## Portable Proteomics Pipeline (P3) Labelled (iTRAQ4) Benchmark

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April 29, 2016

This document is to benchmark and as a sanity check to the iTRAQ4 method for the Portable Proteomics Pipelines (P3) containers.

## Results from PNNL

For this first purpose, 1 first file TCGA calculated by PNNL used as the comparasion:

```
show(mzml.sp)
## [1] "TCGA_13-1489_42-2590_36-2529_117C_W_PNNL_B2S5_f01_ReporterIons.txt"
dim(msexp.sp) # The size of table
## [1] 16058
                12
head(msexp.sp)[1:3,]
##
     Dataset ScanNumber Collision. Mode ParentIonMZ BasePeakIntensity
## 1
      322257
                      48
                                              329.45
                                                              13848.50
                                    hcd
## 2
     322257
                      49
                                              438.94
                                                               6742.62
                                    hcd
                                              412.25
## 3
      322257
                      50
                                                               6918.29
                                    hcd
##
     BasePeakMZ ReporterIonIntensityMax Ion_114 Ion_115 Ion_116 Ion_117
## 1
       390.9035
                                 7211.01 2909.30 3358.80 7211.01 3741.59
## 2
       585.8510
                                 5977.44 2457.31 2793.67 5977.44 2964.12
## 3
       412.2520
                                 2242.89 1554.30 2242.89 1474.75 1375.35
##
     Weighted.Avg.Pct.Intensity.Correction
## 1
                                        3.0
```

## Result from P3 container

## 2

## 3

For comparation, 1 file were also loaded and then merged with the identification data using the MSnBase package. A similar filter also applied: (1) unidentified spectrum with msgf e-value < e evalue\_treshold and (2) spectrum with multiple identification were removed

3.2

3.2



```
idSummary(msexp.id)
##
                                                spectrumFile
## 1 TCGA_13-1489_42-2590_36-2529_117C_W_PNNL_B2S5_f01.mzML
                                                      idFile coverage
## 1 TCGA_13-1489_42-2590_36-2529_117C_W_PNNL_B2S5_f01.mzid
                                                                0.977
dim(fData(msexp.id)) # before filter
## [1] 16058
                30
head(fData(msexp.id)[order(fData(msexp.id)[,"scan number(s)"]),])[1:3,cols]
##
                                                 pepseq ms-gf:evalue
                                 accession
## X1.1
                           ref|NP_775799.2 DKGKLLIQRSR
                                                             5.595577
## X2.1 ref|NP 775808.2;ref|NP 001003795.1 LSSIEKIKQLR
                                                            61.405174
           ref|NP_006010.2;ref|NP_006044.1 HRRRLRRRPADR
                                                            17.495434
dim(fData(msexp.filter2)) # After Filter
## [1] 8836
              30
head(fData(msexp.id)[order(fData(msexp.filter2)[,"scan number(s)"]),])[1:3,cols]
##
                                pepseq ms-gf:evalue
                 accession
## X1.1
           ref|NP_775799.2 DKGKLLIQRSR
                                           5.595577
## X2354.1 ref|NP 055625.4
                              RFNMEKRR
                                           9.515514
## X2918.1 ref|NP_065868.1
                                           2.306805
                              NSLSGLRR
Comparing the results
Here is the results from the P3 container. Quantification used "MAX".
head(qnt.id[order(qnt.id$accession),])[1:3,cols.p3]
                                          pepseq ms-gf:evalue iTRAQ4.114
                     accession
```

```
##
## 5445 Contaminant ALBU BOVIN FGERALKAWSVARLSQK
                                                   0.7894632
                                                               27540.73
## 8719 Contaminant_ALBU_BOVIN LCVLHEKTPVSEKVTK
                                                   1.6668929
                                                               94342.88
## 404
              ref|NP 000005.2
                                   GVPIPNKVIFIR
                                                  11.7619490 266568.03
       iTRAQ4.115 iTRAQ4.116 iTRAQ4.117
##
## 5445
         27540.73
                    27540.73
                               27540.73
## 8719
         94342.88
                    94342.88
                               94342.88
## 404
        266568.03 266568.03 266568.03
```



To put in contrast, the data aboved is merged with the data from PNNL. iTRAQ4.114 - iTRAQ4.117 indicates results from P3, while Ion\_114 - Ion\_117 indicate results from PNNL.

```
head(qnt.merged[,c(cols.p3, "Ion_114", "Ion_115", "Ion_116", "Ion_117")])[1:3,]
##
                                pepseq ms-gf:evalue iTRAQ4.114 iTRAQ4.115
           accession
## 1 ref|NP_775799.2
                                            5.595577 13848.4980 13848.4980
                           DKGKLLIQRSR
## 2 ref|NP_002211.1
                              RPWEEGNR
                                            3.909789 1169.3190 1169.3190
## 3 ref|NP_056025.2 VDSITAAGGEGPFPTSR
                                           56.255220
                                                       686.2188
                                                                  686.2188
     iTRAQ4.116 iTRAQ4.117 Ion_114 Ion_115 Ion_116 Ion_117
## 1 13848.4980 13848.4980 2909.30 3358.80 7211.01 3741.59
## 2 1169.3190 1169.3190 711.88 665.36 1191.37 745.25
## 3
       686.2188
                  686.2188
                             52.28
                                     55.44 177.77
                                                      41.19
To see if the results coherence, correlation among each ions is computed:
# Correlation for Ion Reporter 114
cor(qnt.merged$iTRAQ4.114, qnt.merged$Ion_114, method = "spearman", "pairwise.complete.obs")
## [1] 0.6087464
```

```
# Correlation for Ion Reporter 115
cor(qnt.merged$iTRAQ4.115, qnt.merged$Ion_115, method = "spearman", "pairwise.complete.obs")
```

```
## [1] 0.6163082
```

```
# Correlation for Ion Reporter 116
cor(qnt.merged$iTRAQ4.116, qnt.merged$Ion_116, method = "spearman", "pairwise.complete.obs")
```

## [1] 0.6040497

```
# Correlation for Ion Reporter 117
cor(qnt.merged$iTRAQ4.117, qnt.merged$Ion_117, method = "spearman", "pairwise.complete.obs")
```

## [1] 0.6067853

## References

- 1. Domon B, Aebersold R: Mass spectrometry and protein analysis. science 2006, **312**:212–217.
- 2. Deutsch EW: Mass spectrometer output file format mzML. Proteome Bioinformatics 2010:319–331.
- 3. Gatto L, Gibb S: MSnbase: Labelled and label-free mS2 data pre-processing, visualisation and quantification. 2016.