Portable Proteomics Pipeline (P3) Spectrum Count Quantification

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Portable Proteomics Pipeline (P3) is a series of protein mass spectrometry data pre-processing pipelines in Docker containers. The packages includes protein identification, filtering, and quantification for both labeled and label-free mass spectrometry data.

This document discuss the kristiyanto/p3:scquant container, a spectrum count quantification pipeline for label-free mass spectrometry protein quantification. The pipeline is based on MSnID pipeline [1].

Input

The container takes the following files:

```
p3.config # Configuration file
*.mzid # mzIdentML files resulted from p3:msgf container or other identification tools.
```

p3.config may contain various parameters for p3 related containers. To run spectrum count quantification, p3.config must contain the following information:

```
[spectrum_count]
score_treshold = 7.0  # The scoring treshold
error_treshold = 20  # Error Treshold
fdr = 0.01  # False Discovery Rate
iteration = 5000  # Number of iteration
```

Running the Container

to run the container, a docker engine must be installed. A more information about installing Docker engine is available at https://docs.docker.com/engine/installation/. Input files must be mounted to /root/data within the container. This can be done by using -v switch. For MacOS and Windows users, the folder should be located under C:\Users or /Users/. More information about volumens in Docker containers is available at http://container-solutions.com/understanding-volumes-docker/

```
# Download/update the container from DockerHub
docker pull kristiyanto/p3:squant
# Run the container
docker run --rm -v /Users/path/files:/root/data kristiyanto/p3:scquant
```



Output

Once the quantification process is completed, LabelFreeQuant.txt and msnset.rda are generated. LabelFreeQuant.txt is a tab delimited file with the quantification results, with each column represented from each of the mzid file provided. msnset.rda is an msnset object for the result that can be easily imported to R for additional analysis.

Pipeline

kristiyanto/p3:scquant is based on R, and it uses MSnID package and pipeline. A more detailed information about the pipeline is available at http://bioconductor.org/packages/release/bioc/vignettes/MSnID/inst/doc/msnid_vignette.pdf.

For this documentation, three mzid files are processed.

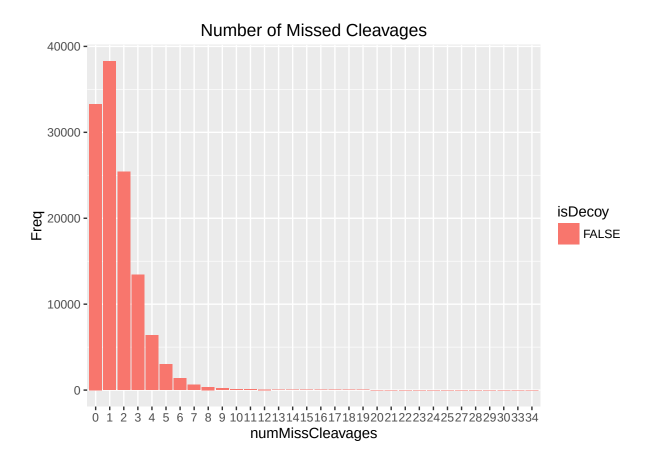
```
# Files
print(mz.files)

## [1] "Biodiversity_mix_01_280ct15_Arwen_15-07-13.mzid"
## [2] "Biodiversity_mix_02_280ct15_Arwen_15-07-13.mzid"
## [3] "Biodiversity_mix_03_280ct15_Arwen_15-07-13.mzid"

prop.table(table(mzid$numIrregCleavages))

##
## 0 1 2
## 0.9668602085 0.0327185017 0.0004212899
```





Filter

Filtering is done by using the parameter defined in the p3.config file. The filtering is including the error calculation (variable: error_treshold), scoring (variable: score_treshold), and filter optimization (variable: iteration and fdr) [3].

```
show(mzid)
```

```
## MSnID object
## Working directory: "."
## #Spectrum Files: 3
## #PSMs: 795177 at 0 % FDR
## #peptides: 122960 at 0 % FDR
## #accessions: 167543 at 0 % FDR
```

show(fObj)

```
## MSnIDFilter object
## (msmsScore > 7) & (massError < 20)</pre>
```



```
evaluate_filter(prj, f0bj, level="PSM")
##
      fdr
## PSM 0 589592
evaluate_filter(prj, f0bj, level="peptide")
##
           fdr
## peptide
           0 49166
evaluate_filter(prj, f0bj, level="accession")
            fdr
## accession 0 102866
show(prj)
## MSnID object
## Working directory: "."
## #Spectrum Files: 3
## #PSMs: 622994 at 0 % FDR
## #peptides: 66562 at 0 % FDR
## #accessions: 121791 at 0 % FDR
```

Quantification

Quantification is done by using the MSnBase package [4].

Combined 46483 features into 39394 using sum

Results

```
exprs.table <- exprs(msnset)
exprs.table <- cbind(Protein=row.names(exprs.table), as.data.frame(exprs.table))
head(exprs.table)</pre>
```



```
##
                                      Protein
## Contaminant_Trypa5
                           Contaminant_Trypa5
## sp|AOA321|ATPF_COFAR sp|AOA321|ATPF_COFAR
## sp|AOA348|CYF_COFAR
                          sp|AOA348|CYF_COFAR
## sp|AOA393|YCF1_COFAR sp|AOA393|YCF1_COFAR
## sp|AOA4Z3|A3LT2_RAT
                         sp|AOA4Z3|A3LT2_RAT
## sp|AOAAS4|SMS1 PIG
                           sp|AOAAS4|SMS1_PIG
##
                        Biodiversity_mix_01_280ct15_Arwen_15-07-13.mzXML
## Contaminant_Trypa5
                                                                         0
## sp|AOA321|ATPF_COFAR
## sp|AOA348|CYF_COFAR
                                                                         0
## sp|AOA393|YCF1_COFAR
                                                                         1
## sp|AOA4Z3|A3LT2_RAT
                                                                         0
## sp|AOAAS4|SMS1_PIG
##
                        Biodiversity_mix_02_280ct15_Arwen_15-07-13.mzXML
## Contaminant_Trypa5
## sp|AOA321|ATPF_COFAR
                                                                         0
## sp|AOA348|CYF_COFAR
                                                                         1
## sp|AOA393|YCF1_COFAR
                                                                         3
## sp|AOA4Z3|A3LT2 RAT
                                                                         0
## sp|AOAAS4|SMS1_PIG
##
                        Biodiversity_mix_03_280ct15_Arwen_15-07-13.mzXML
## Contaminant_Trypa5
## sp|AOA321|ATPF_COFAR
                                                                         1
## sp|AOA348|CYF_COFAR
                                                                         1
## sp|AOA393|YCF1_COFAR
                                                                         2
## sp|AOA4Z3|A3LT2_RAT
                                                                         1
## sp|AOAAS4|SMS1_PIG
```

References

- 1. Laurent Gatto VP with contributions from: MSnID: Utilities for Exploration and Assessment of Confidence of LC-MSn Proteomics Identifications.
- 2. Petyuk VA: MSnID package for handling mS/MS identifications. 2015.
- 3. Deutsch EW: Mass spectrometer output file format mzML. Proteome Bioinformatics 2010:319–331.
- 4. Gatto L, Lilley K: MSnbase an r/Bioconductor package for isobaric tagged mass spectrometry data visualization, processing and quantitation. *Bioinformatics* 2012, **28**:288–289.