protViz: Visualizing and Analyzing Mass Spectrometry Related Data in Proteomics

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Vignette for package version $\geq v.0.1.48$

Recent changes, updates, and new features:

- 0.2.05 export aa2mass
- 0.2.00 peakplot margin bug fix

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1 Preliminary Note

protViz is an R package to do quality checks, visualizations and analysis of mass spectrometry data, coming from proteomics experiments. The package is developed, tested and used at the Functional Genomics Center Zurich. We use this package mainly for prototyping, teaching, and having fun with proteomics data. But it can also be used to do solid data analysis for small scale data sets. Nevertheless, if one is patient, it also handles large data sets.

2 Related Work

The methode of choice in proteomics is mass spectrometry. There are already packages in R which deal with mass spec related data. Some of them are listed here:

- MSnbase package (basic functions for mass spec data including quant aspect with iTRAQ data)
 - http://www.bioconductor.org/packages/release/bioc/html/MSnbase.html
- plgem spectral counting quantification, applicable to MudPIT experiments http://www.bioconductor.org/packages/release/bioc/html/plgem.html
- synapter MSe (Hi3 = Top3 Quantification) for Waters Q-tof data aquired in MSe mode
 - http://www.bioconductor.org/packages/release/bioc/html/synapter.html
- mzR
 http://www.bioconductor.org/packages/release/bioc/html/mzR.html
- isobar iTRAQ/TMT quantification package http://www.bioconductor.org/packages/release/bioc/html/isobar.html
- readMzXmlData http://cran.r-project.org/web/packages/readMzXmlData/
- msQC http://bioconductor.org/packages/3.0/bioc/html/msQC.html

3 Get Data In - Preprocessing

The most time consuming and challenging part for data analysis and visualization is shaping the data that they can easily be processed. In this package, we intentionally left this part away because it is very infrastructure dependent. Moreover we use also commercial tools to analyze data and export the data into R accessible formats. We provide different kind of importers if these formats are available, but with very little effort, one can bring other exports in a similar format which will make it easy to use our package for a variety of tools.

3.1 Identification - In-silico from Proteins to Peptides

For demonstration we use a sequence of peptides derived from a tryptics digest using the Swissprot FETUA_BOVIN Alpha-2-HS-glycoprotein precursor (Fetuin-A) (Asialofetuin) protein.

fcat and tryptic-digest are commandline programs which are included in the package. fcat removes the lines starting with > and all 'new line' character within the protein sequence while tryptic-digest is doing the triptic digest of a protein sequence applying the rule: cleave after arginine (R) and lysine (K) except followed by proline(P).

\$ cat Fetuin.fasta

Ι

MKSFVLLFCLAQLWGCHSIPLDPVAGYKEPACDDPDTEQAALAAVDYINKHLPRGYKHTL NQIDSVKVWPRRPTGEVYDIEIDTLETTCHVLDPTPLANCSVRQQTQHAVEGDCDIHVLK QDGQFSVLFTKCDSSPDSAEDVRKLCPDCPLLAPLNDSRVVHAVEVALATFNAESNGSYL QLVEISRAQFVPLPVSVSVEFAVAATDCIAKEVVDPTKCNLLAEKQYGFCKGSVIQKALG GEDVRVTCTLFQTQPVIPQPQPDGAEAEAPSAVPDAAGPTPSAAGPPVASVVVGPSVVAV PLPLHRAHYDLRHTFSGVASVESSSGEAFHVGKTPIVGQPSIPGGPVRLCPGRIRYFKI

```
$ cat Fetuin.fasta | fcat | tryptic-digest
SFVLLFCLAQLWGCHSIPLDPVAGYK
EPACDDPDTEQAALAAVDYINK
HLPR
GYK
HTLNQIDSVK
VWPR
RPTGEVYDIEIDTLETTCHVLDPTPLANCSVR
QQTQHAVEGDCDIHVLK
QDGQFSVLFTK
CDSSPDSAEDVR
LCPDCPLLAPLNDSR
VVHAVEVALATFNAESNGSYLQLVEISR
AQFVPLPVSVSVEFAVAATDCIAK
EVVDPTK
CNLLAEK
QYGFCK
GSVIQK
ALGGEDVR
VTCTLFQTQPVIPQPQDGAEAEAPSAVPDAAGPTPSAAGPPVASVVVGPSVVAVPLPLHR
AHYDLR.
HTFSGVASVESSSGEAFHVGK
TPIVGQPSIPGGPVR
LCPGR
IR
YFK
```

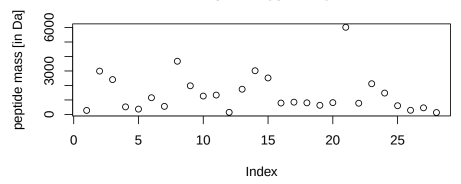
4 Peptide Identification

The currency in proteomics are the peptides. In proteomics, proteins are digested to so-called peptides since peptides are much easier to handle biochemically than proteins. Proteins are very different in nature some are very sticky while others are soluble in aqueous solutions while again are only sitting in membranes. Therefore, proteins are chopped up into peptides because it is fair to assume, that for each protein, there will be a number of peptides behaving well, so that they can actually be measured with the mass spectrometer. This step introduces another problem, the so-called protein inference problem. In this package here, we do not at all touch upon the protein inference.

