SWATH2stats example script

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Example R code showing the usage of the SWATH2stats package. The data processed is the publicly available dataset of *S.pyogenes* (Röst et al. 2014) (http://www.peptideatlas.org/PASS/PASS00289). The results file 'rawOpenSwathResults_1pcnt_only.tsv' can be found on PeptideAtlas (ftp://PASS00289@ftp.peptideatlas. org/../Spyogenes/results/). This is a R Markdown file, showing the result of processing this data. The lines shaded in grey represent the R code executed during this analysis.

The stable release package SWATH2stats can be directly installed from Bioconductor using the commands below. This file here was generated using the current development release SWATH2stats v.1.1.14 that can be downloaded from http://bioconductor.org/packages/devel/bioc/html/SWATH2stats.html.

```
## try http:// if https:// URLs are not supported
source('https://bioconductor.org/biocLite.R')
biocLite('SWATH2stats')

## Conversely, install from github
devtools::install_github("abelew/SWATH2stats")
```

Part 1: Loading and annotation

Load the SWATH-MS example data from the package, this is a reduced file in order to limit the file size of the package.

```
library(SWATH2stats)
library(data.table)
data('Spyogenes', package='SWATH2stats')
```

Alternatively the original file downloaded from the Peptide Atlas can be loaded from the working directory.

```
data <- data.frame(fread('rawOpenSwathResults_1pcnt_only.tsv', sep='\t', header=TRUE))</pre>
```

Extract the study design information from the file names. Alternatively, the study design table can be provided as an external table.

```
## Filename Condition BioReplicate Run

## 1 Strep0_Repl1_R02/split_hroest_K120808 Strep0 1 1

## 2 Strep0_Repl2_R02/split_hroest_K120808 Strep0 2 2

## 3 Strep10_Repl1_R02/split_hroest_K120808 Strep10 1 3

## 4 Strep10_Repl2_R02/split_hroest_K120808 Strep10 2 4
```

The SWATH-MS data is annotated using the study design table.

Not checking that the files are identical between the annotation and data.

Remove the decoy peptides for a subsequent inspection of the data.

data.annotated.nodecoy <- subset(data.annotated, decoy==FALSE)</pre>

Part 2: Analyze correlation, variation and signal

Count the different analytes for the different injections.

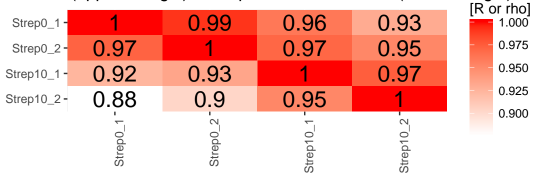
count_analytes(data.annotated.nodecoy)

##		run_id	transition_group_id	fullpeptidename	proteinname
##	1	Strep0_1_1	10229	8377	1031
##	2	Strep0_2_2	9716	7970	1003
##	3	Strep10_1_3	8692	7138	943
##	4	Strep10 2 4	8424	6941	910

Plot the correlation of the signal intensity.

intensity correlation between samples:

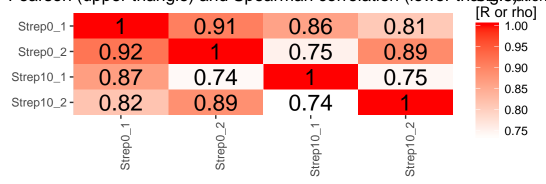
Pearson (upper triangle) and Spearman correlation (lower triangle) ation



Plot the correlation of the delta_rt, which is the deviation of the retention time from the expected retention time.

delta_rt correlation between samples:

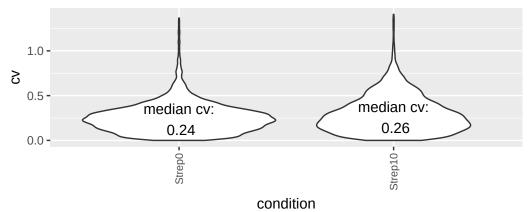
Pearson (upper triangle) and Spearman correlation (lower triangle) ation



Plot the variation of the signal across replicates.

