VI. Bioinformatics in R (presentation)

Center for Health Data Science, University of Copenhagen

01 September, 2021

Bioconductor

Bioconductor provides tools for computational biology and bioinformatics analysis in R - it is open source and open development and it has an active user community.

Mostly when we install R-packages we use install.packages('name_of_package'). When we use this command we refer to the CRAN repository of packages, however sometimes we want a package from Bioconductor instead. For this we use the command BiocManager::install('name_of_package'). In order to use this installer, you need to download the R-package BiocManager e.g. install.packages('BiocManager').

Gene Expression Analysis in R with DEseq2

DEseq2 is one of the many packages/frameworks which exists for analysis of bulk gene expression data in R. For more information on DEseq2, please have a look at the original publication here.

Other highly used packages for differential expression analysis DEA are:

- limma
- edgeR
- NOIseq

DEseq2 has many advantages over classical models and post hoc tests, as it is specifically developed for handling common issues and biases in expression data, including differences in sequencing depth and highly variable dispersion of counts between genes.

In brief, DEseq2 fits a generalized linear model (GLM) for each gene in the dataset. In the case where we compare two groups i.e. treatment vs control, the GLM fit returns coefficients indicating the overall expression strength of a gene, along with the log-2 fold change between groups. DEseq2 adjusts variable gene dispersion estimates using an empirical Bayes approach which borrows information across genes and shrinks gene-wise dispersions towards a common dispersion trend to increase accuracy of differential expression testing.

About the Dataset

The dataset used for this presentation was acquired from the following github tutorial on RNAseq analysis: https://combine-australia.github.io/RNAseq-R/06-rnaseq-day1.html.

RNA sequencing data generated from luminal and basal cell sub-populations in the mammary gland of three groups of mice:

- Control
- Pregnant
- Lactating

The objective of the original study (found here) was to identify genes specifically expressed in lactating mammary glands, the gene expression profiles of luminal and basal cells from different developmental stages were compared.

Load R-packages:

```
# Data Wrangling
# install.packages("tidyverse")
# install.packages("readxl")
library(tidyverse)
library(readxl)

# For Plotting
# install.packages("ggplot2")
library(ggplot2)

# For DEA
# install.packages("BiocManager")
# BiocManager::install("DESeq2")
library(BiocManager)
library(DESeq2)
```

Importing Data

Reading in data:

```
exprDat <- read_excel("MouseRNAseq.xlsx")
exprInfo <- read_excel("MouseSampleInfo.xlsx")

# Look at the data:
head(exprDat, n=5)</pre>
```

```
## # A tibble: 5 x 14
     EntrezGeneID GeneName MCL1.DG MCL1.DH MCL1.DI MCL1.DJ MCL1.DK MCL1.DL MCL1.LA
##
                   <chr>>
                                       <dbl>
                                                        <dbl>
                                                                <dbl>
                                                                         <dbl>
                                                                                 <dbl>
##
     <chr>>
                              <dbl>
                                               <dbl>
## 1 497097
                   Xkr4
                                438
                                         300
                                                  65
                                                          237
                                                                  354
                                                                           287
                                                                                     0
## 2 19888
                                                   0
                                                                    0
                                                                             0
                                                                                    10
                   Rp1
                                  1
                                           1
                                                            0
## 3 20671
                   Sox17
                                106
                                         182
                                                  82
                                                          105
                                                                   43
                                                                            82
                                                                                    16
                                 309
                                         234
                                                          300
                                                                           270
                                                                                    560
## 4 27395
                   Mrpl15
                                                 337
                                                                  290
## 5 18777
                  Lypla1
                                652
                                         515
                                                 948
                                                          935
                                                                  928
                                                                           791
                                                                                   826
## # ... with 5 more variables: MCL1.LB <dbl>, MCL1.LC <dbl>, MCL1.LD <dbl>,
## #
      MCL1.LE <dbl>, MCL1.LF <dbl>
```

```
dim(exprDat)
```

```
## [1] 23151 14
```

