US HUPO 2017 Short course - Section 7 : Protein significance analysis with MSstats

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This section describes steps and considerations 1) to properly format data processed by Skyline, prior to the MSstats analysis. 2) to properly use MSstats for significance analysis.

In the following example, the quantified peak intensities data from skyline file for **iPRG_10ppm_2rt_15cut_nosingle** of ABRF 2015 is used.

0. Prerequisites

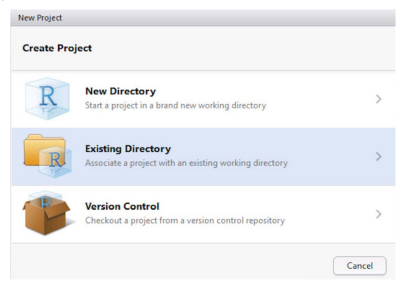
Export the report from Skyline.

File > Export > Report > select MSstat Input > save .csv file.

todo: add capture from Skyline add which skyline file, locate the directory in USB

1. Create a new Rstudio project

From the menu, select **File** > **New Project...**, then select **Existing Directory** and choose the directory where you downloaded 1) this script, 2) MSstats input report from Skyline, and 3) the example datasets for this tutorial. All the output files we'll be creating in this tutorial will be saved in the 'working directory' that now has been set by Rstudio.



Let's verify the working directory path with the get working directory command.

```
getwd()
```

[1] "/Users/meenachoi/Dropbox/visits/2017/03/US Hupo/Section7_MSstats"

2. Read Skyline output

The required input data is generated automatically when using MSstats report format in Skyline. We first load and access the dataset processed by Skyline. The name of saved file from Skyline using MSstats report format is 'iPRG_10ppm_2rt_15cut_nosingle.csv'. or you can use the published data from todo: add link, This file is available in panorama.

```
# Read output from skyline : Cox.skyline.csv
raw <- read.csv(file="iPRG_10ppm_2rt_15cut_nosingle.csv")</pre>
```

We can read csv file. Here we will load R data file which is the exactly same data in iPRG_10ppm_2rt_15cut_nosingle.csv file.

```
# Load R data, which is convered from csv file, output from skyline : Cox.skyline.csv
load("iprg.skyline.RData")
raw <- iprg.skyline
head(raw)</pre>
```

```
##
                      ProteinName PeptideSequence PeptideModifiedSequence
## 1 DECOY_sp|POCF18|YMO85_YEAST
                                        KDMYGNPFQK
                                                            KDM[+16]YGNPFQK
## 2 DECOY_sp|POCF18|YM085_YEAST
                                        KDMYGNPFQK
                                                            KDM[+16]YGNPFQK
## 3 DECOY_sp|POCF18|YM085_YEAST
                                        KDMYGNPFQK
                                                            KDM[+16]YGNPFQK
## 4 DECOY sp|POCF18|YM085 YEAST
                                        KDMYGNPFQK
                                                            KDM[+16]YGNPFQK
## 5 DECOY_sp|POCF18|YM085_YEAST
                                        KDMYGNPFQK
                                                            KDM[+16]YGNPFQK
                                                            KDM[+16]YGNPFQK
## 6 DECOY sp|POCF18|YM085 YEAST
                                        KDMYGNPFQK
##
     PrecursorCharge PrecursorMz FragmentIon ProductCharge ProductMz
## 1
                    3
                         415.1974
                                    precursor
                                                            3 415.1974
## 2
                    3
                                                              415.1974
                         415.1974
                                                            3
                                    precursor
                    3
## 3
                         415.1974
                                    precursor
                                                            3
                                                               415.1974
                    3
                                                            3
## 4
                         415.1974
                                     precursor
                                                               415.1974
## 5
                    3
                         415.1974
                                    precursor
                                                            3
                                                               415.1974
                    3
## 6
                         415.1974
                                    precursor
                                                               415.1974
##
     IsotopeLabelType Condition BioReplicate
                                                                 FileName
## 1
                light
                              NA
                                            NA JD_06232014_sample1_B.raw
## 2
                light
                                            NA JD_06232014_sample1_C.raw
                              NA
## 3
                light
                              NA
                                            NA JD_06232014_sample1-A.raw
## 4
                                            NA JD_06232014_sample2_A.raw
                light
                              NA
## 5
                light
                              NA
                                            NA JD_06232014_sample2_B.raw
## 6
                                            NA JD_06232014_sample2_C.raw
                light
                              NA
##
        Area StandardType Truncated annotation QValue
## 1
     147327
                        NA
                               False
## 2 1373397
                        NA
                               False
                                                     NA
## 3
       71765
                        NA
                               False
                                                     NA
## 4
       66387
                        NA
                               False
                                                     NA
## 5
                        NA
                               False
                                                     NA
      107736
## 6
      380812
                        NA
                               False
                                                     NA
```

3. Load MSstats

Load MSstats first. Then you are ready to start MSstats.

```
library(MSstats)
?MSstats
```

4. Preprocessing for skyline output

4.1. Set annotation file

Annotation information is required to fill in Condition and BioReplicate for corresponding Run information. Users have to prepare as csv or txt file like 'iPRG_skyline_annotation.csv', which includes Run, Condition, and BioReplicate information, and load it in R.

```
annot <- read.csv("iPRG_skyline_annotation.csv", header=TRUE)
annot</pre>
```

```
##
                             Run Condition BioReplicate
## 1
      JD_06232014_sample1-A.raw Condition1
                                                       1
      JD_06232014_sample2_A.raw Condition2
                                                       2
      JD_06232014_sample4_B.raw Condition4
                                                       4
      JD_06232014_sample1_B.raw Condition1
                                                       1
## 4
## 5
      JD_06232014_sample1_C.raw Condition1
                                                       1
      JD 06232014 sample2 B.raw Condition2
                                                       2
      JD_06232014_sample2_C.raw Condition2
                                                       2
## 7
      JD_06232014_sample3_A.raw Condition3
                                                       3
                                                       3
      JD_06232014_sample3_B.raw Condition3
## 10 JD_06232014_sample3_C.raw Condition3
                                                       3
                                                       4
## 11 JD_06232014_sample4-A.raw Condition4
## 12 JD_06232014_sample4_C.raw Condition4
```

4.2. Preprocessing with SkylinetoMSstatsFormat

The input data for MSstats is required to contain variables of ProteinName, PeptideSequence, PrecursorCharge, FragmentIon, ProductCharge, IsotopeLabelType, Condition, BioReplicate, Run, Intensity. These variable names should be fixed. MSstats input from Skyline adapts the column scheme of the dataset so that it fits MSstats input format. However there are several extra column names and also some of them need to be changed. SkylinetoMSstatsFormat function helps pre-processing for making right format of MSstats input from Skyline output. For example, it renames some column name, and replace truncated peak intensities with NA. Another important step is to handle isotopic peaks before using dataProcess. The output from Skyline for DDA experiment has several measurements of peak area from the monoisotopic, M+1 and M+2 peaks. To get a robust measure of peptide intensity, we can sum over isotopic peaks per peptide or use the highest peak. Here we take a summation per peptide ion.