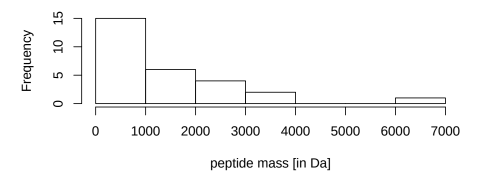
4.1 Computing the Parent Ion Mass

```
> library(protViz)
> op<-par(mfrow=c(1,1))
> fetuin<-c('MK', 'SFVLLFCLAQLWGCHSIPLDPVAGYK',
+ 'EPACDDPDTEQAALAAVDYINK',
+ 'HLPR', 'GYK', 'HTLNQIDSVK', 'VWPR',
+ 'RPTGEVYDIEIDTLETTCHVLDPTPLANCSVR',
+ 'QQTQHAVEGDCDIHVLK', 'QDGQFSVLFTK',
+ 'CDSSPDSAEDVR', 'K', 'LCPDCPLLAPLNDSR',
+ 'VVHAVEVALATFNAESNGSYLQLVEISR',
+ 'AQFVPLPVSVSVEFAVAATDCIAK',
+ 'EVVDPTK', 'CNLLAEK', 'QYGFCK',
+ 'GSVIQK', 'ALGGEDVR',
+ 'VTCTLFQTQPVIPQPQPDGAEAEAPSAVPDAAGPTPSAAGPPVASVVVGPSVVAVPLPLHR',
+ 'AHYDLR', 'HTFSGVASVESSSGEAFHVGK',
+ 'TPIVGQPSIPGGPVR', 'LCPGR', 'IR', 'YFK', 'I')
> (pm<-parentIonMass(fetuin))</pre>
     278.1533 2991.5259 2406.0765 522.3147
                                              367.1976 1154.6164 557.3194
 [8] 3671.7679 1977.9447 1269.6474 1337.5274 147.1128 1740.8407 3016.5738
[15] 2519.3214 787.4196 847.4342 802.3552
                                              631.3773 816.4210 6015.1323
[22] 774.3893 2120.0043 1474.8376 602.3079
                                              288.2030 457.2445 132.1019
> op < -par(mfrow = c(2,1))
> plot(pm, ylab="peptide mass [in Da]",
      main="Fetuin Peptide tryptic digested.")
> hist(pm, xlab="peptide mass [in Da]")
```

Fetuin Peptide tryptic digested.



Histogram of pm



4.2 In-silico Peptide Fragmentation

The fragment ions of a peptide can be computed following the rules proposed in [4]. Beside the b and y ions the FUN argument of fragmentIon defines which ions are computed. the default ions beeing computed are defined in the function defaultIon. The are no limits for defining other forms of fragment ions for ETD (c and z ions) CID (b and y ions).

> defaultIon

```
function (b, y)
{
    Hydrogen <- 1.007825
    Oxygen <- 15.994915
    Nitrogen <- 14.003074
    c <- b + (Nitrogen + (3 * Hydrogen))
    z <- y - (Nitrogen + (3 * Hydrogen))
    return(cbind(b, y, c, z))
}
<environment: namespace:protViz>
> peptides<-c('HTLNQIDSVK', 'ALGGEDVR', 'TPIVGQPSIPGGPVR')
> pim<-parentIonMass(peptides)</pre>
```

```
> fi<-fragmentIon(peptides)</pre>
> par(mfrow=c(3,1));
> for (i in 1:length(peptides)){
       plot(0,0,
+
           xlab='m/Z',
            ylab='',
+
            xlim=range(c(fi[i][[1]]$b,fi[i][[1]]$y)),
            ylim=c(0,1),
+
           type='n',
+
           axes=FALSE,
            sub=paste( pim[i], "Da"));
+
       box()
       axis(1,fi[i][[1]]$b,round(fi[i][[1]]$b,2))
       pepSeq<-strsplit(peptides[i],"")</pre>
+
       axis(3,fi[i][[1]]$b,pepSeq[[1]])
+
       abline(v=fi[i][[1]]$b, col='red',lwd=2)
+
       abline(v=fi[i][[1]]$c, col='orange')
       abline(v=fi[i][[1]]$y, col='blue',lwd=2)
+
       abline(v=fi[i][[1]]$z, col='cyan')
+ }
                        466.24
                                           822.41
                  352.2
                                     707.38
                                                     1008.51
      138.07 239.11
                               594.3
                                                            1136.61
                                   m/Z
                               1154.616401 Da
                                  Е
                    G
                        G
                                           D
                L
                                                642.31
      72.04
              185.13
                       299.17
                                 428.21
                                         543.24
                                                            798.41
                               816.420981 Da
                    V G
                           Q
                                           P G G
      102.05
              312.19
                     468.28 596.34
                                 780.43
                                          990.56
                                                  1201.66
                                                            1456.83
                               1474.837601 Da
```

The next lines compute the singly and doubly charged fragment ions of the HTLNQIDSVK peptide. Which are usually the ones that can be used to make an identification.

```
> Hydrogen<-1.007825
> (fi.HTLNQIDSVK.1<-fragmentIon('HTLNQIDSVK'))[[1]]</pre>
           b
                                         z
                     У
1
    138.0662
             147.1128
                       155.0927 130.0863
2
    239.1139
             246.1812 256.1404 229.1547
3
   352.1979 333.2132 369.2245 316.1867
4
    466.2409 448.2402 483.2674 431.2136
5
    594.2994
             561.3242 611.3260 544.2977
6
   707.3835 689.3828 724.4100 672.3563
7
    822.4104 803.4258 839.4370 786.3992
8
   909.4425 916.5098 926.4690 899.4833
   1008.5109 1017.5575 1025.5374 1000.5309
10 1136.6058 1154.6164 1153.6324 1137.5899
> (fi.HTLNQIDSVK.2<-(fi.HTLNQIDSVK.1[[1]] + Hydrogen) / 2)</pre>
           b
                               С
1
    69.53701 74.06031
                       78.05028 65.54704
  120.06085 123.59452 128.57412 115.08124
  176.60288 167.11053 185.11615 158.59726
  233.62434 224.62400 242.13761 216.11073
5
  297.65363 281.16603 306.16691 272.65276
  354.19566 345.19532 362.70894 336.68205
7
  411.70913 402.21679 420.22241 393.70351
8 455.22515 458.75882 463.73842 450.24554
  504.75935 509.28266 513.27262 500.76938
10 568.80683 577.81211 577.32010 569.29884
```

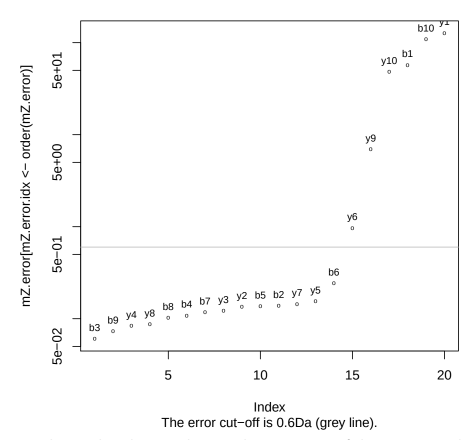
4.3 Peptide Sequence - Fragment Ion Matching

