Analysis and Exploration of Proteomics Data

R Markdown

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Proteomics Data Analysis

Below are shown the steps typically used to perform Differential Gene Expression analysis. The analysis steps have been applied over the *EGF-driven protein synthesis* transcriptomics data from D.A. Rothenberg et al. A Proteomics Approach to Profiling the Temporal Translational Response to Stress and Growth. iScience. 2018; 9:367-381.

Loading of R-Packages

We start by loading the R-packages we need to use for our analysis:

```
library(edgeR)
library(lattice)
library(biomaRt)
library(gplots)
library(gridExtra)
library(openxlsx)
library(tools)
library(ggplot2)
library(dplyr)
library(Glimma)
library(openxlsx)
```

Loading of the data

We load the generated **subread_counts** object, define the experimental conditions for each available sample and prepare the data in the DGEObj format.

```
load("course.Rdata")
groups <- factor(c("PBS","EGF","PBS","EGF"))

label <- c("PBS2", "EGF1", "PBS1", "EGF2")

print(colnames(subread_counts$counts))

## [1] "SRR7451179.Rep.2.MP60.2.mRNA.Seq.PBS.bam"

## [2] "SRR7451182.Rep.1.ME60.1.mRNA.Seq.EGF.bam"

## [3] "SRR7451187.Rep.1.MP60.1.mRNA.Seq.PBS.bam"

## [4] "SRR7451201.Rep.2.ME60.2.mRNA.Seq.EGF.bam"

print(colnames(subread_counts$counts))</pre>
```

```
## [2] "SRR7451182.Rep.1.ME60.1.mRNA.Seq.EGF.bam"
## [3] "SRR7451187.Rep.1.MP60.1.mRNA.Seq.PBS.bam"
## [4] "SRR7451201.Rep.2.ME60.2.mRNA.Seq.EGF.bam"
DGEObj <- DGEList(group=groups, counts=subread_counts$counts, genes=subread_counts$annotation[,c("Gene I
print(DGEObj)
## An object of class "DGEList"
## $counts
                   SRR7451179.Rep.2.MP60.2.mRNA.Seq.PBS.bam
## ENSG00000223972
## ENSG00000227232
                                                           14
## ENSG00000278267
                                                            0
## ENSG00000243485
                                                            0
                                                            0
## ENSG00000284332
##
                   SRR7451182.Rep.1.ME60.1.mRNA.Seq.EGF.bam
## ENSG00000223972
## ENSG00000227232
                                                           24
## ENSG00000278267
                                                            0
                                                            0
## ENSG00000243485
## ENSG00000284332
                   SRR7451187.Rep.1.MP60.1.mRNA.Seq.PBS.bam
## ENSG00000223972
## ENSG00000227232
                                                           10
## ENSG00000278267
                                                            0
## ENSG00000243485
                                                            0
## ENSG00000284332
                                                            0
##
                   SRR7451201.Rep.2.ME60.2.mRNA.Seq.EGF.bam
## ENSG00000223972
                                                           22
## ENSG00000227232
## ENSG00000278267
                                                            0
## ENSG00000243485
                                                            0
## ENSG00000284332
                                                            0
## 58879 more rows ...
## $samples
                                             group lib.size norm.factors
##
## SRR7451179.Rep.2.MP60.2.mRNA.Seq.PBS.bam
                                              PBS 26653631
                                                                        1
## SRR7451182.Rep.1.ME60.1.mRNA.Seq.EGF.bam
                                               EGF 30763088
                                                                        1
## SRR7451187.Rep.1.MP60.1.mRNA.Seq.PBS.bam
                                               PBS 28377976
                                                                        1
## SRR7451201.Rep.2.ME60.2.mRNA.Seq.EGF.bam
                                               EGF 31739139
                                                                        1
##
## $genes
##
                             GeneID Length
## ENSG00000223972 ENSG00000223972
                                      1735
## ENSG00000227232 ENSG00000227232
                                      1351
## ENSG00000278267 ENSG00000278267
                                        68
## ENSG00000243485 ENSG00000243485
                                      1021
## ENSG00000284332 ENSG00000284332
                                       138
## 58879 more rows ...
```

Experimental Design

We make an experimental design matrix where we indicate the condition and replicate ID's for each sample. We then prepare the needed EGF_vs_PBS contrast.

```
design <- model.matrix(~0+groups)</pre>
colnames(design) <- levels(groups)</pre>
print(design)
     EGF PBS
## 1
       0
          1
## 2
      1
## 3
       0 1
## 4
       1
## attr(,"assign")
## [1] 1 1
## attr(,"contrasts")
## attr(,"contrasts")$groups
## [1] "contr.treatment"
contrasts <- makeContrasts(</pre>
  EGF_vs_PBS = EGF - PBS,
  levels=design
)
counts <- DGEObj$counts</pre>
```

DGEObj Processing

We simplify a bit the names of the samples.

head(DGEObj\$samples)

```
##
                                              group lib.size norm.factors
## SRR7451179.Rep.2.MP60.2.mRNA.Seq.PBS.bam PBS 26653631
## SRR7451182.Rep.1.ME60.1.mRNA.Seq.EGF.bam
                                                EGF 30763088
                                                                          1
## SRR7451187.Rep.1.MP60.1.mRNA.Seq.PBS.bam
                                                PBS 28377976
                                                                          1
## SRR7451201.Rep.2.ME60.2.mRNA.Seq.EGF.bam
                                                EGF 31739139
rownames(DGEObj$samples) <- gsub("\\.", "_", rownames(DGEObj$samples))</pre>
rownames(DGEObj$samples) <- gsub("KD_", "", rownames(DGEObj$samples))</pre>
rownames(DGEObj$samples) <- gsub(" circSLC8A1", "", rownames(DGEObj$samples))</pre>
rownames(DGEObj$samples) <- gsub("_bam", "", rownames(DGEObj$samples))</pre>
rownames(DGEObj$counts) <- gsub("_1_mRNA_Seq_PBS", "", rownames(DGEObj$counts))</pre>
rownames(DGEObj$counts) <- gsub("_1_mRNA_Seq_EGF", "", rownames(DGEObj$counts))</pre>
rownames(DGEObj$counts) <- gsub("_2 mRNA_Seq_PBS", "", rownames(DGEObj$counts))
rownames(DGEObj$counts) <- gsub("_2 mRNA_Seq_EGF", "", rownames(DGEObj$counts))
colnames(DGEObj$counts) <- gsub("\\.", "_", colnames(DGEObj$counts))</pre>
colnames(DGEObj$counts) <- gsub("KD_", "", colnames(DGEObj$counts))</pre>
colnames(DGEObj$counts) <- gsub("_1_mRNA_Seq_PBS", "", colnames(DGEObj$counts))</pre>
colnames(DGEObj$counts) <- gsub("_1_mRNA_Seq_EGF", "", colnames(DGEObj$counts))</pre>
colnames(DGEObj$counts) <- gsub("_2_mRNA_Seq_PBS", "", colnames(DGEObj$counts))</pre>
colnames(DGEObj$counts) <- gsub("_2_mRNA_Seq_EGF", "", colnames(DGEObj$counts))</pre>
colnames(DGEObj$counts) <- gsub("_bam", "", colnames(DGEObj$counts))</pre>
DGEObj$samples
```

```
group lib.size norm.factors
##
## SRR7451179_Rep_2_MP60_2_mRNA_Seq_PBS
                                           PBS 26653631
## SRR7451182 Rep 1 ME60 1 mRNA Seq EGF
                                            EGF 30763088
## SRR7451187_Rep_1_MP60_1_mRNA_Seq_PBS
                                           PBS 28377976
                                                                     1
## SRR7451201_Rep_2_ME60_2_mRNA_Seq_EGF
                                           EGF 31739139
head(DGEObj$counts)
                   SRR7451179_Rep_2_MP60 SRR7451182_Rep_1_ME60
##
## ENSG00000223972
                                        0
## ENSG00000227232
                                       14
                                                              24
## ENSG00000278267
                                        0
                                                               0
## ENSG00000243485
                                        0
                                                                0
## ENSG00000284332
                                        0
                                                                0
## ENSG00000237613
                                        0
                                                                0
##
                   SRR7451187_Rep_1_MP60 SRR7451201_Rep_2_ME60
## ENSG00000223972
## ENSG00000227232
                                       10
                                                              22
## ENSG00000278267
                                        0
                                                                0
                                                               0
                                        0
## ENSG00000243485
## ENSG00000284332
                                        0
                                                                0
## ENSG00000237613
                                        0
                                                               0
head(DGEObj$genes)
                             GeneID Length
##
## ENSG00000223972 ENSG00000223972
## ENSG00000227232 ENSG00000227232
                                      1351
## ENSG00000278267 ENSG00000278267
                                        68
## ENSG00000243485 ENSG00000243485
                                      1021
## ENSG00000284332 ENSG00000284332
                                       138
## ENSG00000237613 ENSG00000237613
                                      1219
DGEOb j
## An object of class "DGEList"
## $counts
##
                   SRR7451179_Rep_2_MP60 SRR7451182_Rep_1_ME60
## ENSG00000223972
                                        0
                                                               0
## ENSG00000227232
                                       14
                                                              24
## ENSG00000278267
                                        0
                                                               0
## ENSG00000243485
                                        0
                                                                0
## ENSG00000284332
                                        0
                                                                0
                   SRR7451187_Rep_1_MP60 SRR7451201_Rep_2_ME60
## ENSG00000223972
                                        0
                                                               0
## ENSG00000227232
                                       10
                                                              22
## ENSG00000278267
                                        0
                                                               0
## ENSG00000243485
                                        0
                                                               0
## ENSG00000284332
                                        0
                                                                0
## 58879 more rows ...
##
## $samples
                                         group lib.size norm.factors
##
                                           PBS 26653631
## SRR7451179_Rep_2_MP60_2_mRNA_Seq_PBS
## SRR7451182_Rep_1_ME60_1_mRNA_Seq_EGF
                                           EGF 30763088
                                                                     1
## SRR7451187_Rep_1_MP60_1_mRNA_Seq_PBS
                                           PBS 28377976
                                                                     1
```

```
## SRR7451201_Rep_2_ME60_2_mRNA_Seq_EGF
                                           EGF 31739139
##
## $genes
##
                            GeneID Length
## ENSG00000223972 ENSG00000223972
                                      1735
## ENSG00000227232 ENSG00000227232
                                      1351
## ENSG00000278267 ENSG00000278267
                                       68
## ENSG00000243485 ENSG00000243485
                                      1021
## ENSG00000284332 ENSG00000284332
                                      138
## 58879 more rows ...
```

Data Filtering

We filter the data by keeping only candidates with > 2 counts per million mapped reads AND in 3 libs.

```
keep <- rowSums(cpm(DGEObj) > 1) >= 3

# select sub set from above
DGEObj <- DGEObj[keep, keep.lib.sizes = FALSE]

dim(DGEObj)</pre>
```

```
## [1] 14691 4
```

edgeR Analysis

We calculate the normalization factor, estimate the dispersion, fit the linear model, and then summarize the results in an edgeR table format.

```
DGEObj <- calcNormFactors(DGEObj) # recalc norm factors

DGEObj <- estimateDisp(DGEObj, design) # estimate dispersion

DGEObj$common.dispersion</pre>
```

```
## [1] 0.001686058

fit <- glmFit(DGEObj, design) # fit generalized linear model
lrt <- glmLRT(fit, contrast = contrasts[, "EGF_vs_PBS"])

# main data table of edgeR results
edgerTable <- topTags(lrt, n = nrow(DGEObj))$table

y <- cpm(DGEObj, log = TRUE, prior.count = 1)
lcpm <- cpm(DGEObj, log = TRUE)</pre>
```

Mapping

We do mapping of the Ensembl gene ID's.

```
values = edgerTable$GeneID, mart = ensembl)
edgerTable.idx <- match(edgerTable$GeneID, genemap$ensembl_gene_id)
edgerTable.idx2 <- match(edgerTable$GeneID, genemap2$ensembl_gene_id)
# add biomart data to results tablesTRUE
edgerTable$external_gene_name <- genemap$external_gene_name[edgerTable.idx]</pre>
# edgerTable$entrezgene <- genemap$entrezgene[edgerTable.idx]
edgerTable$description <- genemap$description[edgerTable.idx]</pre>
edgerTable$GO <- genemap2$go_id[edgerTable.idx2]</pre>
edgerTable$ensembl_gene_id <- genemap$ensembl_gene_id[edgerTable.idx]</pre>
lrt$table$gene <- genemap$external_gene_name[edgerTable.idx]</pre>
gene_names <- data.frame(lrt$table$gene)</pre>
edgerTable <- edgerTable[c("external_gene_name", "GeneID", "Length", "logFC",
                           "logCPM", "PValue", "FDR", "GO", "description")]
ttop_dge <- edgerTable
head(ttop_dge)
##
                   external_gene_name
                                                GeneID Length
                                                                 logFC
## ENSG00000125740
                                 FOSB ENSG00000125740 5553 4.962212 6.902998
## ENSG0000138166
                                DUSP5 ENSG00000138166 2535 3.983957 6.291272
## ENSG00000119508
                                NR4A3 ENSG00000119508 6314 3.302585 8.107344
## ENSG0000175592
                                FOSL1 ENSG00000175592 1887 3.307155 5.967628
## ENSG0000171223
                                 JUNB ENSG00000171223
                                                       1830 3.283124 7.112127
## ENSG0000162772
                                 ATF3 ENSG00000162772 4040 3.119576 7.369775
                                                        GO
##
                          PValue
                                           FDR
## ENSG00000125740 0.000000e+00 0.000000e+00 GD:0003677
## ENSG00000138166 0.000000e+00 0.000000e+00 GD:0016791
## ENSG00000119508 0.000000e+00 0.000000e+00 GD:0003677
## ENSG00000175592 5.605306e-296 2.058689e-292 GD:0003677
## ENSG00000171223 1.089103e-281 3.200002e-278 GO:0003677
## ENSG00000162772 9.076298e-280 2.222331e-276 GD:0003677
                                                                                                   descri
## ENSG00000125740 FosB proto-oncogene, AP-1 transcription factor subunit [Source:HGNC Symbol;Acc:HGNC:
## ENSG0000138166
                                            dual specificity phosphatase 5 [Source: HGNC Symbol; Acc: HGNC:
## ENSG0000119508
                            nuclear receptor subfamily 4 group A member 3 [Source: HGNC Symbol; Acc: HGNC:
                           FOS like 1, AP-1 transcription factor subunit [Source: HGNC Symbol; Acc: HGNC: 1
## ENSG0000175592
## ENSG00000171223 JunB proto-oncogene, AP-1 transcription factor subunit [Source: HGNC Symbol; Acc: HGNC:
                                          activating transcription factor 3 [Source: HGNC Symbol; Acc: HGNC
## ENSG0000162772
```