Here is the summary of pre-processing steps in SkylinetoMSstatsFormat function (in orange box below).

In Skyline

Remove duplicated rows: exactly same values in some rows

Remove decoy proteins

Remove protein which has only one peptide per protein

In MSstats

SkylinetoMSstatsFormat

- Rename column names
- Replace NA for truncated rows
- Sum of isotopic peaks per peptide and charge
- Replace intensity = 1 with zero if intensity < 1
- Log 2 transform for intensity
- Normalization
- Extra step for imputation : Distinguish missing at random and censored missing
 - Decide cutoff for censored missing values among all log2(intensity) > 0.
 - If log2(intensity) < cutoff, log2(intensity) replaces with zero and is considered as censored missing values (censoredInt='0'), then, will be imputed.
 - NA will be remained as NA, which are missing at random.

```
# reformating and pre-processing for Skyline output.
quant <- SkylinetoMSstatsFormat(raw, annotation=annot)</pre>
```

```
## ** Proteins, which names include DECOY, are removed.
```

Peptides, that are used in more than one proteins, are removed.

Warning in SkylinetoMSstatsFormat(raw, annotation = annot): NAs introduced
by coercion

** Truncated peaks are replaced with NA.

** For DDA datasets, three isotopic peaks per feature and run are summed.

head(quant)

##		ProteinName	Pe	eptideSequence	PrecursorCharge	FragmentIon
##	1	sp P38915 SPT8_YEAST	AAAA	GAGGAGDSGDAVTK	2	sum
##	2	sp P40457 MLP2_YEAST		QLLDESSEQK	2	sum
##	3	sp P42943 TCPH_YEAST		GGAEQVIAEVER	2	sum
##	4	sp P40202 CCS1_YEAST		DYSFLGVIAR	2	sum
##	5	sp Q12049 THP3_YEAST	LSWNNVNQVS	SNPLMVTPLPGLQK	3	sum
##	6	sp P47122 NPA3_YEAST	7	TPPYVINLDPAVLR	3	sum
##		ProductCharge Isotope	eLabelType	Condition Bio	oReplicate	
##	1	NA	L	Condition1	1	
##	2	NA	L	Condition1	1	
##	3	NA	L	Condition1	1	
##	4	NA	L	Condition1	1	
##	5	NA	L	Condition1	1	
##	6	NA	L	Condition1	1	
##			Run Inter	nsity		
##	1	JD_06232014_sample1_H	3.raw	0		
##	2	JD_06232014_sample1_H	3.raw 668	36797		

```
## 3 JD_06232014_sample1_B.raw 284344776
## 4 JD_06232014_sample1_B.raw 200172818
## 5 JD_06232014_sample1_B.raw 2841395
## 6 JD_06232014_sample1_B.raw 5602300
```

For further details, visit the help file using the following code.

?SkylinetoMSstatsFormat

5. Processing the data for analysis: Normalizing and summarizing data with dataProcess

! Always pay attention to the default options

After reading the datasets, MSstats performs 1) logarithm transformation of Intensity column, 2) normalization, 3) feature selection, (all features vs subset of features), 4) imputation for censored missing value, which are below the cutoff and undetectable, 5) run-level summarization.

To get started with this function, visit the help section of dataProcess first:

Default normalization and summarization options

Then (1) normalization will be performed first. The default option for normalization is equalizeMedians. 'equalizeMedians' fits for label-based SRM experiments, which we can use reference signals. There are three more options for normalization. Depending on the suitable assumption for your experiment, you can choose one of them.

Then, (2) run level summarization will be performed including missing value imputation by accerelated failure model and robust parameter estimation by TMP (Tukey's median polish).

Below show the default for all options in dataProcess except skylineReport and censoredInt. For input from Skyline, skylineReport=TRUE and censoredInt='NA'