variation <- plot_variation(data.annotated.nodecoy)</pre>

intensity cv across conditions



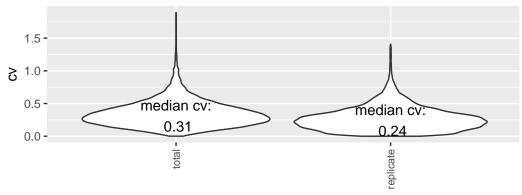
head(variation)

```
##
                                  transition_group_id condition
                         1_FSWISTGGGASMELLEGK/2_run0
## 1
                                                         Strep0 135206 119997
## 2
                         1_FSWISTGGGASMELLEGK/2_run0
                                                        Strep10 147766 110436
                   1000_TGIFSQDDENALENSIGFSSK/3_run0
## 3
                                                         Strep0
                                                                   6946
                                                                          4161
                              10000_DIVEAVIPR/2_run0
## 4
                                                         Strep0 163405
                                                                         67537
                              10000_DIVEAVIPR/2_run0
## 5
                                                        Strep10
                                                                 53345
                                                                         20963
## 6 10001_SSGYNLGGEQSGHVIIMDYNTTGDGQLTAIQLAK/3_run0
                                                         Strep0
                                                                 10798 10876
##
## 1 0.084281039
## 2 0.204462368
## 3 0.354603833
## 4 0.587064396
## 5 0.616287124
## 6 0.005089446
```

Plot the total variation versus variation within replicates.

variation_total <- plot_variation_vs_total(data.annotated.nodecoy)</pre>

intensity coefficient of variation - total versus within replicates



variation_total[[2]]

scope mode_cv mean_cv median_cv

```
## 1 replicate 0.2209867 0.2728681 0.2438041
## 2 total 0.2655678 0.3439050 0.3139993
```

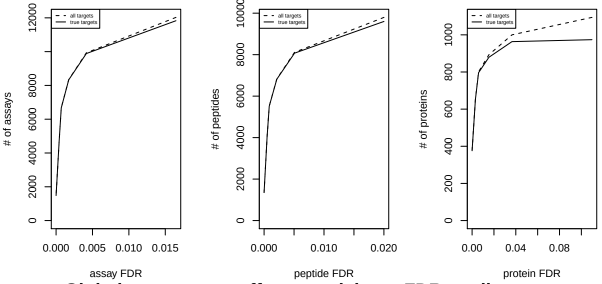
Calculate the summed signal per peptide and protein across samples.

```
peptide_signal <- write_matrix_peptides(data.annotated.nodecoy)
protein_signal <- write_matrix_proteins(data.annotated.nodecoy)
head(protein_signal)</pre>
```

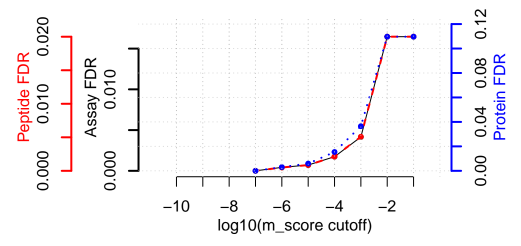
```
##
                         proteinname Strep0_1_1 Strep0_2_2 Strep10_1_3
## 1
     Spyo_Exp3652_DDB_SeqID_1571119
                                         265206
                                                    163326
                                                                  51831
## 2
     Spyo_Exp3652_DDB_SeqID_1579753
                                         185725
                                                    150672
                                                                  21483
## 3 Spyo_Exp3652_DDB_SeqID_1631459
                                         176686
                                                    132415
                                                                  42165
## 4 Spyo_Exp3652_DDB_SeqID_1640263
                                           3310
                                                      6617
                                                                  98550
## 5 Spyo_Exp3652_DDB_SeqID_1709452
                                                    747772
                                                                 503581
                                         852502
## 6 Spyo_Exp3652_DDB_SeqID_17244480
                                          17506
                                                      29578
                                                                   7607
     Strep10_2_4
##
## 1
           45021
## 2
          144314
## 3
           32735
## 4
           45169
## 5
          504761
## 6
            2482
```

Part 3: FDR estimation

Estimate the overall FDR across runs using a target decoy strategy.



Global m-score cutoff connectivity to FDR quality



According to this FDR estimation one would need to filter the data with a lower mscore threshold to reach an overall protein FDR of 5%.

```
mscore4protfdr(data, FFT=0.25, fdr_target=0.05)
## Target protein FDR: 0.05
## Required overall m-score cutoff: NA
```

achieving protein FDR: NA

[1] NA

Part 4: Filtering

Number of proteins: 830

Number of peptides: 4642

##

##

Filter data for values that pass the 0.001 mscore criteria in at least two replicates of one condition.

data.filtered <- filter_mscore_condition(data.annotated, 0.001, n.replica = 2)

Fraction of peptides selected: 0.67

Original dimension: 37061, new dimension: 29835, difference: 7226.