Given a peptide sequence and a tandem mass spectrum. For the assignment of a canditate peptide an in-silico fragment ion spectra fi is computed. The function findNN determines for each fragment ion the closesed peak in the MS2. If the difference between the in-silico mass and the measured mass is inside the 'accuracy' mass window of the mass spec device the in-silico fragment ion is considered as potential hit.

```
peptideSequence<-'HTLNQIDSVK'
>
>
      spec<-list(scans=1138,</pre>
          title="178: (rt=22.3807) [20080816_23_fetuin_160.RAW]",
+
          rtinseconds=1342.8402,
+
+
          charge=2,
+
          mZ=c(195.139940, 221.211970, 239.251780, 290.221750,
      316.300770, 333.300050, 352.258420, 448.384360, 466.348830,
+
      496.207570, 509.565910, 538.458310, 547.253380, 556.173940,
+
```

```
560.358050, 569.122080, 594.435500, 689.536940, 707.624790,
+
      803.509240, 804.528220, 822.528020, 891.631250, 909.544400,
+
      916.631600, 973.702160, 990.594520, 999.430580, 1008.583600,
+
      1017.692500, 1027.605900),
+
          intensity=c(931.8, 322.5, 5045, 733.9, 588.8, 9186, 604.6,
+
      1593, 531.8, 520.4, 976.4, 410.5, 2756, 2279, 5819, 2.679e+05,
+
      1267, 1542, 979.2, 9577, 3283, 9441, 1520, 1310, 1.8e+04,
+
      587.5, 2685, 671.7, 3734, 8266, 3309))
+
>
      fi<-fragmentIon(peptideSequence)</pre>
>
      n<-nchar(peptideSequence)</pre>
      by.mZ<-c(fi[[1]]$b, fi[[1]]$y)
>
      by.label<-c(paste("b",1:n,sep=''), paste("y",n:1,sep=''))</pre>
>
>
      # should be a R-core function as findInterval!
      idx<-findNN(by.mZ, spec$mZ)</pre>
>
>
      mZ.error<-abs(spec$mZ[idx]-by.mZ)</pre>
>
      plot(mZ.error[mZ.error.idx<-order(mZ.error)],</pre>
+
          main="Error Plot",
+
          pch='o',
+
          cex=0.5,
+
          sub='The error cut-off is 0.6Da (grey line).',
+
          log='y')
      abline(h=0.6,col='grey')
>
>
      text(1:length(by.label),
          mZ.error[mZ.error.idx],
+
          by.label[mZ.error.idx],
+
          cex=0.75, pos=3)
```

Error Plot



The graphic above is showing the mass error of the assingment between the MS2 spec and the singly charged fragment ions of HTLNQIDSVK. The function psm is doing the peptide sequence matching. Of course, the more theoretical ions match (up to a small error tolerance, given by the system) the actually measured ion series, the more likely it is, that the measured spectrum indeed is from the inferred peptide (and therefore the protein is identified)

4.4 Modifications

```
[[2]]
[4] "000000000000100100"
> fi<-fragmentIon(c('TAFDEAIAELDTLSEESYK',</pre>
     'TAFDEAIAELDTLNEESYK', 'TAFDEAIAELDTLSEESYK',
     'TAFDEAIAELDTLNEESYK'),
+
         modified=c('0000000000000200000',
+
         '0000000000000000000'),
     modification=m$mono)
+
> #bh<-c('TAFDEAIAELDTLNEESYK', 'TAFDEAIAELDTLSEESYK')
> #fi<-fragmentIon(rep('HTLNQIDSVK',2),
      modified=c('0000000100','0000000000'),
> #
      modification=m[,2])
     Labeling Peaklists
4.5
The labeling of the spectra can be done with the peakplot function.
> data(msms)
> op < -par(mfrow = c(2,1))
> peakplot("TAFDEAIAELDTLNEESYK", msms[[1]])
$mZ.Da.error
[1] 232.331344 161.294234 14.225824
                                      -0.032616
                                                -0.143306 0.032244
[7]
      0.054604 -0.004076 -0.071746
                                      -0.084536 -0.097076 -0.038856
[13]
     -0.061816 0.004554 -0.122336
                                     -0.139626 -1.071256 -18.783686
[19] -146.878646 187.273499 24.210169 0.048669 0.177779 0.027939
[25]
     0.049579 0.052379 0.044579 0.036749 0.043189 -0.035101
[31]
    -0.061011
                 0.000729 -0.092081
                                       2.011029 -8.412111
                                                           7.195579
[37] -63.841531 -164.889211 215.304795 144.267685 -2.800725 -17.059165
                                       1.292875 -0.003965 -13.612585
[43]
     2.034875
                 2.264105 4.008125
[49]
     -0.060925 -17.065405
                                       3.000405 -17.148885 -17.166175
                            3.897535
[55] -18.097805 -35.810235 -163.905195 204.300048 41.236718 17.075218
[61]
    -0.843372 -1.091812 0.129908 17.078928 -0.372162 -16.539502
[67]
    -1.044962 -1.000952 -1.409062
                                                16.934468
                                      -2.995122
                                                           19.037578
                24.222128 -46.814982 -147.862662
[73]
    8.614438
$mZ.ppm.error
[1] 2.276532e+06 9.318407e+05 4.443342e+04 -7.494702e+01 -2.539851e+02
[6] 5.075660e+01 7.296574e+01 -4.974443e+00 -7.564705e+01 -7.963713e+01
[11] -8.250960e+01 -3.041352e+01 -4.445040e+01 3.026484e+00 -7.488007e+01
```