Let's check output from dataProcess

```
##
                      PROTEIN
                                                     PEPTIDE TRANSITION
## 27092 sp|D6VTK4|STE2 YEAST EGEVEPVDM[+16]YTPDTAADEEARK 3
                                                                 sum NA
## 10700 sp|D6VTK4|STE2 YEAST
                                   EGEVEPVDMYTPDTAADEEARK 3
## 20407 sp|D6VTK4|STE2_YEAST
                                      FYPGTLSSFQTDSINNDAK_2
                                                                 sum NA
## 2213 sp|D6VTK4|STE2 YEAST
                                                IGPFADASYK 2
                                                                 sum NA
         sp|D6VTK4|STE2 YEAST
                                                   KETTSDK 2
                                                                 sum NA
## 26116 sp|D6VTK4|STE2 YEAST
                                             NOFYOLPTPTSSK 2
                                                                 sum NA
                                      FEATURE LABEL GROUP ORIGINAL
## 27092 EGEVEPVDM[+16]YTPDTAADEEARK_3_sum_NA
                                                         Condition1
## 10700
              EGEVEPVDMYTPDTAADEEARK_3_sum_NA
                                                         Condition1
## 20407
                 FYPGTLSSFQTDSINNDAK_2_sum_NA
                                                         Condition1
## 2213
                          IGPFADASYK_2_sum_NA
                                                         Condition1
                                                   L
## 938
                             KETTSDK_2_sum_NA
                                                   L
                                                         Condition1
## 26116
                                                  L
                       NQFYQLPTPTSSK_2_sum_NA
                                                         Condition1
         SUBJECT_ORIGINAL RUN GROUP SUBJECT SUBJECT_NESTED INTENSITY
## 27092
                            1
                                  1
                                          1
                                                              5222795
                        1
## 10700
                        1
                            1
                                  1
                                          1
                                                        1.1 182195648
## 20407
                                                        1.1 86229170
## 2213
                                                        1.1 157996653
                        1
                                  1
                                          1
## 938
                        1
                                  1
                                          1
                                                        1.1 177684007
## 26116
                        1
                                          1
                                                        1.1 140368798
        ABUNDANCE METHOD
                                        originalRUN censored
## 27092 22.07353
                        1 JD_06232014_sample1_B.raw
                                                        FALSE
                        1 JD 06232014 sample1 B.raw
## 10700 27.19805
                                                        FALSE
## 20407 26.11881
                        1 JD 06232014 sample1 B.raw
                                                        FALSE
                        1 JD_06232014_sample1_B.raw
## 2213
          26.99246
                                                        FALSE
## 938
          27.16188
                        1 JD_06232014_sample1_B.raw
                                                        FALSE
## 26116 26.82179
                        1 JD_06232014_sample1_B.raw
                                                        FALSE
# This table includes run-level summarized log2 intensities. (column : LogIntensities)
# Now one summarized log2 intensities per Protein and Run.
# NumMeasuredFeature : show how many features are used for run-level summarization.
          If there is no missing value, it should be the number of features in certain protein.
# MissingPercentage : the number of missing features / the number of features in certain protein.
head(quant.processed$RunlevelData)
##
                      Protein LogIntensities NumMeasuredFeature
       1 sp|D6VTK4|STE2_YEAST
## 1
                                    26.81232
       2 sp|D6VTK4|STE2 YEAST
                                    26.60786
                                                               8
                                                               8
       3 sp|D6VTK4|STE2 YEAST
                                    26.58301
                                    26.83563
                                                               8
## 4
       4 sp|D6VTK4|STE2_YEAST
## 5
       5 sp|D6VTK4|STE2_YEAST
                                    26.79430
       6 sp|D6VTK4|STE2_YEAST
                                    26.60863
     MissingPercentage more50missing NumImputedFeature
## 1
                     0
                               FALSE
## 2
                     0
                               FALSE
                                                      0
## 3
                     0
                               FALSE
                                                      0
## 4
                     0
                               FALSE
                                                      0
## 5
                     0
                               FALSE
                                                      0
## 6
                     0
                               FALSE
                                                      0
                   originalRUN GROUP GROUP ORIGINAL SUBJECT ORIGINAL
## 1 JD_06232014_sample1_B.raw
                                   1
                                         Condition1
                                                                    1
## 2 JD_06232014_sample1_C.raw
                                   1
                                         Condition1
                                                                    1
## 3 JD_06232014_sample1-A.raw
                                   1
                                         Condition1
                                                                    1
```

Condition2

2

4 JD_06232014_sample2_A.raw

```
## 5 JD_06232014_sample2_B.raw
                                           Condition2
## 6 JD_06232014_sample2_C.raw
                                           Condition2
     SUBJECT NESTED SUBJECT
## 1
                1.1
## 2
                1.1
## 3
                1.1
                           1
## 4
                           2
                2.2
                           2
## 5
                2.2
## 6
                2.2
# show which summarization method is used.
head(quant.processed$SummaryMethod)
## [1] "TMP"
```

6. Visualization of processed data

6.1. Quality control plots

Now let's look at what the equalize medians procedure did to our data. QC plot is good to see the distribution of intensities per MS run and outliers. So, it is good visualization to check normalization. However, not good to see individual intensities.

Then, iPRG_skyline_equalizeNorm_QCPlot.pdf are generated in the currect directory.

Now the median log2 intensities per run across MS runs are the same.

6.2. Profile plots

Profile plot is good visualization to check individual measurements. Each dot means one intensity. The dots are linked with line per feature. If line is disconnected, that means there is no value (missing value). Color means different peptides and charge stages. different line type means different transition.

iPRG_skyline_equalizeNorm_ProfilePlot.pdf and iPRG_skyline_equalizeNorm_ProfilePlot_wSummarization.pdf are generated in the current directory.

Then, Let's go though profile plots to see overall quality of data.

All proteins

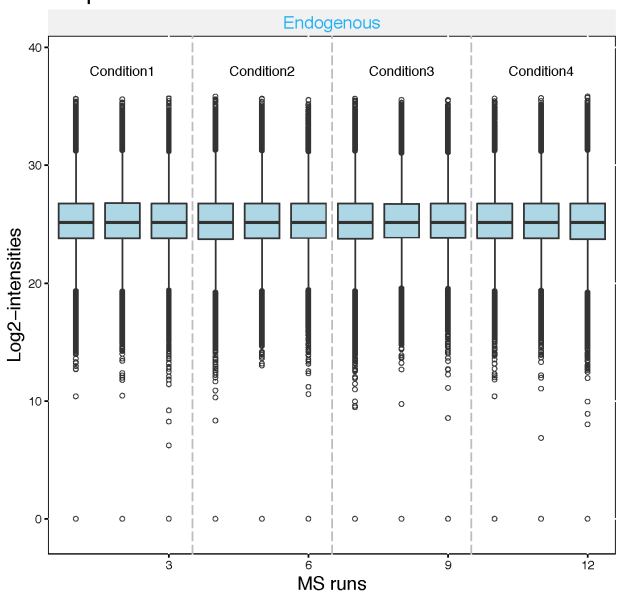


Figure 1:

There are two pdfs for each protein, first is profile plot with normalized data and second plot is profile plot with normalized data and summarized data. This profile plot shows each peptide transition across runs, grouped per condition. Ech peptide has a different colour/type layout.

spID6VTK4ISTE2_YEAST

• Detected data

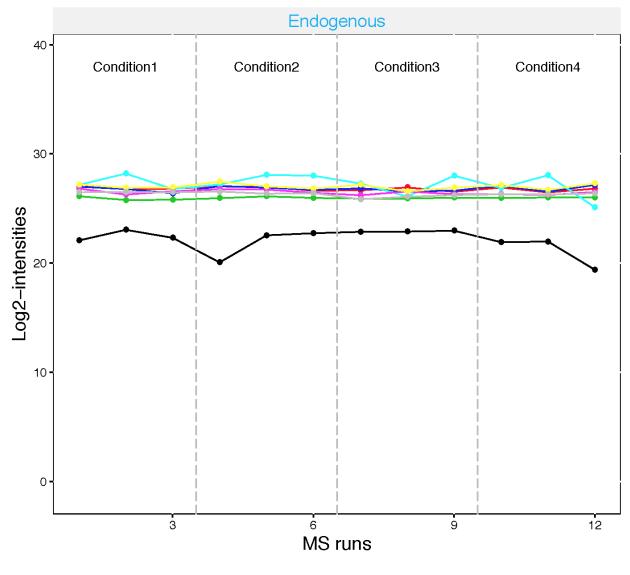


Figure 2:

This plot shows The panel on the right shows the same transitions in grey, with the values as summarized by the model overlayed in red.

