

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.

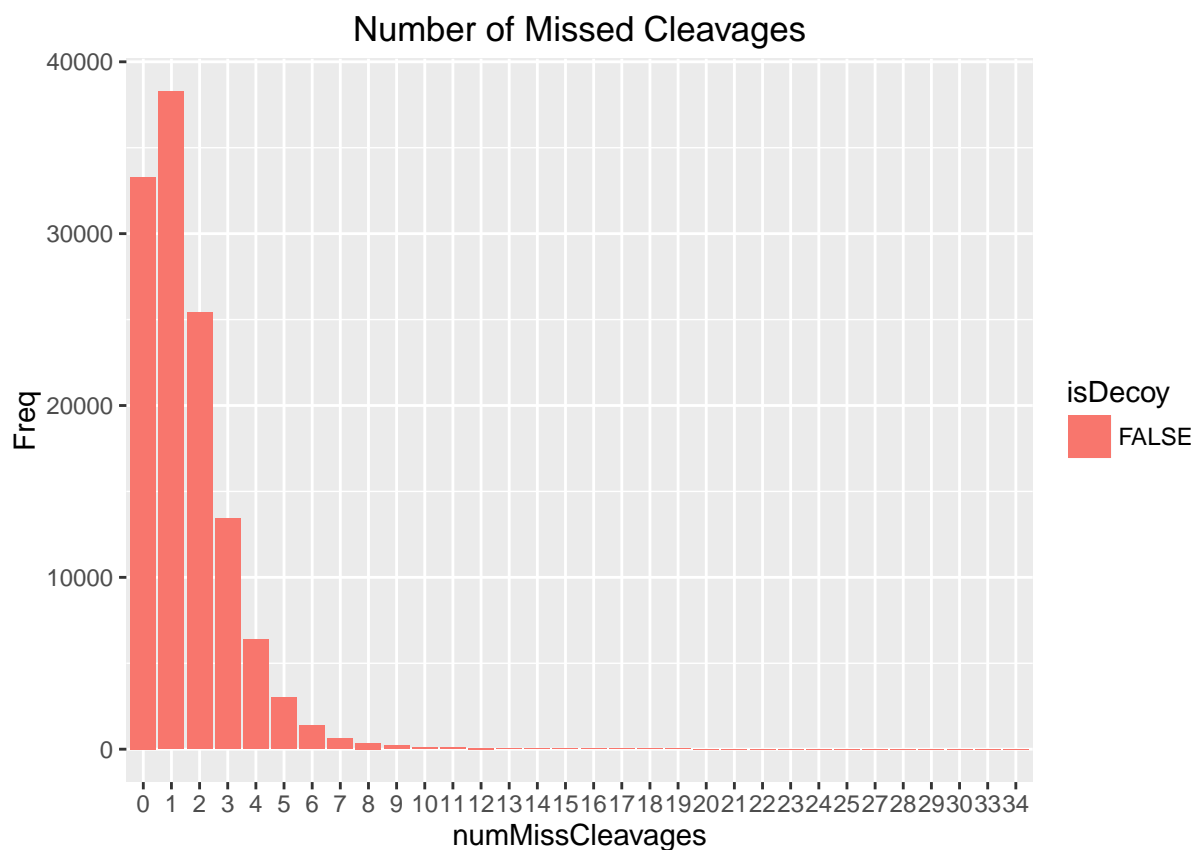
```
# Files
```

```
print(mz.files)
```

```
## [1] "Biodiversity_mix_01_28Oct15_Arwen_15-07-13.mzid"
## [2] "Biodiversity_mix_02_28Oct15_Arwen_15-07-13.mzid"
## [3] "Biodiversity_mix_03_28Oct15_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)
```

```
evaluate_filter(prj, fObj, level="PSM")
```

```
##      fdr      n
## PSM    0 589592
```

```
evaluate_filter(prj, fObj, level="peptide")
```

```
##      fdr      n
## peptide  0 49166
```

```
evaluate_filter(prj, fObj, level="accession")
```

```
##      fdr      n
## 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].

```
msnset <- as(prj, "MSnSet")
msnset <- combineFeatures(msnset, fData(msnset)$accession, redundancy.handler="unique",
                          fun="sum", cv=FALSE)
```

```
## 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)
```

```

##                                     Protein
## Contaminant_Trypa5      Contaminant_Trypa5
## sp|AOA321|ATPF_CO FAR  sp|AOA321|ATPF_CO FAR
## sp|AOA348|CYF_CO FAR   sp|AOA348|CYF_CO FAR
## sp|AOA393|YCF1_CO FAR  sp|AOA393|YCF1_CO FAR
## sp|AOA4Z3|A3LT2_RAT    sp|AOA4Z3|A3LT2_RAT
## sp|AOAAS4|SMS1_PIG     sp|AOAAS4|SMS1_PIG
##                                     Biodiversity_mix_01_28Oct15_Arwen_15-07-13.mzXML
## Contaminant_Trypa5                                           2
## sp|AOA321|ATPF_CO FAR                                         0
## sp|AOA348|CYF_CO FAR                                           0
## sp|AOA393|YCF1_CO FAR                                           1
## sp|AOA4Z3|A3LT2_RAT                                           0
## sp|AOAAS4|SMS1_PIG                                           1
##                                     Biodiversity_mix_02_28Oct15_Arwen_15-07-13.mzXML
## Contaminant_Trypa5                                           1
## sp|AOA321|ATPF_CO FAR                                         0
## sp|AOA348|CYF_CO FAR                                           1
## sp|AOA393|YCF1_CO FAR                                           3
## sp|AOA4Z3|A3LT2_RAT                                           0
## sp|AOAAS4|SMS1_PIG                                           0
##                                     Biodiversity_mix_03_28Oct15_Arwen_15-07-13.mzXML
## Contaminant_Trypa5                                           1
## sp|AOA321|ATPF_CO FAR                                           1
## sp|AOA348|CYF_CO FAR                                           1
## sp|AOA393|YCF1_CO FAR                                           2
## sp|AOA4Z3|A3LT2_RAT                                           1
## sp|AOAAS4|SMS1_PIG                                           0

```

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.