[16] -7.920687e+01 -5.791093e+02 -9.331667e+03 -6.860308e+04 1.272993e+06 [21] 7.805297e+04 1.225277e+02 3.378218e+02 4.263587e+01 6.444386e+01 [26] 5.935833e+01 4.532837e+01 3.345395e+01 3.564687e+01 -2.618263e+01

\$idx

[1]	1	1	1	3	14	21	38	49	64	87	91	97	102	106	110	113	115	116	116	
[20]	1	1	2	12	25	41	53	70	89	94	99	104	107	108	111	114	116	116	116	
[39]	1	1	1	3	16	24	41	52	67	88	93	97	104	107	110	113	115	116	116	
[58]	1	1	2	11	22	40	53	68	88	93	98	103	106	108	111	114	116	116	116	

\$label

```
[1] "b1" "b2" "b3" "b4"
                           "b5" "b6" "b7" "b8"
                                                  "b9"
                                                       "b10" "b11" "b12"
[13] "b13" "b14" "b15" "b16" "b17" "b18" "b19" "y1"
                                                  "y2"
                                                       "y3" "y4"
                                                                   "y5"
          "y7"
                           "y10" "y11" "y12" "y13" "y14" "y15" "y16" "y17"
[25] "y6"
               "y8" "y9"
[37] "v18" "v19" "c1"
                     "c2"
                           "c3"
                                "c4" "c5" "c6" "c7" "c8" "c9"
[49] "c11" "c12" "c13" "c14" "c15" "c16" "c17" "c18" "c19" "z1"
                                                             "z2"
                                                                   "z3"
                           "z8" "z9" "z10" "z11" "z12" "z13" "z14" "z15"
[61] "z4" "z5" "z6" "z7"
[73] "z16" "z17" "z18" "z19"
```

\$score

[1] -1

\$sequence

[1] "TAFDEAIAELDTLNEESYK"

\$fragmentIon

	b	У	С	Z
1	102.0550	147.1128	119.0815	130.0863
2	173.0921	310.1761	190.1186	293.1496
3	320.1605	397.2082	337.1870	380.1816
4	435.1874	526.2508	452.2140	509.2242
5	564.2300	655.2933	581.2566	638.2668
6	635.2671	769.3363	652.2937	752.3097
7	748.3512	882.4203	765.3777	865.3938
8	819.3883	983.4680	836.4148	966.4415
9	948.4309	1098.4950	965.4574	1081.4684
10	1061.5149	1211.5790	1078.5415	1194.5525
11	1176.5419	1340.6216	1193.5684	1323.5951
12	1277.5896	1411.6587	1294.6161	1394.6322
13	1390.6736	1524.7428	1407.7002	1507.7162

```
14 1504.7165 1595.7799 1521.7431 1578.7533
15 1633.7591 1724.8225 1650.7857 1707.7959
16 1762.8017 1839.8494 1779.8283 1822.8229
17 1849.8338 1986.9178 1866.8603 1969.8913
18 2012.8971 2057.9549 2029.9236 2040.9284
19 2140.9920 2159.0026 2158.0186 2141.9761
> peakplot("TAFDEAIAELDTLSEESYK", msms[[2]])
$mZ.Da.error
 [1]
     245.264254 174.227144
                              27.158734
                                           14.444434
                                                       0.021404
                                                                  -0.111266
 [7]
      -0.039926
                 -0.021626
                              -0.121916
                                          -8.079236
                                                      -0.158376
                                                                  -0.153156
[13]
                 -0.022946
                              -0.186736
                                          -0.092226
                                                      -0.120456
                                                                  -0.151686
      -0.094316
[19] -128.246646 200.206409
                              37.143079
                                           0.078909
                                                       0.062269
                                                                   0.129769
[25]
       0.103729
                   0.060869
                              -0.051451 -18.048351
                                                      -0.027511
                                                                  -0.025601
[31]
      -0.006211
                   0.020529
                             -0.048781
                                          -0.024771
                                                      -9.166311
                                                                   6.953579
[37]
     -45.209531 -146.257211
                             228.237705 157.200595
                                                      10.132185
                                                                  -2.582115
[43]
       1.626855
                   2.722405
                               9.009025
                                          -1.130895
                                                       1.216385
                                                                  13.347315
[49]
                   0.960295 -17.120865
                                                     -17.213285 -17.118775
      -3.671525
                                           3.020205
[55]
     -17.147005 -17.178235 -145.273195 217.232958
                                                      54.169628
                                                                  17.105458
[61]
                 -1.260332
                              -0.899352
                                         -3.098942
                                                      -1.173512
                                                                  -1.021802
      -0.833452
[67]
      -0.939162
                  -1.007752
                             -1.377062
                                          -3.022622
                                                      16.977768
                                                                  17.001778
[73]
       7.860238
                  23.980128 -28.182982 -129.230662
$mZ.ppm.error
[6] -1.751484e+02 -5.335196e+01 -2.639286e+01 -1.285450e+02 -7.611043e+03
[11] -1.346114e+02 -1.198789e+02 -6.782037e+01 -1.552813e+01 -1.162198e+02
[16] -5.313198e+01 -6.608212e+01 -7.638202e+01 -6.066594e+04 1.360904e+06
[21] 1.197483e+05 1.986591e+02 1.183257e+02 1.980319e+02 1.397352e+02
[26]
```

- [1] 2.403257e+06 1.006558e+06 8.482850e+04 3.319130e+04 3.793488e+01

- 7.115774e+01 -5.379332e+01 -1.684426e+04 -2.322450e+01 -1.948903e+01
- [31] -4.485617e+00 1.370673e+01 -3.109508e+01 -1.458996e+01 -5.056331e+03
- [36] 3.547913e+03 -2.226035e+04 -6.860121e+04 1.916651e+06 8.268554e+05
- [41] 3.004915e+04 -5.709941e+03 2.798859e+03 4.173588e+03 1.177069e+04
- [46] -1.352074e+03 1.259905e+03 1.237534e+04 -3.076091e+03 7.417604e+02
- [51] -1.216230e+04 2.020566e+03 -1.060078e+04 -9.766434e+03 -9.319787e+03
- [56] -8.576627e+03 -6.817113e+04 1.669915e+06 1.847849e+05 4.499286e+04
- [61] -1.636709e+03 -1.974616e+03 -1.239974e+03 -3.696333e+03 -1.249174e+03
- [66] -9.690310e+02 -8.043928e+02 -7.772361e+02 -1.006903e+03 -2.041339e+03 [71] 1.094110e+04 1.011538e+04 4.376983e+03 1.234257e+04 -1.399411e+04
- [76] -6.110297e+04