Instead of making all profiles plots for all proteins, we can make plot for individual protein. Here is the example of spike-in protein, sp|P44015|VAC2_YEAST

spID6VTK4ISTE2_YEAST

Processed feature–level data → Run summary
 Detected data

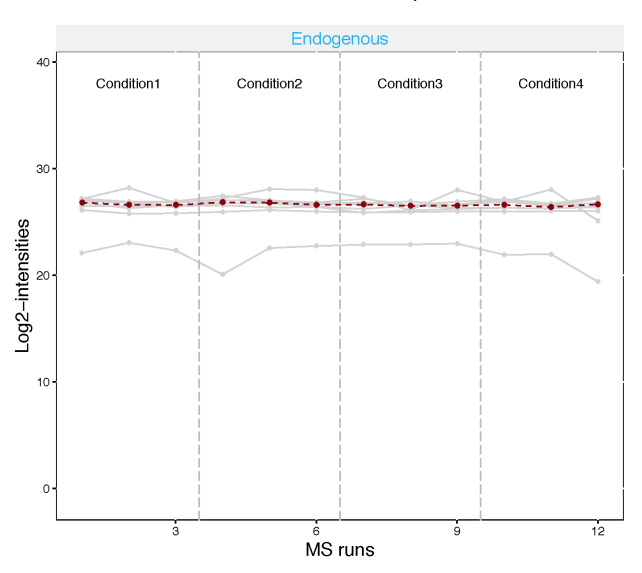


Figure 3:

```
width=7, height=7,
which.Protein = 'sp|P44015|VAC2_YEAST',
address="iPRG_skyline_equalizeNorm_P44015_")
```

```
## Warning: Ignoring unknown aesthetics: linetype
## Drew the Profile plot for sp|P44015|VAC2_YEAST ( 1 of 1 )
## Warning: Ignoring unknown aesthetics: linetype
## Drew the Profile plot with summarization for sp|P44015|VAC2_YEAST ( 1 of 1 )
```

spIP44015IVAC2_YEAST

• Detected data O Censored missing data

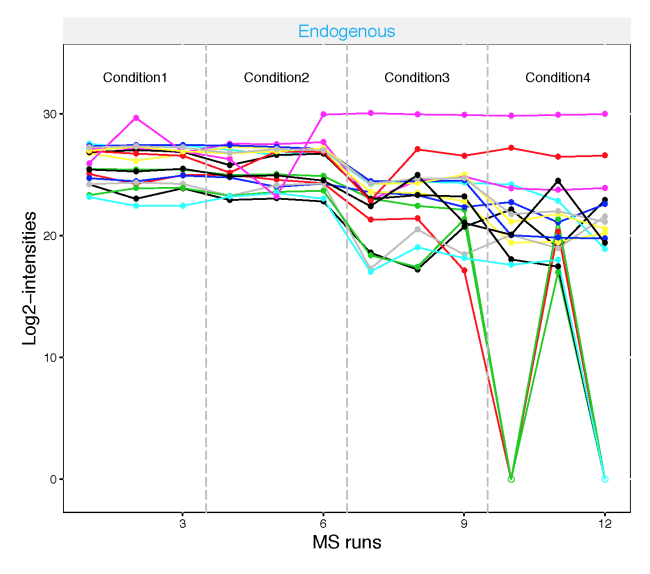


Figure 4:

This is the study design. Let's check visualization for those proteins.

spIP44015IVAC2_YEAST

Processed feature–level data → Run summary
 Detected data ○ Censored missing data

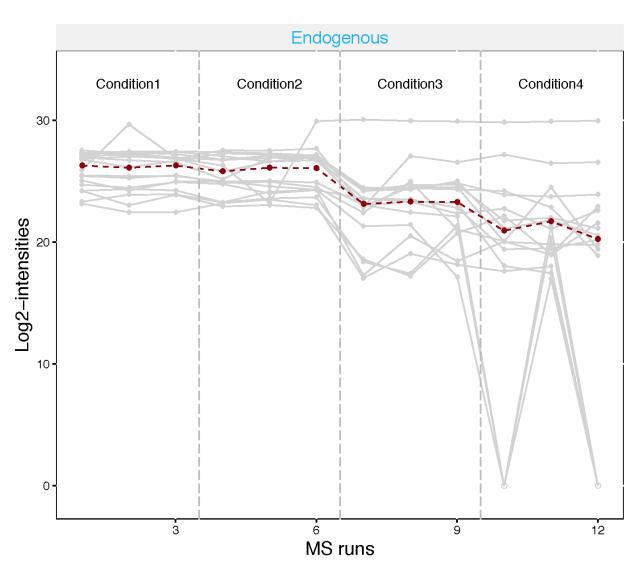


Figure 5:

	Samples				
Protein name	1	2	3	4	
sp P44015 VAC2_YEAST	65	55	15	2	
sp P55752 ISCB_YEAST	55	15	2	65	
sp P44374 SFG2_YEAST	15	2	65	55	
sp P44983 UTR6_YEAST	2	65	55	15	
sp P44683 PGA4_YEAST	11	0.6	10	500	
sp P55249 ZRT4_YEAST	10	500	11	0.6	

Figure 6:

MSstats needs at least one measurement per run. If not, can not get run-summarized value.

6.3. Condition plots

Condition plots illustrate the systematic difference between conditions. The dots indicates the mean of all measurements (Peptide ion and multiple MS runs) in each condition and default error bar is CI with 0.95 significant level. However, it is not related with model-based analysis.

This is the condition plot for protein, sp|P44015|VAC2_YEAST.

7. Finding differentially abundant proteins across conditions

7.1 Assign contrast matrix

After we normalized the data and summarized each protein's behaviour across conditions in dataProcess step, we are all set to compare protein changes between groups of conditions. Within MSstats we can do this with the groupComparison function, which takes as input the output of the dataProcess function.

```
?groupComparison
```

We have to tell groupComparison which are the conditions we would like to compare. You can make your contrast.matrix in R in a text editor. We define our contrast matrix by adding a column for every condition, in alphabetical order. We add a row for every comparison we would like to make between groups of conditions.