head(exprInfo)

```
## # A tibble: 6 x 5
##
    SampleName CellType Status
                                 Status.Type
                                                CellType.colors
                                 <chr>
                                                <chr>
##
    <chr>
           <chr> <chr>
## 1 MCL1.DG
               basal
                        control control.basal #79ADDC
## 2 MCL1.DH
             basal control control.basal #79ADDC
## 3 MCL1.DI basal pregnant pregnant.basal #79ADDC
## 4 MCL1.DJ
             basal pregnant pregnant.basal #79ADDC
                       lactate lactate.basal #79ADDC
               basal
## 5 MCL1.DK
## 6 MCL1.DL
               basal
                        lactate lactate.basal #79ADDC
Convert character columns to factor types:
exprInfo <- exprInfo %>%
 mutate(CellType = as.factor(CellType),
        Status = factor(Status, levels = c("control", "pregnant", "lactate")),
        Status.Type = as.factor(Status.Type))
head(exprInfo)
## # A tibble: 6 x 5
    SampleName CellType Status
                                 Status.Type
                                                CellType.colors
    <chr>
              <fct>
##
                        <fct>
                                 <fct>
                                                <chr>>
               basal
## 1 MCL1.DG
                        control control.basal #79ADDC
## 2 MCL1.DH basal control control.basal #79ADDC
## 3 MCL1.DI basal pregnant pregnant.basal #79ADDC
             basal pregnant pregnant.basal #79ADDC basal lactate lactate.basal #79ADDC
## 4 MCL1.DJ
## 5 MCL1.DK
## 6 MCL1.DL
               basal lactate lactate.basal #79ADDC
```

Initial Data Check & Filtering:

Let's try to sample 12 (n) random genes and plot their count distribution.

```
expr12 <- exprDat %>%
  dplyr::select(-EntrezGeneID, -GeneName) %>%
  sample_n(.,12) %>%
  t() %>%
  as_tibble() %>%
  rename_at(vars(names(.)), ~pasteO("Gene", seq(1:12))) %>%
  gather() %>%
  mutate(valuelog2 = log2(value+1))

# Give it a look:
expr12

## # A tibble: 144 x 3
```

```
##
         value valuelog2
     key
     <chr> <dbl>
                    <dbl>
##
## 1 Gene1 1405
                    10.5
## 2 Gene1 1354
                   10.4
## 3 Gene1 1845
                   10.9
## 4 Gene1 1459
                    10.5
## 5 Gene1 1078
                   10.1
```

```
6 Gene1
                         10.2
##
              1141
##
    7 Gene1
               814
                          9.67
                775
                          9.60
##
    8 Gene1
##
    9 Gene1
               874
                          9.77
## 10 Gene1
                749
                          9.55
## # ... with 134 more rows
Plot:
ggplot(expr12, aes(valuelog2)) +
  geom_histogram(color="black", fill="grey80", bins=20) +
  theme_minimal() +
  facet_wrap(~key)
               Gene1
                                      Gene<sub>10</sub>
                                                              Gene11
                                                                                     Gene12
  12.5
  10.0
   7.5
   5.0
    2.5
   0.0
                                       Gene3
                                                              Gene4
               Gene2
                                                                                      Gene5
  12.5
  10.0
   7.5
   5.0
    2.5
   0.0
                Gene6
                                       Gene7
                                                              Gene8
                                                                                      Gene9
  12.5
  10.0
   7.5
    5.0
    2.5
    0.0
                                                                              0
        0
              5
                    10
                          15
                                0
                                      5
                                           10
                                                 15
                                                       0
                                                             5
                                                                   10
                                                                         15
                                                                                    5
                                                                                          10
                                                                                                15
                                                valuelog2
```

We will filter out genes with too many zero counts. Exclude the columns with gene information:

```
# Count number of Os across samples. Filter samples where at least four samples has a count great than
exprDat <- exprDat %>%
  mutate(nzeros = rowSums(dplyr::select(.,-EntrezGeneID, -GeneName)==0)) %>%
  filter(nzeros <= 8) %>%
  dplyr::select(-nzeros)

#How many genes do we have left:
dim(exprDat)
```

Differential Expression Analysis- DESeq2

We will now make a DESeq2 object. For this we use the function DESeqDataSetFromMatrix from the DEseq2 package. As input we give our count matrix, our gene IDs and our meta data (exprInfo). Additionally we include a design for DE contrasts. In this case we add CellType (luminal or basal) and Status (control, pregnant or lactating).