\$idx

[1] 39 45 64 90 96 106 116 121 126 129 131 133 133 1 1 1 3 11 20 [20] 1 2 7 24 38 49 65 90 97 110 115 122 123 127 130 132 133 133 47 98 108 116 122 126 129 131 133 133 [39] 1 1 1 3 13 23 40 67 2 6 21 36 62 90 95 108 113 121 123 127 130 132 133 133 [58] 47

\$label

```
[1] "b1" "b2" "b3" "b4" "b5" "b6" "b7" "b8" "b9" "b10" "b11" "b12" [13] "b13" "b14" "b15" "b16" "b17" "b18" "b19" "y1" "y2" "y3" "y4" "y5" [25] "y6" "y7" "y8" "y9" "y10" "y11" "y12" "y13" "y14" "y15" "y16" "y17" [37] "y18" "y19" "c1" "c2" "c3" "c4" "c5" "c6" "c7" "c8" "c9" "c10" [49] "c11" "c12" "c13" "c14" "c15" "c16" "c17" "c18" "c19" "z1" "z2" "z3" [61] "z4" "z5" "z6" "z7" "z8" "z9" "z10" "z11" "z12" "z13" "z14" "z15" [73] "z16" "z17" "z18" "z19"
```

\$score

[1] -1

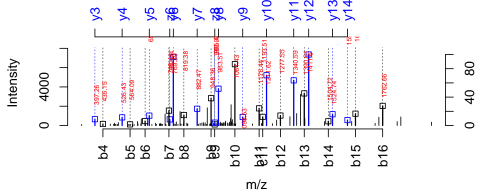
\$sequence

[1] "TAFDEAIAELDTLSEESYK"

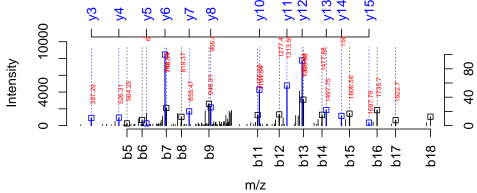
\$fragmentIon

```
b
                    У
    102.0550
             147.1128 119.0815 130.0863
2
   173.0921
             310.1761
                       190.1186 293.1496
3
   320.1605 397.2082 337.1870 380.1816
4
   435.1874 526.2508 452.2140 509.2242
   564.2300 655.2933 581.2566 638.2668
5
6
   635.2671 742.3254 652.2937 725.2988
7
   748.3512 855.4094 765.3777 838.3829
   819.3883 956.4571 836.4148 939.4306
8
   948.4309 1071.4841 965.4574 1054.4575
10 1061.5149 1184.5681 1078.5415 1167.5416
11 1176.5419 1313.6107 1193.5684 1296.5842
12 1277.5896 1384.6478 1294.6161 1367.6213
13 1390.6736 1497.7319 1407.7002 1480.7053
14 1477.7056 1568.7690 1494.7322 1551.7424
15 1606.7482 1697.8116 1623.7748 1680.7850
16 1735.7908 1812.8385 1752.8174 1795.8120
17 1822.8229 1959.9069 1839.8494 1942.8804
18 1985.8862 2030.9440 2002.9127 2013.9175
19 2113.9811 2131.9917 2131.0077 2114.9652
```

> par(op)



Scan 3246 (rt=67.4676) [/p474/Proteomics/ORBI_1/jonas_20080530_bhdaten_de



Scan 3353 (rt=70.3158) [/p474/Proteomics/ORBI_1/jonas_20080530_bhdaten_dc

5 Quantification

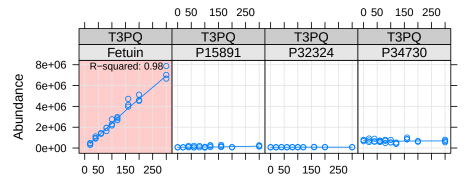
For an overview on Quantitative Proteomics read [1, 2]. The authors are aware that meaningful statistics usually require much higher number of biological replicates. In almost all cases there are not more than three to six repitions. For the moment there are limited options due to the availability of machine time and the limits of the technologies.

5.1 Label-free methods on protein level

The data set fetuinLFQ contains a subset of our results descriped in [3]. The example below shows a visualization using trellis plots. It graphs the abundance of four protein in dependency from the fetuin concentration spiked into the sample.

> library(lattice)
> data(fetuinLFQ)
> cv<-1-1:7/10
> t<-trellis.par.get("strip.background")
> t\$col<-(rgb(cv,cv,cv))
> trellis.par.set("strip.background",t)
> print(xyplot(abundance~conc|prot*method,
+ groups=prot,

```
xlab="Fetuin concentration spiked into experiment [fmol]",
+
      ylab="Abundance",
      aspect=1,
      data=fetuinLFQ$t3pq[fetuinLFQ$t3pq$prot
+
          %in% c('Fetuin', 'P15891', 'P32324', 'P34730'),],
     panel = function(x, y, subscripts, groups) {
+
          if (groups[subscripts][1] == "Fetuin") {
              panel.fill(col="#ffcccc")
+
          }
          panel.grid(h=-1,v=-1)
         panel.xyplot(x, y)
         panel.loess(x,y, span=1)
          if (groups[subscripts][1] == "Fetuin") {
              panel.text(min(fetuinLFQ$t3pq$conc),
                  max(fetuinLFQ$t3pq$abundance),
                  paste("R-squared:",
                  round(summary(lm(x~y))$r.squared,2)),
                  cex=0.75,
                  pos=4)
          }
      }
+ ))
```



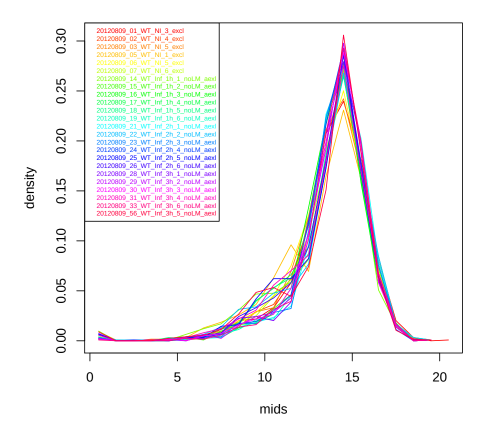
Fetuin concentration spiked into experiment [fmol]

The plot shows the estimated concentration of the four proteins using the top three most intense peptides. The Fetuin peptides are spiked in with increasing concentration while the three other yeast proteins are kept stable in the background.