^{**!} Extreme case: no measurement in certain run.**

spIP44015IVAC2_YEAST

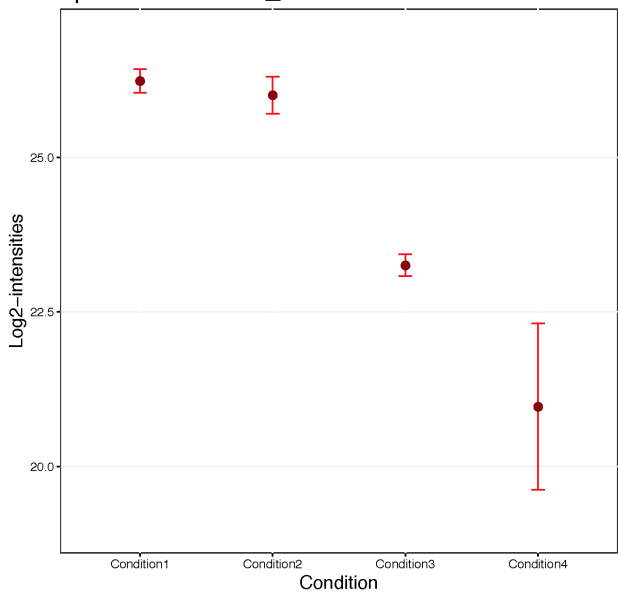


Figure 7:

 ${f 0}$ is for conditions we would like to ignore. ${f 1}$ is for conditions we would like to put in the numerator of the ratio or fold-change. ${f -1}$ is for conditions we would like to put in the denumerator of the ratio or fold-change.

If you have multiple groups, you can assign any gruop comparisons you are interested in. For example, if you have 4 different conditions, Condition1, Condition2, Condition3, Condition4, there are many possible comparisons.

```
# check unique conditions and check order of condition information
# In this case, Disease and Healthy
unique(quant.processed$ProcessedData$GROUP ORIGINAL)
## [1] Condition1 Condition2 Condition3 Condition4
## Levels: Condition1 Condition2 Condition3 Condition4
comparison1<-matrix(c(-1,1,0,0),nrow=1)
comparison2<-matrix(c(-1,0,1,0),nrow=1)
comparison3 < -matrix(c(-1,0,0,1),nrow=1)
comparison4 < -matrix(c(0,-1,1,0),nrow=1)
comparison5<-matrix(c(0,-1,0,1),nrow=1)
comparison6 < -matrix(c(0,0,-1,1),nrow=1)
comparison<-rbind(comparison1, comparison2, comparison3, comparison4, comparison5, comparison6)
row.names(comparison)<-c("C2-C1","C3-C1","C4-C1","C3-C2","C4-C2","C4-C3")
comparison
         [,1] [,2] [,3] [,4]
##
## C2-C1
           -1
                 1
## C3-C1
           -1
                 0
                      1
## C4-C1
          -1
                0
                      Ω
                           1
## C3-C2
           0
               -1
                      1
                           0
## C4-C2
           0
              -1
                      0
                           1
## C4-C3
                 0
                     -1
                           1
```

7.2. Comparing conditions with groupComparison

groupComparison uses the run-level summarized data (\$RunlevelData from dataProcess function) for hypothesis testing.

```
test <- groupComparison(contrast.matrix=comparison, data=quant.processed)</pre>
```

Let's check the output.

```
##
                   Protein Label
                                      log2FC
## 1
      sp|013297|CET1 YEAST C2-C1 -0.11119732 0.07749333 -1.4349278
## 13 sp|013329|F0B1_YEAST C2-C1 -0.16209422 0.29090086 -0.5572146
      sp|013539|THP2_YEAST C2-C1 -0.43742107 0.82871351 -0.5278315
## 25 sp|013547|CCW14 YEAST C2-C1 -0.05774773 0.14672405 -0.3935805 8
## 31 sp|013563|RPN13 YEAST C2-C1 -0.16945187 0.09518591 -1.7802201 8
         pvalue adj.pvalue issue MissingPercentage ImputationPercentage
## 1 0.4396100 0.9991155
                                                                     0
## 7 0.1892233 0.9991155
                             NA
                                                                     0
                                                0
## 13 0.5926243 0.9991155
                             NA
                                                0
                                                                     0
                                                                     0
## 19 0.6119387 0.9991155
                             NA
                                                0
## 25 0.7041724 0.9991155
                             NA
                                                0
                                                                     0
                                                                     0
## 31 0.1129126 0.9991155
                             NA
                                                0
# After fitting linear model, residuals and fitted values can be shown.
head(test$ModelQC)
##
    RJJN
                     PROTEIN ABUNDANCE NumMeasuredFeature MissingPercentage
## 1
       1 sp|D6VTK4|STE2 YEAST
                             26.81232
      2 sp|D6VTK4|STE2_YEAST
                              26.60786
                                                        8
                                                                          0
      3 sp|D6VTK4|STE2_YEAST
                              26.58301
                                                        8
                                                                          0
      4 sp|D6VTK4|STE2 YEAST
                                                                          0
## 4
                              26.83563
                                                        8
      5 sp|D6VTK4|STE2 YEAST
                                                        8
## 5
                              26.79430
                                                                          0
      6 sp|D6VTK4|STE2_YEAST 26.60863
                                                        8
                                                                          0
     more50missing NumImputedFeature
                                                  originalRUN GROUP
## 1
            FALSE
                                  0 JD_06232014_sample1_B.raw
                                                                  1
## 2
            FALSE
                                  0 JD_06232014_sample1_C.raw
                                                                  1
## 3
                                  0 JD_06232014_sample1-A.raw
            FALSE
                                                                  1
## 4
            FALSE
                                  0 JD_06232014_sample2_A.raw
                                                                  2
## 5
            FALSE
                                  0 JD_06232014_sample2_B.raw
                                                                  2
## 6
            FALSE
                                  0 JD_06232014_sample2_C.raw
     GROUP_ORIGINAL SUBJECT_ORIGINAL SUBJECT_NESTED SUBJECT
## 1
        Condition1
                                               1.1
                                                         1 0.14458899
                                  1
        Condition1
## 2
                                  1
                                               1.1
                                                         1 -0.05986794
## 3
        Condition1
                                  1
                                               1.1
                                                         1 -0.08472105
## 4
         Condition2
                                  2
                                               2.2
                                                         2 0.08944428
## 5
         Condition2
                                  2
                                               2.2
                                                         2 0.04811446
## 6
         Condition2
                                  2
                                               2.2
                                                         2 -0.13755874
##
      fitted
## 1 26.66773
## 2 26.66773
## 3 26.66773
## 4 26.74619
## 5 26.74619
## 6 26.74619
# Fitted model per protein
head(test$fittedmodel)
## [[1]]
##
## lm(formula = ABUNDANCE ~ GROUP, data = data2)
## Coefficients:
```

```
## (Intercept)
                     GROUP2
                                   GROUP3
                                                 GROUP4
      26.66773
##
                     0.07846
                                 -0.11566
                                               -0.11841
##
##
## [[2]]
##
## lm(formula = ABUNDANCE ~ GROUP, data = data2)
##
## Coefficients:
## (Intercept)
                      GROUP2
                                   GROUP3
                                                 GROUP4
      24.70386
##
                   -0.11120
                                 -0.14451
                                               -0.09633
##
##
## [[3]]
##
## Call:
## lm(formula = ABUNDANCE ~ GROUP, data = data2)
## Coefficients:
## (Intercept)
                      GROUP2
                                   GROUP3
                                                 GROUP4
##
       23.6207
                    -0.1621
                                   0.1209
                                                -0.3511
##
##
## [[4]]
## Call:
## lm(formula = ABUNDANCE ~ GROUP, data = data2)
## Coefficients:
## (Intercept)
                      GROUP2
                                   GROUP3
                                                 GROUP4
##
       26.4006
                     -0.4374
                                  -0.4318
                                                -0.5024
##
##
## [[5]]
## Call:
## lm(formula = ABUNDANCE ~ GROUP, data = data2)
## Coefficients:
## (Intercept)
                      GROUP2
                                   GROUP3
                                                 GROUP4
      27.04775
##
                   -0.05775
                                 -0.29114
                                               -0.09482
##
##
## [[6]]
##
## Call:
## lm(formula = ABUNDANCE ~ GROUP, data = data2)
## Coefficients:
## (Intercept)
                      GROUP2
                                   GROUP3
                                                 GROUP4
      26.18595
                   -0.16945
                                 -0.07609
                                               -0.21625
```