Convert to exprDat to a dataframe and make GeneNames column into rownames:

```
# Pull out GeneNames and EntrezGeneID for later use
GeneNames <- exprDat %>%
  dplyr::select(EntrezGeneID, GeneName)
exprDat <- exprDat %>%
  dplyr::select(-EntrezGeneID) %>%
  column_to_rownames(., var = "GeneName")
Make a DESeq2 object:
exprObj <- DESeqDataSetFromMatrix(countData = exprDat,</pre>
                              colData = exprInfo,
                              design= ~CellType+Status)
expr0bj
## class: DESeqDataSet
## dim: 17308 12
## metadata(1): version
## assays(1): counts
## rownames(17308): Xkr4 Rp1 ... Uty Gm47283,
## rowData names(0):
## colnames(12): MCL1.DG MCL1.DH ... MCL1.LE MCL1.LF
## colData names(5): SampleName CellType Status Status.Type
     CellType.colors
##
```

Next, we use DEseq() to estimate dispersion, gene-wise and mean-dispersion, fitting model(s):

```
expr0bj <- DESeq(expr0bj)</pre>
```

Preliminary analysis:

Let's have a look at the library sizes:

21675050 21457888 24419457 24366629

```
colSums(assay(expr0bj))

## MCL1.DG MCL1.DH MCL1.DI MCL1.DJ MCL1.DK MCL1.DL MCL1.LA MCL1.LB

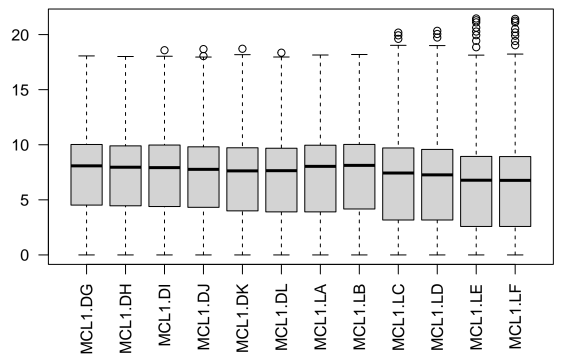
## 22634514 21155013 23488082 22100122 21057113 19583106 19698631 20944796

## MCL1.LC MCL1.LD MCL1.LE MCL1.LF
```

The count distributions may be dominated by a few genes with very large counts. These genes will drive plotting e.g. heatmaps, PCA analysis etc. Let's see if we have any genes with high large counts and in turn,

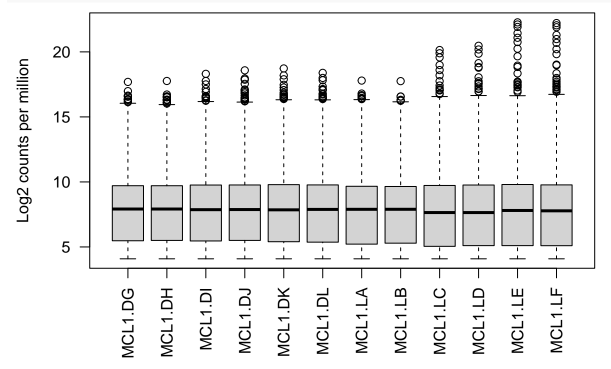
dispersion in our dataset. For convenience I am using the base R boxplot function:

```
#boxplot(assay(exprObj), las=2)
boxplot(log2(assay(exprObj)+1), las=2)
```



We perform variance stabilizing transformation to obtain log2 counts per million read mapped, overcoming issues with outlier genes and sequencing depth:

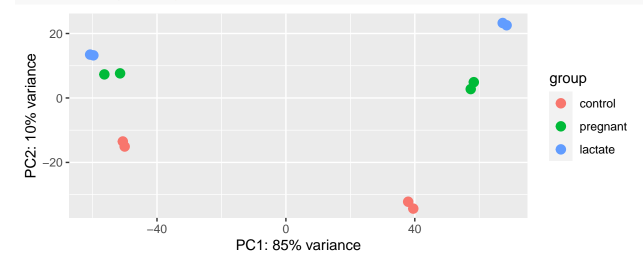
```
expr0bjvst <- vst(expr0bj,blind=FALSE)
boxplot(assay(expr0bjvst), xlab="", ylab="Log2 counts per million",las=2)</pre>
```