5.2 pgLFQ – LCMS based label-free quantification

LCMS based label-free quantification is a very popular method to extract relative quantitative information from mass spectrometry experiments. At the FGCZ we use the software ProgenesisLCMS for this workflow http://www.nonlinear.com/products/progenesis/lc-ms/overview/. Progenesis is a graphical software which does the aligning between several LCMS experiments, extracts signal intensities from LCMS maps and annotates the mastermap with peptide and protein labels.

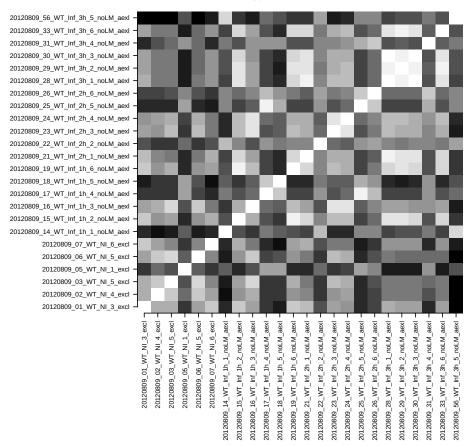
```
> data(pgLFQfeature)
> data(pgLFQprot)
> featureDensityPlot<-function(data, n=ncol(data), nbins=30){</pre>
      my.col<-rainbow(n);</pre>
      mids<-numeric()</pre>
+
      density<-numeric()</pre>
      for (i in 1:n) {
+
           h<-hist(data[,i],nbins, plot=FALSE)
           mids<-c(mids, h$mids)</pre>
+
           density<-c(density, h$density)</pre>
      }
+
+
      plot(mids,density, type='n')
      for (i in 1:n) {
+
          h<-hist(data[,i],nbins, plot=FALSE)</pre>
           lines(h$mids,h$density, col=my.col[i])
      }
+
      legend("topleft", names(data), cex=0.5,
+
           text.col=my.col
      )
+
+ }
> par(mfrow=c(1,1));
> featureDensityPlot(asinh(pgLFQfeature$"Normalized abundance"),
      nbins=25)
```



The featureDensityPlot shows the normalized signal intensity distribution (asinh transformed) over 24 LCMS runs which are aligned in this experiment.

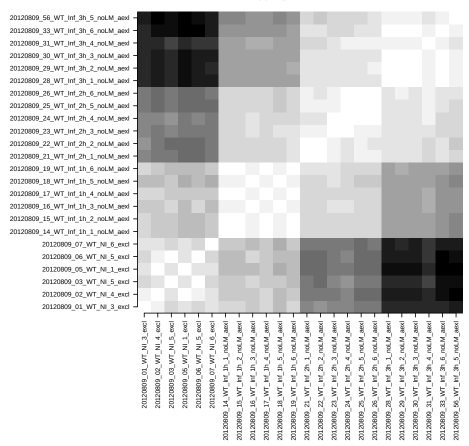
```
> op<-par(mfrow=c(1,1),mar=c(18,18,4,1),cex=0.5)
> samples<-names(pgLFQfeature$"Normalized abundance")
> image(cor(asinh(pgLFQfeature$"Normalized abundance")),
+ col=gray(seq(0,1,length=20)),
+ main='pgLFQfeature correlation',
+ axes=FALSE)
> axis(1,at=seq(from=0, to=1,
+ length.out=length(samples)),
+ labels=samples, las=2)
> axis(2,at=seq(from=0, to=1,
+ length.out=length(samples)), labels=samples, las=2)
> par(op)
```

pgLFQfeature correlation

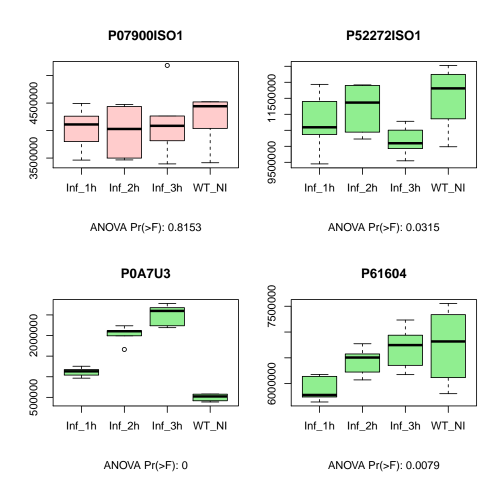


This image plot shows the correlation between runs on feature level (values are asinh transformed). White is perfect correlation while black indicates a poor correlation.

pgLFQprot correlation



This figure shows the correlation between runs on protein level (values are asinh transformed). White is perfect correlation while black indicates a poor correlation. Stricking is the fact that the six biological replicates for each condition cluster very well.

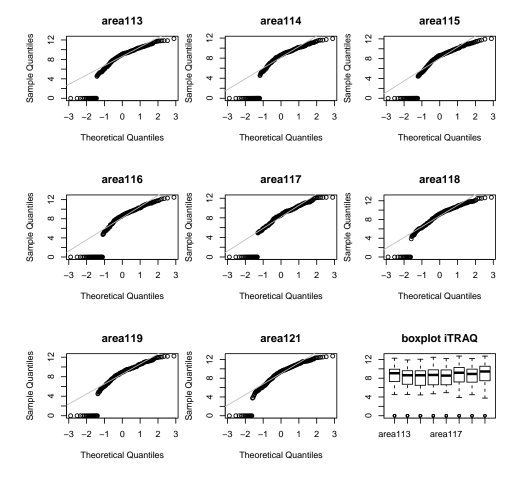


This figure shows the result for four proteins which either differ significantly in expression accross conditions (green boxplots) using an analysis of variance test, or non differing protein expression (red boxplot).

5.3 iTRAQ – Two Group Analysis

The data for the next section is an iTRAQ-8-plex experiment where two conditions are compared (each condition has 4 biological replicates)

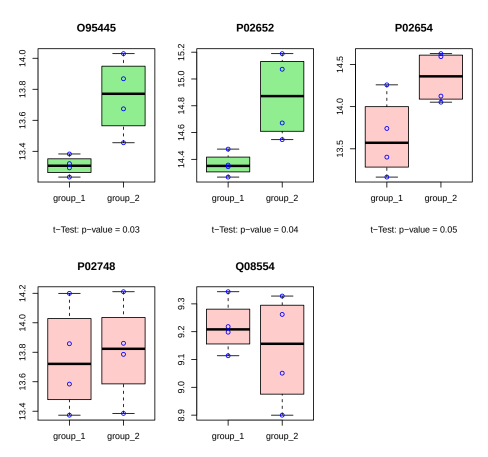
5.3.1 Sanity Check



A first quality check to see if all reporter ion channels are having the same distributions. Shown in the figure are Q-Q plots of the individual reporter channels against a normal distribution. The last is a boxplot for all individual channels.

5.3.2 On Protein Level

```
boxplot.color='lightgreen'
+
+
      b<-boxplot(as.numeric(group1Protein[i,]),</pre>
          as.numeric(group2Protein[i,]),
+
          main=row.names(group1Protein)[i],
          sub=paste("t-Test: p-value =", round(tt.p_value,2)),
+
          col=boxplot.color,
          axes=FALSE)
+
      axis(1, 1:2, c('group_1', 'group_2')); axis(2); box()
+
      points(rep(1,b$n[1]), as.numeric(group1Protein[i,]), col='blue')
+
      points(rep(2,b$n[2]), as.numeric(group2Protein[i,]), col='blue')
+ }
```



This figure shows five proteins which are tested if they differ accross conditions using the four biological replicates with a t-test.

t-Test: p-value = 0.49

5.3.3 On Peptide Level

t-Test: p-value = 0.82

The same can be done on peptide level using the protViz function iTRAQ2GroupAnalysis.

```
> data(iTRAQ)
```

> q<-iTRAQ2GroupAnalysis(data=iTRAQ,

```
group1=c(3,4,5,6),
+
+
      group2=7:10,
      INDEX=paste(iTRAQ$prot,iTRAQ$peptide),
      plot=FALSE)
> q[1:10,]
                              name p_value Group1.area113 Group1.area114
                   095445 AFLLTPR
                                     0.056
1
                                                   1705.43
                                                                    1459.10
2
                   095445 DGLCVPR
                                     0.161
                                                   2730.41
                                                                   1852.90
3
                 095445 MKDGLCVPR
                                     0.039
                                                  28726.38
                                                                  15409.81
4
                095445 NQEACELSNN
                                     0.277
                                                   4221.31
                                                                   4444.28
5
                 095445 SLTSCLDSK
                                     0.036
                                                  20209.66
                                                                  14979.02
6
     PO2652 AGTELVNFLSYFVELGTQPA
                                     0.640
                                                   4504.97
                                                                   4871.88
7
    PO2652 AGTELVNFLSYFVELGTQPAT
                                     0.941
                                                  67308.30
                                                                  46518.21
   PO2652 AGTELVNFLSYFVELGTQPATQ
                                     0.338
                                                   4661.54
                                                                   3971.82
9
     PO2652 EPCVESLVSQYFQTVTDYGK
                                     0.115
                                                   4544.56
                                                                   4356.51
10
                  P02652 EQLTPLIK
                                     0.053
                                                                  22015.94
                                                  24596.42
   Group1.area115 Group1.area116 Group2.area117 Group2.area118 Group2.area119
1
           770.65
                           3636.40
                                           3063.48
                                                           4046.73
                                                                           2924.49
2
          1467.65
                           2266.88
                                           2269.57
                                                           3572.32
                                                                           2064.82
3
         19050.13
                         58185.02
                                          51416.05
                                                          70721.05
                                                                          38976.42
4
          2559.23
                          6859.71
                                           5545.12
                                                          11925.66
                                                                           6371.50
5
         12164.94
                         37572.56
                                          30687.57
                                                          39176.99
                                                                          34417.66
6
          2760.53
                           9213.41
                                           6728.62
                                                          14761.96
                                                                           7796.29
7
         33027.14
                        111629.30
                                          94531.76
                                                         168775.00
                                                                          83526.72
8
          2564.39
                          8269.73
                                           6045.30
                                                          13724.92
                                                                           7426.84
9
          2950.48
                          6357.90
                                           6819.99
                                                          10265.84
                                                                           7012.92
         18424.56
10
                         49811.91
                                          33197.47
                                                          67213.62
                                                                          40030.86
   Group2.area121
1
          5767.87
2
          2208.92
3
         60359.72
4
         15656.92
5
         54439.22
6
         18681.60
7
        168032.50
8
         17214.87
9
         14279.22
10
         87343.38
```

6 Pressure Profiles QC

A common problem with mass spec setup is the pure reliability of the high pressure pump. The following graphics provide visualizations for quality control.

On overview of the pressure profile data can be seen by using the ppp function.

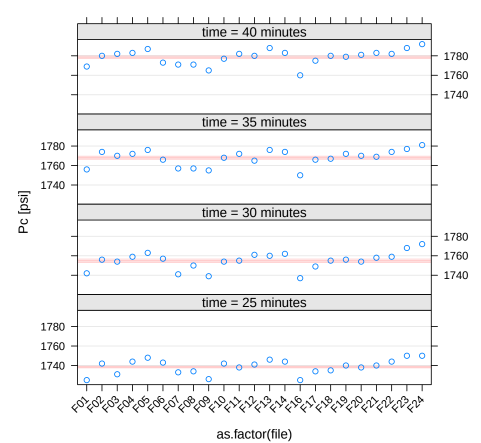
```
> data(pressureProfile)
```

> ppp(pressureProfile)

The lines plots the pressure profiles data on a scatter plot 'Pc' versus 'time' grouped by time range (no figure because of too many data items).