Let's save the testing result as .csv file.

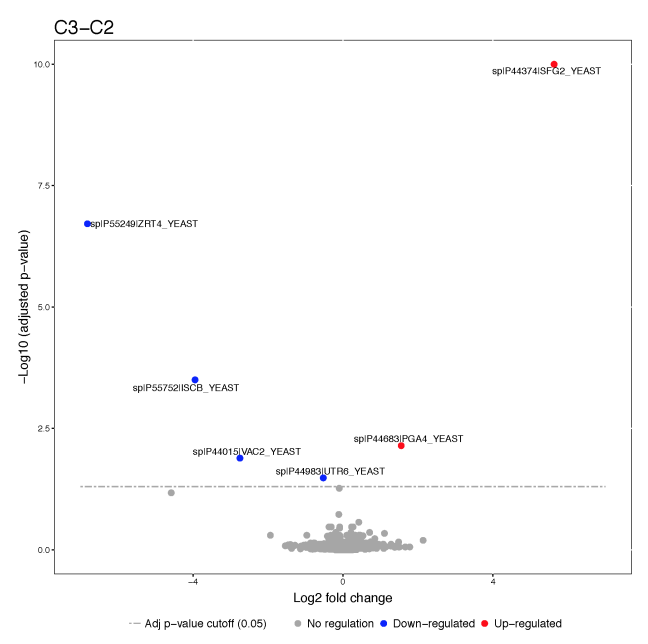
```
Skyline.intensity.comparison.result <- test$ComparisonResult
write.csv(Skyline.intensity.comparison.result, file='testResult_iprg_skyline.csv')
Let's inspect the results to see what proteins are changing significantly between Diseased and Healthy.
head(Skyline.intensity.comparison.result)
##
                    Protein Label
                                       log2FC
                                                              Tvalue DF
## 1
       sp|D6VTK4|STE2_YEAST C2-C1  0.07846162  0.09648008  0.8132416
       sp|013297|CET1 YEAST C2-C1 -0.11119732 0.07749333 -1.4349278 8
## 13 sp|013329|F0B1 YEAST C2-C1 -0.16209422 0.29090086 -0.5572146 8
       sp|013539|THP2_YEAST C2-C1 -0.43742107 0.82871351 -0.5278315
## 25 sp|013547|CCW14 YEAST C2-C1 -0.05774773 0.14672405 -0.3935805 8
## 31 sp|013563|RPN13_YEAST C2-C1 -0.16945187 0.09518591 -1.7802201 8
         pvalue adj.pvalue issue MissingPercentage ImputationPercentage
##
## 1 0.4396100 0.9991155
                              NA
## 7 0.1892233 0.9991155
                              NA
                                                  0
                                                                       0
## 13 0.5926243 0.9991155
                              NA
                                                  0
                                                                       0
                                                                       0
## 19 0.6119387
                 0.9991155
                              NA
                                                  0
## 25 0.7041724
                 0.9991155
                              NA
                                                  0
                                                                       0
## 31 0.1129126 0.9991155
                              NA
                                                  0
                                                                       0
SignificantProteins <-
  Skyline.intensity.comparison.result[Skyline.intensity.comparison.result$adj.pvalue < 0.05 ,]
nrow(SignificantProteins)
## [1] 29
SignificantProteinsUpInDiseased <- SignificantProteins[SignificantProteins$log2FC > 2 ,]
nrow(SignificantProteinsUpInDiseased)
## [1] 14
```

8. Visualization of differentially abundant proteins

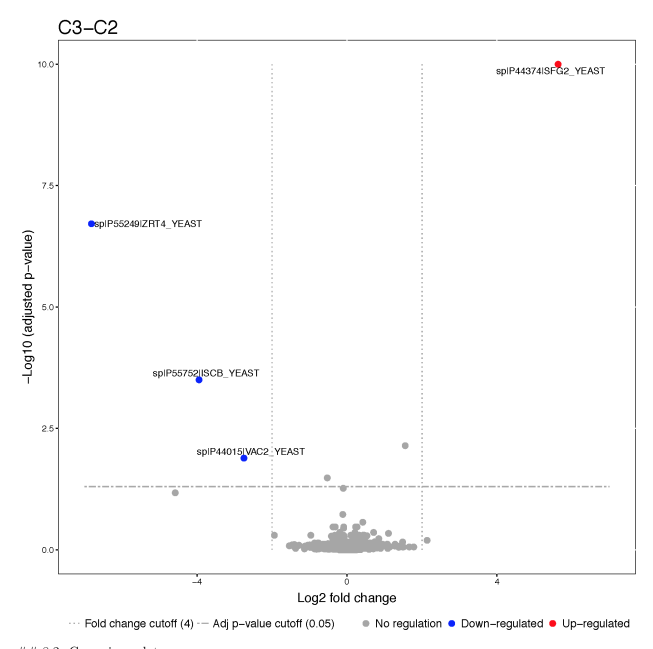
?groupComparisonPlots

8.1. Volcano plot

Volcano plots allow us to visually separate strong changes, which are not significant, from strong and significant changes. Look for these subjects in the upper right and upper left quadrants of the plot. Protein name will be shown only for significant proteins.