Select only the 10 peptides showing strongest signal per protein.

data.filtered2 <- filter_on_max_peptides(data.filtered, n_peptides = 10)

Before filtering:

Number of proteins: 884

Number of peptides: 6594

##

Percentage of peptides removed: 29.6%

##

After filtering:

```
Filter for proteins that are supported by at least two peptides.
```

```
data.filtered3 <- filter_on_min_peptides(data.filtered2, n_peptides = 2)

## Before filtering:
## Number of proteins: 830

## Number of peptides: 4642

##

## Percentage of peptides removed: 0.3%

##

## After filtering:
## Number of proteins: 716

## Number of peptides: 4628</pre>
```

Part 5: Conversion

5 28903_EKAEAAIYQFLEAIGENPNR/3_y6

6 28903_EKAEAAIYQFLEAIGENPNR/3_y6

bioreplicate condition run

1

1

2

Strep0

Strep0

3

2

Strep10

1

2

3

```
Convert the data into a transition-level format (one row per transition measured).
```

```
data.transition <- disaggregate(data.filtered3)</pre>
## The library contains between 4 and 6 transitions per precursor.
## The data table was transformed into a table containing one row per transition.
## 4 row(s) was(were) removed because they did not contain data due to different number of transitions
Convert the data into the format required by MSstats.
MSstats.input <- convert_MSstats(data.transition)
## One or several columns required by MSstats were not in the data. The columns were created and filled
## Missing columns: productcharge, isotopelabeltype
## isotopelabeltype was filled with light.
## Warning in convert_MSstats(data.transition): Intensity values which were 0
## have been replaced by NA.
head(MSstats.input)
##
                        proteinname
                                       peptidesequence precursorcharge
## 1 Spyo_Exp3652_DDB_SeqID_1571119
                                          SLPEEDLDKNEK
## 2 Spyo_Exp3652_DDB_SeqID_1571119
                                          SLPEEDLDKNEK
                                                                      2
## 3 Spyo Exp3652 DDB SeqID 1571119
                                                                      2
                                         TIFDDEPISEETK
## 4 Spyo_Exp3652_DDB_SeqID_1571119
                                                                      2
                                        TIFDDEPISEETK
## 5 Spyo_Exp3652_DDB_SeqID_1571119 LSLPSQEPLLAAFHGEK
## 6 Spyo_Exp3652_DDB_SeqID_1571119 LSLPSQEPLLAAFHGEK
##
                         fragmention productcharge isotopelabeltype intensity
## 1
             118149_AHIAYLPSDGR/2_y8
                                                               light
                                                                           4036
                                                 NA
## 2
             118149_AHIAYLPSDGR/2_y8
                                                               light
                                                                           1642
                                                 NA
## 3
             118149_AHIAYLPSDGR/2_y8
                                                               light
                                                                           2405
                                                 NA
             118149_AHIAYLPSDGR/2_y8
                                                 NA
                                                                light
                                                                            720
```

NA

NA

light

light

3410

1984

```
## 4 2 Strep10 4
## 5 1 Strep0 1
## 6 2 Strep10 4
```

Convert the data into the format required by mapDIA.

```
mapDIA.input <- convert_mapDIA(data.transition)
head(mapDIA.input)</pre>
```

```
proteinname
                                                 peptidesequence
## 1 Spyo_Exp3652_DDB_SeqID_1571119
                                                    SLPEEDLDKNEK
## 2 Spyo_Exp3652_DDB_SeqID_1571119
                                                    TIFDDEPISEETK
## 3 Spyo_Exp3652_DDB_SeqID_1571119
                                               LSLPSQEPLLAAFHGEK
## 4 Spyo_Exp3652_DDB_SeqID_1571119
                                                      SLETEGKVDK
## 5 Spyo_Exp3652_DDB_SeqID_1579753 TLIDAYEAFC[160]PLDLSMEGDVK
## 6 Spyo_Exp3652_DDB_SeqID_1579753
                                             SDTAGTIVSLNTDLPNQSK
##
                          fragmention Strep0_1 Strep0_2 Strep10_1 Strep10_2
## 1
             118149_AHIAYLPSDGR/2_y8
                                           4036
                                                               1642
                                                     NaN
                                                                           NaN
## 2
             118149_AHIAYLPSDGR/2_y8
                                            NaN
                                                    2405
                                                                NaN
                                                                           720
                                                                          1984
## 3 28903_EKAEAAIYQFLEAIGENPNR/3_y6
                                           3410
                                                     NaN
                                                                NaN
## 4 28903_EKAEAAIYQFLEAIGENPNR/3_y6
                                            NaN
                                                    2185
                                                                NaN
                                                                          NaN
      97491_LALAPNTPGQIVALELGEK/3_y7
                                                    4099
                                                               3060
                                                                          2301
## 5
                                           5681
## 6
         56597_LNDGAFLALDGSAQYK/2_y9
                                                                           860
                                           3349
                                                    2552
                                                                \mathtt{NaN}
```