The Trellis xyplot shows the Pc development over each instrument run to a specified relative run time (25,30,...).

```
> pp.data<-pps(pressureProfile, time=seq(25,40,by=5))</pre>
 print(xyplot(Pc ~ as.factor(file) | paste("time =",
      as.character(time), "minutes"),
      panel = function(x, y){
+
          m < -sum(y)/length(y)
          m5 < -(max(y) - min(y)) * 0.05
          panel.abline(h=c(m-m5,m,m+m5),
              col=rep("#ffcccc",3),lwd=c(1,2,1))
          panel.grid(h=-1, v=0)
          panel.xyplot(x, y)
      },
+
      ylab='Pc [psi]',
      layout=c(1,4),
+
      sub='The three red lines indicate the average plus min 5%.',
      scales = list(x = list(rot = 45)),
+
      data=pp.data))
```



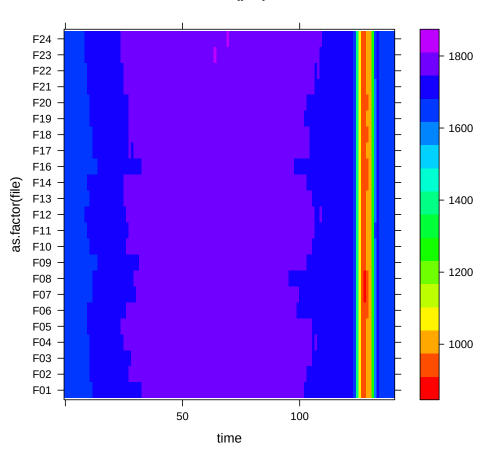
The three red lines indicate the average plus min 5%.

While each panel in the xyplot above shows the data to a given point in time, we try to use the levelplot to get an overview of the whole pressure profile data.

```
> pp.data<-pps(pressureProfile, time=seq(0,140,length=128))</pre>
```

- > print(levelplot(Pc ~ time * as.factor(file),
- main='Pc(psi)',
- + data=pp.data,
- + col.regions=rainbow(100)[1:80]))

Pc(psi)



7 Session information

> sessionInfo()

R Under development (unstable) (2014-09-17 r66626) Platform: x86_64-unknown-linux-gnu (64-bit)

locale:

[1]	LC_CTYPE=en_US.UTF-8	LC_NUMERIC=C
[3]	LC_TIME=en_US.UTF-8	LC_COLLATE=C

[5] LC_MONETARY=en_US.UTF-8 LC_MESSAGES=en_US.UTF-8

[7] LC_PAPER=en_US.UTF-8 LC_NAME=C

[9] LC_ADDRESS=C LC_TELEPHONE=C

```
[11] LC_MEASUREMENT=en_US.UTF-8 LC_IDENTIFICATION=C
attached base packages:
[1] stats
              graphics grDevices utils datasets methods
                                                                base
other attached packages:
[1] lattice_0.20-29 protViz_0.2.06
loaded via a namespace (and not attached):
[1] Rcpp_0.11.2 grid_3.2.0 tools_3.2.0
> packageDescription('protViz')
Package: protViz
Type: Package
Title: Visualizing and Analyzing Mass Spectrometry Related Data in
       Proteomics
Version: 0.2.06
Date: 2014-09-18
Author: Christian Panse <cp@fgcz.ethz.ch>, Jonas Grossmann
        <jg@fgcz.ethz.ch>, Simon Barkow-Oesterreicher <sb@fgcz.ethz.ch>
Maintainer: Christian Panse <cp@fgcz.ethz.ch>
Depends: R (>= 3.0.2), methods
Imports: Rcpp (>= 0.9.9)
Suggests: lattice, RUnit, BiocGenerics
Description: This R package helps with quality checks, visualizations
        and analysis of mass spectrometry data, coming from proteomics
        experiments. The package is developed, tested and used at the
        Functional Genomics Center Zurich. We use this package mainly
        for prototyping, teaching, and having fun with proteomics data.
        But it can also be used to do solid data analysis for small
        scale data sets.
License: GPL-3
URL: http://fgcz-data.uzh.ch/~cpanse/protViz/
Collate: aa2mass.R deisotoper.R de_novo.R findNN_.R findNN.R
        fragmentIon.R .....
LazyData: true
Built: R 3.2.0; x86_64-unknown-linux-gnu; 2014-09-18 07:02:00 UTC; unix
```

References

[1] M. Bantscheff, S. Lemeer, M. M. Savitski, and B. Kuster. Quantitative mass spectrometry in proteomics: critical review update from 2007 to the present. *Anal Bioanal Chem*, 404(4):939–965, Sep 2012.

-- File: /scratch/RtmpGo1mnA/Rinst48896ccade22/protViz/Meta/package.rds

- [2] S. Cappadona, P. R. Baker, P. R. Cutillas, A. J. Heck, and B. van Breukelen. Current challenges in software solutions for mass spectrometry-based quantitative proteomics. *Amino Acids*, 43(3):1087–1108, Sep 2012.
- [3] J. Grossmann, B. Roschitzki, C. Panse, C. Fortes, S. Barkow-Oesterreicher, D. Rutishauser, and R. Schlapbach. Implementation and evaluation of relative and absolute quantification in shotgun proteomics with label-free methods. *J Proteomics*, 73(9):1740–1746, Aug 2010. [DOI:10.1016/j.jprot.2010.05.011] [PubMed:20576481].
- [4] P. Roepstorff and J. Fohlman. Proposal for a common nomenclature for sequence ions in mass spectra of peptides. *Biomed. Mass Spectrom.*, 11(11):601, Nov 1984. [DOI:10.1002/bms.1200111109] [PubMed:6525415].