We can set up estimated fold change cutoff.



8.2. Comarison plot

Comparison plots illustrate model-based estimates of log-fold changes, and the associated uncertainty, in several comparisons of conditions for one protein. X-axis is the comparison of interest. Y-axis is the log fold change. The dots are the model-based estimates of log-fold change, and the error bars are the model-based 95% confidence intervals (the option sig can be used to change the significance level of significance). For simplicity, the confidence intervals are adjusted for multiple comparisons within protein only, using the Bonferroni approach. For proteins with N comparisons, the individual confidence intervals are at the level of 1-sig/N.

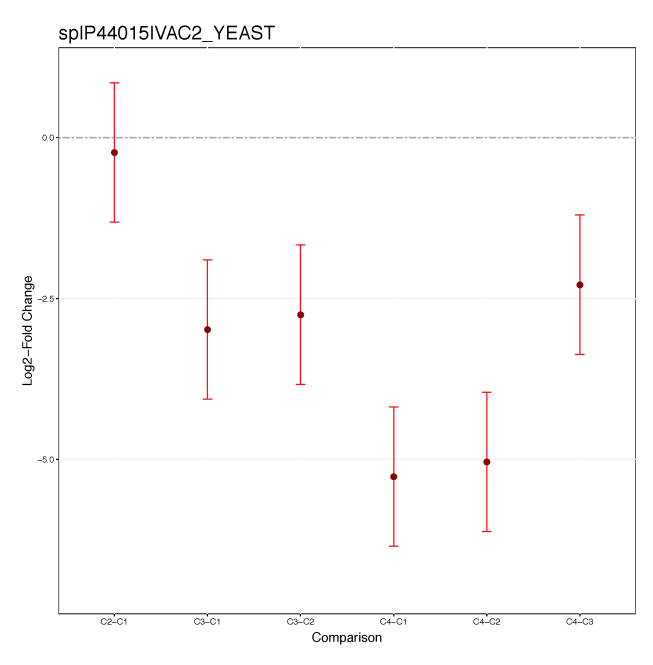


Figure 8:

9. Planning future experimental designs

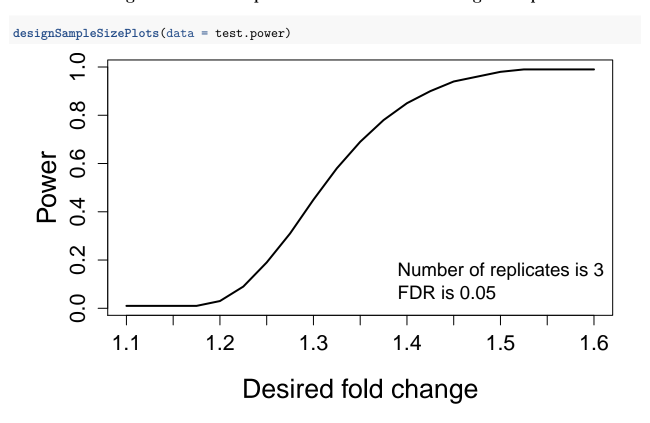
This last analysis step views the dataset as a pilot study of a future experiment, utilizes its variance components, and calculates the minimal number of replicates required in a future experiment to achieve the desired statistical power. The calculation is performed by the function designSampleSize, which takes as input the fitted model in groupComparison. Sample size calculation assumes same experimental design (i.e. group comparison, time course or paired design) as in the current dataset, and uses the model fit to estimate the median variance components across all the proteins. Finally, sample size calculation assumes that a large proportion of proteins (specifically, 99%) will not change in abundance in the future experiment. This assumption also provides conservative results. Using the estimated variance components, the function relates the number of biological replicates per condition (numSample, rounded to 0 decimal), average statistical power across all the proteins (power), minimal fold change that we would like to detect (can be specified as a range, e.g. desiredFC=c(1.1, 2)), and the False Discovery Rate (FDR). The user should specify all these quantities but one, and the function will solve for the remainder. The quantity to solve for should be set to = TRUE.

?designSampleSize

9.1. Calculating statistical power

```
##
      desiredFC numSample
                             FDR power
## 1
           1.100
                          3 0.05
                                  0.01 0.010
## 2
           1.125
                          3 0.05
                                  0.01 0.010
                          3 0.05
## 3
           1.150
                                  0.01 0.010
                          3 0.05
## 4
           1.175
                                  0.01 0.010
## 5
           1.200
                          3 0.05
                                  0.03 0.009
## 6
           1.225
                          3 0.05
                                  0.09 0.009
                          3 0.05
## 7
          1.250
                                  0.19 0.009
## 8
          1.275
                          3 0.05
                                  0.31 0.009
## 9
                          3 0.05
                                  0.45 0.009
          1.300
## 10
          1.325
                          3 0.05
                                  0.58 0.009
                          3 0.05
## 11
          1.350
                                  0.69 0.008
## 12
          1.375
                          3 0.05
                                  0.78 0.008
## 13
          1.400
                          3 0.05
                                  0.85 0.008
## 14
          1.425
                          3 0.05
                                  0.90 0.008
## 15
          1.450
                          3 0.05
                                  0.94 0.008
                          3 0.05
## 16
          1.475
                                  0.96 0.008
## 17
          1.500
                          3 0.05
                                  0.98 0.008
## 18
          1.525
                          3 0.05
                                  0.99 0.007
  19
          1.550
                          3 0.05
                                  0.99 0.007
                          3 0.05
## 20
           1.575
                                  0.99 0.007
                          3 0.05
                                  0.99 0.007
## 21
           1.600
```