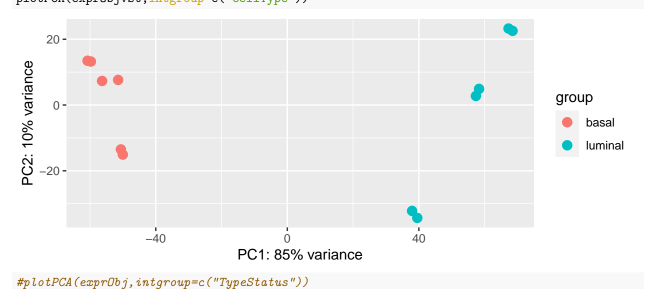
Principal Component Analysis

Before performing DEA it is a good idea to explore how samples cluster together based on there gene expression profile. The expectation here is that samples from the same group (treatment vs control, condition A vs condition B, etc.) will cluster together. A principal component analysis (PCA) plot can also help us to identify outlier samples which might need to be removed from the analysis. We use our vst counts for principal component analysis:





plotPCA(exprObjvst,intgroup=c("CellType"))



Testing

Have a look at the group comparisons:

resultsNames(expr0bj)

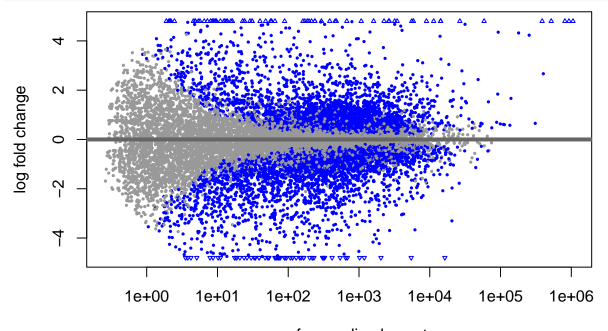
Test for DE genes between the three groups of mice, adjusted for cell type:

(I) lactating and control mice:

```
resLC <- results(expr0bj, contrast = c("Status", "lactate", "control"), independentFiltering = FALSE)</pre>
```

Summary and plot of DE analysis results:

DESeq2::plotMA(resLC)



mean of normalized counts

summary(resLC)

```
##
## out of 17308 with nonzero total read count
## adjusted p-value < 0.1
## LFC > 0 (up) : 3550, 21%
## LFC < 0 (down) : 3474, 20%
## outliers [1] : 0, 0%
## low counts [2] : 0, 0%
## (mean count < 0)
## [1] see 'cooksCutoff' argument of ?results
## [2] see 'independentFiltering' argument of ?results</pre>
```

Custom function to filter results of DEA. We make a function to save writing the same code three times in a row, one time for each comparison:

```
# SIGNIFICANT DE GENES:
# Takes as arguments:
```

```
# my.res = a dataframe of results from the DEseq results()function
# my.LFC = log fold change cutoff, default is 1.0
# my.cof = adjusted p-value cutoff, default is 0.01
SigDE <- function(my.res) {
   my.res <- as.data.frame(my.res) %>%
      rownames_to_column(., var = "GeneName") %>%
      as_tibble() %>%
      mutate(dir = ifelse(log2FoldChange >= 0, 'up', 'down')) %>%
      filter((log2FoldChange >= 1.0 | log2FoldChange <= -1.0) & padj <= 0.01) %>%
      arrange(padj, desc(abs(log2FoldChange)))
   return(my.res)
}
```

Filter DEA results from comparison of lactating vs control mice using custom function:

```
resLC <- SigDE(resLC)</pre>
# Number of DE genes:
dim(resLC)
## [1] 2126
# Give it a look
head(resLC, n=5)
## # A tibble: 5 x 8
##
    GeneName baseMean log2FoldChange lfcSE stat
                                                     pvalue
                                                                 padj dir
     <chr>
                <dbl>
                                                      <dbl>
##
                                <dbl> <dbl> <dbl>
                                                                <dbl> <chr>
                                9.76 0.424 23.0 3.46e-117 5.98e-113 up
## 1 Wap
              387497.
                                8.24 0.414 19.9 6.35e- 88 5.49e- 84 up
## 2 Csn1s2a 1056557.
## 3 Csn1s2b
               26616.
                              10.4 0.580 18.0 2.69e- 72 1.55e- 68 up
## 4 Glycam1
              520318.
                               9.62 0.539 17.9 2.52e- 71 1.09e- 67 up
## 5 Pigr
                                 6.40 0.373 17.2 4.17e- 66 1.44e- 62 up
                11356.
```

Below we perform the same steps as above to get the DE genes between (II) pregnant and control mice and (III) lactating and pregnant mice:

(II) pregnant and control mice:

```
resPC <- results(expr0bj, contrast = c("Status", "pregnant", "control"), independentFiltering = FALSE)
#DESeq2::plotMA(resPC)
#summary(resPC)
resPC <- SigDE(resPC)

# Number of DE genes:
dim(resPC)

## [1] 1206 8

(III) lactating and pregnant mice:
resLP <- results(expr0bj, contrast = c("Status", "lactate", "pregnant"), independentFiltering = FALSE)
#DESeq2::plotMA(resLP)
#summary(resLP)
resLP <- SigDE(resLP)</pre>
```

```
## [1] 790 8
```

Heatmap Visualization

To visually inspect if DE genes identified in our DESeq2 analysis successfully separate the three groups of mice (control, pregnant and lactating), we will make a heatmap. For this we use the heatmap function and package viridis.

It will not make sense to include all DE genes in this heatmap (3000 genes). Instead pick the top 50 most significant DE genes, based on adj. p-value and logFC.

Make a vector of unique EntrezGeneIDs (top50):

```
topDE <- bind_rows(resPC[1:50,], resLC[1:50,], resLP[1:50,]) %>%
  pull(GeneName) %>%
  unique()

length(topDE)
```

[1] 124

The expression counts themselves (not logFC) are needed for the heatmap. We use the topDE vector to extract these from the vst normalized DESeq2 object.

```
head(assay(expr0bjvst), n=5)
```

```
##
          MCL1.DG MCL1.DH MCL1.DI MCL1.DJ MCL1.DK MCL1.DL MCL1.LA MCL1.LB
## Xkr4
         8.536754 8.157576 6.401025 7.987204 8.608154 8.350271 4.088346 4.088346
          4.395113 4.408625 4.088346 4.088346 4.088346 4.088346 5.049024 4.605457
## Sox17 6.823999 7.536586 6.631003 7.018342 6.198459 6.849080 5.290970 5.528569
## Mrpl15 8.084835 7.843863 8.288204 8.288468 8.346705 8.271139 8.881522 8.550341
## Lypla1 9.068321 8.867045 9.679391 9.821810 9.921005 9.717343 9.409346 9.384315
##
          MCL1.LC MCL1.LD MCL1.LE
                                      MCL1.LF
## Xkr4
          4.088346 4.088346 4.088346
                                     4.088346
## Rp1
          5.155437 4.603001 4.088346 4.088346
## Sox17 5.496313 5.102135 4.880034 5.487699
## Mrpl15 8.993366 8.614985 9.157331 9.288091
## Lypla1 9.418084 9.531917 9.945641 10.020675
resVST <- assay(expr0bjvst) %>%
  as.data.frame() %>%
  rownames_to_column(var = "GeneName")
  as_tibble() %>%
  filter(GeneName %in% topDE)
```

The heatmap function in base R wants gene expression data as a matrix (a dataframe with numeric values only). We extract the GeneNames column and convert the tibble into a matrix:

```
HPnames <- resVST %>%
  pull(GeneName)

HPdat <- resVST %>%
  dplyr::select(-GeneName) %>%
  as.matrix()
```

We use the heatmap function to generate a heatmap. We can modify the look of the heatmap as desired, e.g. add column colors, row labels, change color scheme etc.