Convert the data into the format required by aLFQ.

```
aLFQ.input <- convert_aLFQ(data.transition)
head(aLFQ.input)</pre>
```

```
##
                                      protein_id
          run id
                                                        peptide_id
## 1 Strep0_1_1 Spyo_Exp3652_DDB_SeqID_1571119
                                                      SLPEEDLDKNEK
## 2 Strep10_1_3 Spyo_Exp3652_DDB_SeqID_1571119
                                                      SLPEEDLDKNEK
## 3 Strep0_2_2 Spyo_Exp3652_DDB_SeqID_1571119
                                                     TIFDDEPISEETK
## 4 Strep10_2_4 Spyo_Exp3652_DDB_SeqID_1571119
                                                     TIFDDEPISEETK
## 5 StrepO_1_1 Spyo_Exp3652_DDB_SeqID_1571119 LSLPSQEPLLAAFHGEK
## 6 Strep10_2_4 Spyo_Exp3652_DDB_SeqID_1571119 LSLPSQEPLLAAFHGEK
                                             transition_id
##
## 1
                      AHIAYLPSDGR 118149_AHIAYLPSDGR/2_y8
## 2
                      AHIAYLPSDGR 118149_AHIAYLPSDGR/2_y8
## 3
                      AHIAYLPSDGR 118149_AHIAYLPSDGR/2_y8
## 4
                      AHIAYLPSDGR 118149_AHIAYLPSDGR/2_y8
## 5 EKAEAAIYQFLEAIGENPNR 28903_EKAEAAIYQFLEAIGENPNR/3_y6
## 6 EKAEAAIYQFLEAIGENPNR 28903 EKAEAAIYQFLEAIGENPNR/3 v6
         peptide_sequence precursor_charge transition_intensity concentration
##
## 1
              AHIAYLPSDGR
                                          2
                                                            4036
                                                                              ?
## 2
              AHIAYLPSDGR
                                          2
                                                                              ?
                                                            1642
                                                                              ?
## 3
              AHIAYLPSDGR
                                          2
                                                            2405
                                          2
                                                                              ?
## 4
              AHIAYLPSDGR
                                                             720
                                          3
                                                                              ?
## 5 EKAEAAIYQFLEAIGENPNR
                                                            3410
                                                                              ?
## 6 EKAEAAIYQFLEAIGENPNR
                                                            1984
```

Session info on the R version and packages used.

sessionInfo()

```
## R version 3.5.0 (2018-04-23)
## Platform: x86_64-pc-linux-gnu (64-bit)
## Running under: Debian GNU/Linux buster/sid
```

```
##
## Matrix products: default
## BLAS: /usr/lib/x86_64-linux-gnu/openblas/libblas.so.3
## LAPACK: /usr/lib/x86_64-linux-gnu/libopenblasp-r0.2.20.so
## locale:
## [1] LC CTYPE=en US.utf8
                                  LC NUMERIC=C
## [3] LC_TIME=en_US.utf8
                                  LC_COLLATE=en_US.utf8
   [5] LC_MONETARY=en_US.utf8
                                  LC_MESSAGES=en_US.utf8
  [7] LC_PAPER=en_US.utf8
                                  LC_NAME=C
  [9] LC_ADDRESS=C
                                  LC_TELEPHONE=C
## [11] LC_MEASUREMENT=en_US.utf8 LC_IDENTIFICATION=C
## attached base packages:
## [1] stats
                graphics grDevices utils
                                               datasets methods
                                                                   base
##
## other attached packages:
## [1] data.table_1.11.4 SWATH2stats_1.11.3
## loaded via a namespace (and not attached):
## [1] Rcpp_0.12.17
                             knitr_1.20
                                                  magrittr_1.5
## [4] devtools_1.13.5
                             munsell_0.5.0
                                                  colorspace_1.3-2
## [7] rlang_0.2.1
                             stringr_1.3.1
                                                  plyr_1.8.4
## [10] tools_3.5.0
                             grid_3.5.0
                                                  gtable_0.2.0
## [13] withr_2.1.2
                             htmltools_0.3.6
                                                  yaml_2.1.19
## [16] lazyeval_0.2.1
                             rprojroot_1.3-2
                                                  digest_0.6.15
## [19] tibble_1.4.2
                             reshape2_1.4.3
                                                  formatR_1.5
## [22] ggplot2_2.2.1
                                                  evaluate_0.10.1
                             memoise_1.1.0
## [25] rmarkdown_1.10
                             labeling_0.3
                                                  stringi_1.2.3
## [28] pillar_1.2.3
                             compiler_3.5.0
                                                  BiocInstaller_1.30.0
## [31] scales_0.5.0
                             backports_1.1.2
```