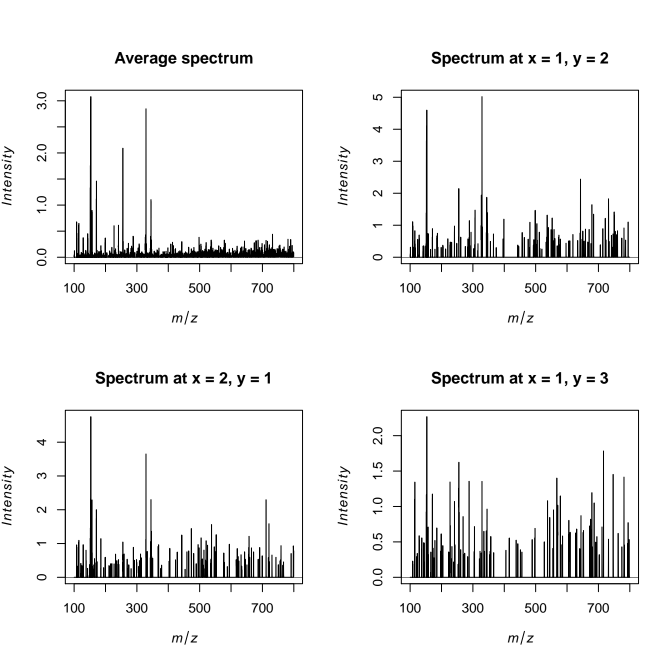


Log2-transformed intensities

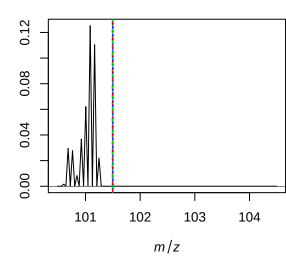


Histogram of m/z values



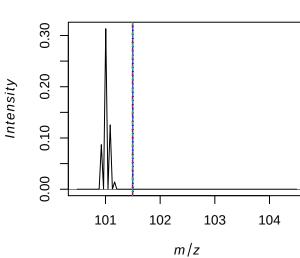




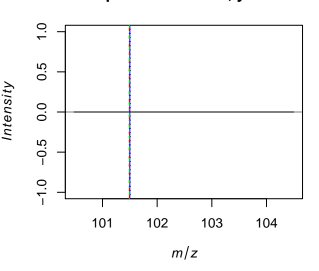


Intensity

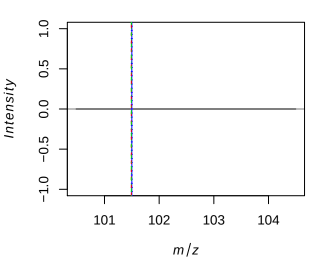
Spectrum at x = 1, y = 2



Spectrum at x = 2, y = 1

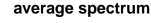


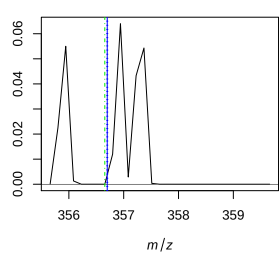
Spectrum at x = 1, y = 3



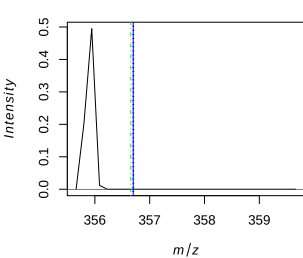
Intensity

Intensity

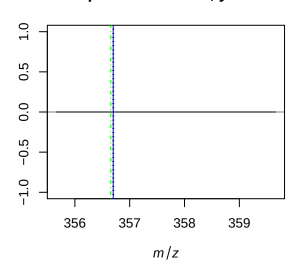




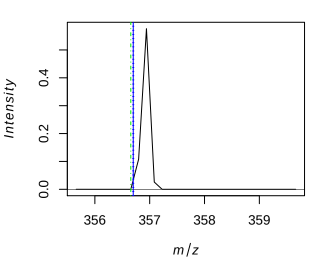
Spectrum at x = 1, y = 2



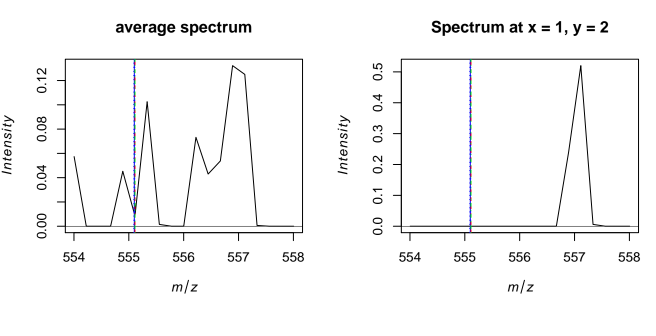
Spectrum at x = 2, y = 1

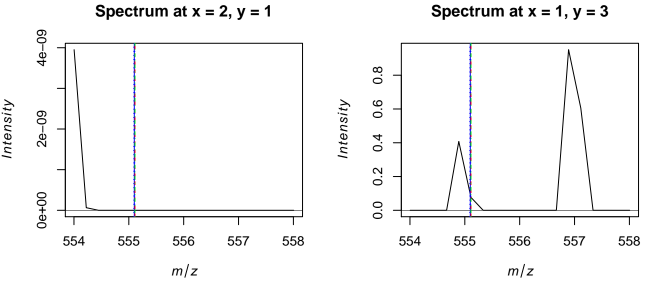


Spectrum at x = 1, y = 3

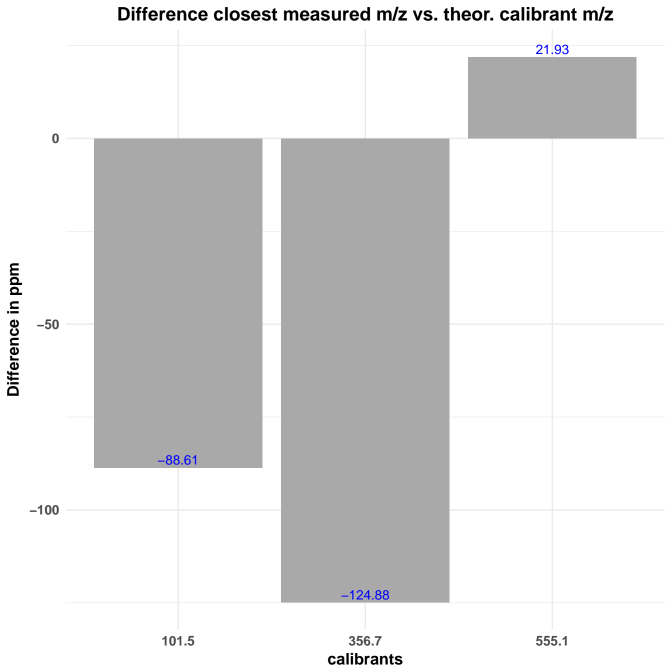


theor. m/z: 101.5 most abundant m/z: 555.1122 closest m/z: 555.1122





Difference m/z with max. average intensity vs. theor. calibrant m/z 21.93 0 Difference in ppm -30 -60 -88.61 -90 101.5 356.7 555.1 calibrants



Difference m/z with max. average intensity vs. theor. m/z (per spectrum)