9.2. Visualizing the relationship between desired fold-change and power



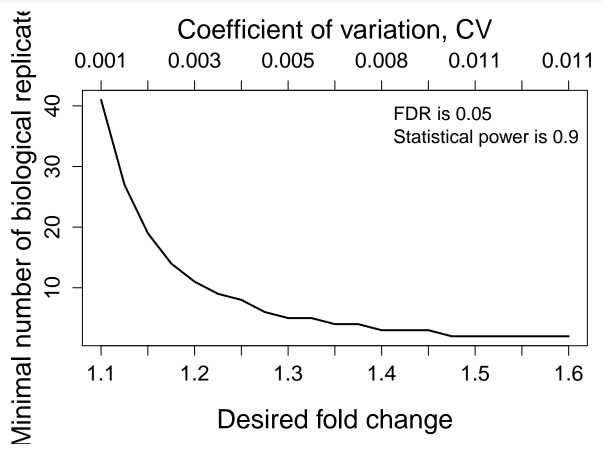
9.3. Designing sample size for desired fold-change

```
desiredFC numSample FDR power
##
## 1
          1.100
                        41 0.05
                                   0.9 0.001
## 2
          1.125
                        27 0.05
                                   0.9 0.001
## 3
          1.150
                        19 0.05
                                   0.9 0.002
## 4
          1.175
                        14 0.05
                                   0.9 0.002
## 5
          1.200
                        11 0.05
                                   0.9 0.003
## 6
          1.225
                         9 0.05
                                   0.9 0.003
## 7
          1.250
                         8 0.05
                                   0.9 0.003
## 8
          1.275
                         6 0.05
                                   0.9 0.004
                         5 0.05
## 9
          1.300
                                   0.9 0.005
                         5 0.05
## 10
          1.325
                                   0.9 0.005
## 11
          1.350
                         4 0.05
                                   0.9 0.006
## 12
          1.375
                         4 0.05
                                   0.9 0.006
## 13
          1.400
                         3 0.05
                                   0.9 0.008
                         3 0.05
                                   0.9 0.008
## 14
          1.425
```

```
## 15
           1.450
                          3 0.05
                                    0.9 0.008
##
  16
           1.475
                          2 0.05
                                    0.9 0.012
##
  17
           1.500
                          2 0.05
                                    0.9 0.011
                          2 0.05
  18
           1.525
                                    0.9 0.011
##
##
  19
           1.550
                          2 0.05
                                    0.9 0.011
## 20
           1.575
                          2 0.05
                                    0.9 0.011
## 21
           1.600
                          2 0.05
                                    0.9 0.011
```

9.4. Visualizing the relationship between desired fold-change and minimum sample size number





10. Protein subject quantification

If there is no technical replicate, subject (or sample) quantification should be the same as run-level summarization. If there are technical replicates, subjet-level summarization (quantification) with run-level summarization will be useful for downstream analysis, such as classification.

?quantification

```
sampleQuant <- quantification(quant.processed)
head(sampleQuant)</pre>
```

```
##
                   Protein Condition1_1 Condition2_2 Condition3_3
     sp|D6VTK4|STE2_YEAST
                               26.60786
                                             26.79430
                                                          26.53029
     sp|013297|CET1_YEAST
                               24.71809
                                             24.57865
                                                          24.62652
     sp|013329|F0B1_YEAST
                               23.47075
                                             23.43427
                                                          23.73741
## 4 sp|013539|THP2_YEAST
                               27.38510
                                             25.90646
                                                          25.91799
                                                          26.75541
## 5 sp|013547|CCW14_YEAST
                               27.11638
                                            26.91302
## 6 sp|013563|RPN13_YEAST
                               26.17056
                                             26.01078
                                                          26.11412
    Condition4_4
## 1
         26.60612
## 2
         24.64886
## 3
         23.16646
## 4
         25.91781
         26.98082
## 6
         26.05415
```

11. msstats.log and sessionInfo.txt

These two files are important to keep the records of package versions and options in functions.

```
"R.version.3.3.2..2016.10.31."
"Platform: x86_64-apple-darwin13.4.0 (64-bit)"
"Running under: OS X El Capitan 10.11.6"
"locale:"
"[1] en_US.UTF-8/en_US.UTF-8/en_US.UTF-8/c/en_US.UTF-8"
"attached base packages:"
"[1] stats
               graphics grDevices utils
                                             datasets methods
                                                                  base
"other attached packages:"
"[1] MSstats_3.7.2"
"loaded via a namespace (and not attached):"
" [1] Rcpp_0.12.7
                          magrittr_1.5
                                              BiocGenerics_0.18.0"
" [4] splines_3.3.2
                          MASS 7.3-45
                                              munsell 0.4.3
" [7] colorspace_1.2-6
                          lattice_0.20-34
                                              minqa_1.2.4
"[10] stringr_1.1.0
                          plyr_1.8.4
                                              caTools_1.17.1
"[13] tools_3.3.2
                          parallel_3.3.2
                                              grid_3.3.2
"[16] Biobase_2.32.0
                                              nlme_3.1-128
                          gtable_0.2.0
"[19] KernSmooth_2.23-15
                          marray_1.50.0
                                              mzR_2.6.3
"[22] gtools_3.5.0
                                              survival_2.39-5
                          ProtGenerics_1.4.0
"[25] lme4_1.1-12
                                              assertthat_0.1
                          lazyeval_0.2.0
"[28] tibble_1.2
                          Matrix_1.2-7.1
                                              reshape2_1.4.1
"[31] nloptr_1.0.4
                          ggplot2_2.2.0
                                              bitops_1.0-6
"[34] codetools_0.2-15
                          ggrepel_0.5
                                              stringi_1.1.1
                                             gplots_3.0.1
"[37] limma_3.28.21
                          gdata_2.17.0
"[40] scales_0.4.1
                          minpack.lm 1.2-0
"MSstats - dataProcess function"
"The required input : provided - okay"
"summaryMethod : TMP"
"cutoffCensored : minFeature"
"censoredInt : 0"
"New input format : made new columns for analysis - okay"
"** There are 2585 intensities which are zero. These intensities are replaced with 1."
"Logarithm transformation: log2 transformation is done - okay"
"fillIncompleteRows = TRUE"
"Balanced data format with NA for missing feature intensities - okay"
"Factorize in columns(GROUP, SUBJECT, GROUP_ORIGINAL, SUBJECT_ORIGINAL, SUBJECT_ORIGINAL_NESTED,
FEATURE, RUN) - okay"
"Normalization : Constant normalization (equalize medians) - okay"
"Between Run Interference Score is not calculated."
"* Use all features that the dataset originally has."
"1 level of Isotope type labeling in this experiment"
"Summary of Features:"
"# of Protein : 3027"
"# of Peptides/Protein : 2-196"
"# of Transitions/Peptide : 1-1"
```

Figure 9: