

Lab 13

Youn Soo Na (PID: A17014731)

```
# library(BiocManager)
# library(DESeq2)
```

Today we will analyze some RNASeq data from Himes et al. on the effects of dexamethasone (dex), a synthetic glucocorticoid steroid on airway smooth muscle cells (ASM).

Data Import

```
counts <- read.csv("airway_scaledcounts.csv", row.names = 1)
metadata <- read.csv("airway_metadata.csv")
```

Q1. How many genes are in this dataset?

```
nrow(counts)
```

[1] 38694

Q2. How many ‘control’ cell lines do we have?

```
table(metadata$dex)
```

```
control treated
        4       4
```

There are 4 control cell lines.

Toy differential expression analysis

Calculate the mean per gene count values for all “control” samples (i.e. columns in `counts`) and do the same for “treated” and then compare them.

1. Find all “control” values/columns in `counts`

```
library(dplyr)
```

Attaching package: 'dplyr'

The following objects are masked from 'package:stats':

filter, lag

The following objects are masked from 'package:base':

intersect, setdiff, setequal, union

```
control <- metadata %>% filter(dex=="control")
control.counts <- counts %>% select(control$id)
```

2. Find the mean per gene across all control columns

```
control.mean <- apply(control.counts, 1, mean)
```

Q3. How would you make the above code in either approach more robust? Is there a function that could help here?

```
# control.mean <- rowSums( control.counts )/4
```

3. Repeat for treated

Q4. Follow the same procedure for the treated samples (i.e. calculate the mean per gene across drug treated samples and assign to a labeled vector called `treated.mean`)

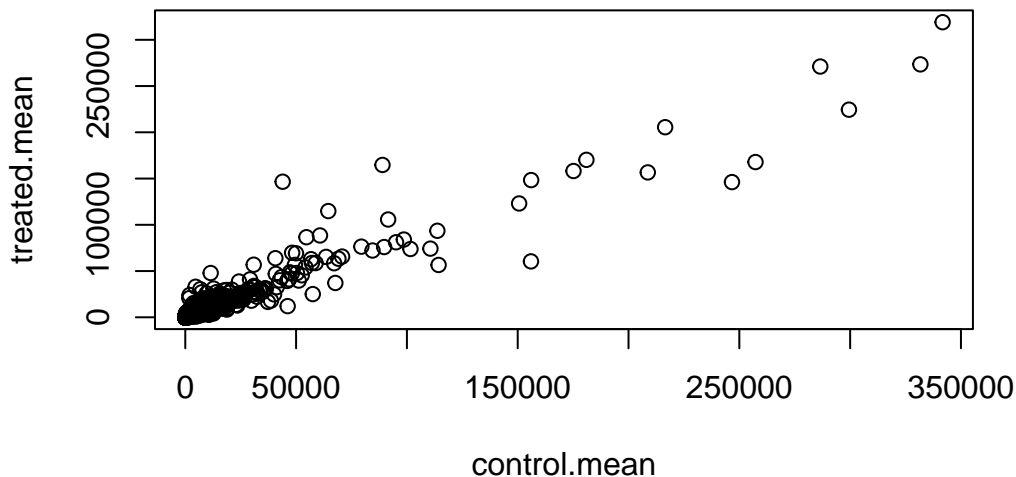
```
treated <- metadata$dex == "treated"
treated.counts <- counts[,treated]
```

```
treated.mean <- apply(treated.counts, 1, mean)
```

```
meancounts <- data.frame(control.mean, treated.mean)
```

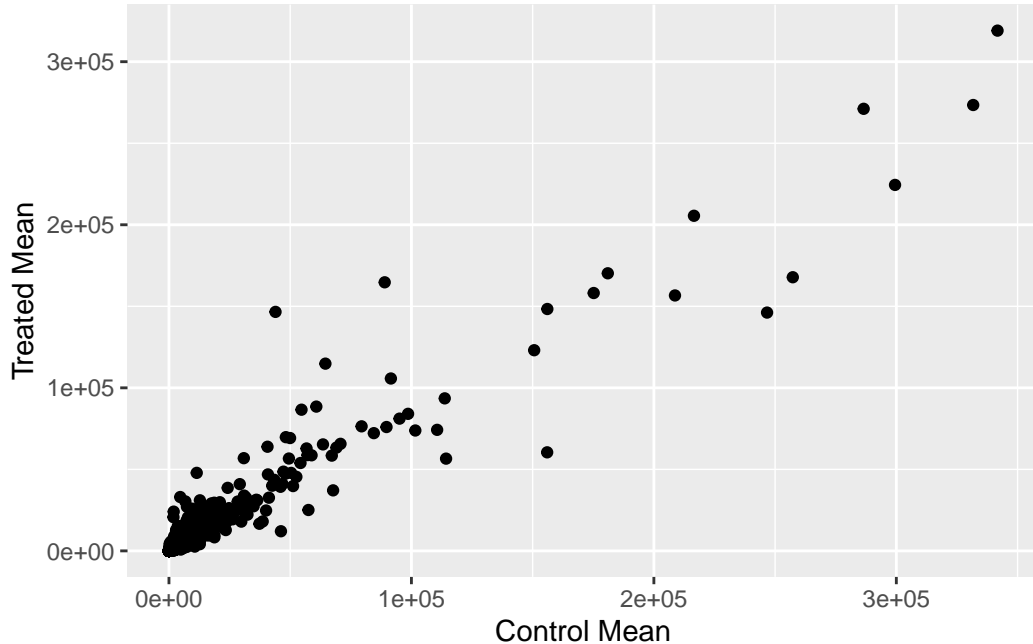
Q5 (a). Create a scatter plot showing the mean of the treated samples against the mean of the control samples. Your plot should look something like the following.

```
plot(meancounts)
```



Q5 (b). You could also use the ggplot2 package to make this figure producing the plot below. What geom_?() function would you use for this plot?

```
library(ggplot2)
ggplot(meancounts, aes(control.mean, treated.mean)) +
  geom_point() +
  labs(x="Control Mean", y="Treated Mean")
```

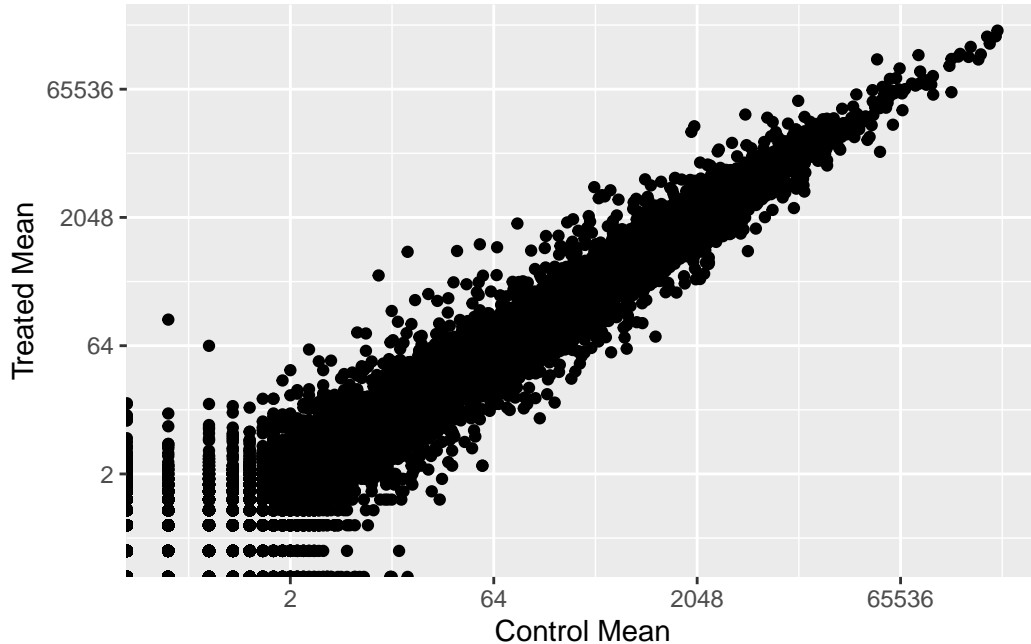


Q6. Try plotting both axes on a log scale. What is the argument to `plot()` that allows you to do this?

```
# plot(meancounts, log="xy")
# or
ggplot(meancounts, aes(control.mean, treated.mean)) +
  geom_point() +
  scale_x_continuous(trans = "log2") +
  scale_y_continuous( trans = "log2" ) +
  labs(x = "Control Mean", y = "Treated Mean")
```

Warning in `scale_x_continuous(trans = "log2")`: log-2 transformation introduced infinite values.

Warning in `scale_y_continuous(trans = "log2")`: log-2 transformation introduced infinite values.



We most frequently use log2 transformations for this type of data.

```
log2(20/10)
```

```
[1] 1
```

These log2 values make the interpretation of “fold-change” a little easier and a rule-of-thumb in the filled is a log2 fold-change of +2 or a -2 is where we start to pay attention.

```
log2(40/10)
```

```
[1] 2
```

Let's calculate the log2(fold-change) and add it to our `meancounts` data.frame.

```
meancounts$log2fc <- log2(meancounts$treated.mean/meancounts$control.mean)
head(meancounts)
```

	control.mean	treated.mean	log2fc
ENSG000000000003	900.75	658.00	-0.45303916
ENSG000000000005	0.00	0.00	NaN
ENSG000000000419	520.50	546.00	0.06900279
ENSG000000000457	339.75	316.50	-0.10226805
ENSG000000000460	97.25	78.75	-0.30441833
ENSG000000000938	0.75	0.00	-Inf

```
to.rm <- rowSums(meancounts[,1:2]==0) > 0
mycounts <- meancounts[!to.rm,]
```

Q7. What is the purpose of the arr.ind argument in the which() function call above? Why would we then take the first column of the output and need to call the unique() function?

Filtering data to remove genes with zero expression

```
zero.vals <- which(meancounts[,1:2]==0, arr.ind=TRUE)

to.rm <- unique(zero.vals[,1])
mycounts <- meancounts[-to.rm,]
head(mycounts)
```

	control.mean	treated.mean	log2fc
ENSG000000000003	900.75	658.00	-0.45303916
ENSG000000000419	520.50	546.00	0.06900279
ENSG000000000457	339.75	316.50	-0.10226805
ENSG000000000460	97.25	78.75	-0.30441833
ENSG000000000971	5219.00	6687.50	0.35769358
ENSG00000001036	2327.00	1785.75	-0.38194109

`arr.ind` returns a matrix of indices that correspond to the non-zero elements (taking out the zero counts). `unique()` prevents accidental repeats when counting the zero entries.

Q. How many genes do I have left after this zero count filtering?

```
nrow(mycounts)
```

[1] 21817

Q. How many genes are “up” regulated upon drug treatment at a threshold of +2 log2-fold-change? Q8. Using the up.ind vector above can you determine how many up regulated genes we have at the greater than 2 fc level?

1. I need to extract the log2fc values
2. I need to find those that are above +2
3. Count them.

```
sum(mycounts$log2fc > 2)
```

[1] 250

Q. How many genes are “down” regulated upon drug treatment at a threshold of -2 log2-fold-change? Q9. Using the down.ind vector above can you determine how many down regulated genes we have at the greater than 2 fc level?

```
sum(mycounts$log2fc < -2)
```

[1] 367

Q10. Do you trust these results? Why or why not?

No, because the statistics side of this data is missing.

Missing the stats. Is the difference in the mean counts significant?

Let’s do this analysis the right way with statistics and use the **DESeq2** package

DESeq2 analysis

```
library(DESeq2)
```

The first function that we will use will setup the data in the way (format) DESeq wants it.

```
dds <- DESeqDataSetFromMatrix(countData = counts,
                                colData = metadata,
                                design = ~dex)
```

converting counts to integer mode

```
Warning in DESeqDataSet(se, design = design, ignoreRank): some variables in  
design formula are characters, converting to factors
```

The function in the package is called `DESeq()` and we can run it on our `dds` object.

```
dds <- DESeq(dds)
```

```
estimating size factors
```

```
estimating dispersions
```

```
gene-wise dispersion estimates
```

```
mean-dispersion relationship
```

```
final dispersion estimates
```

```
fitting model and testing
```

I will get the results from `dds` with the `results()` function:

```
res <- results(dds)  
head(res)
```

```
log2 fold change (MLE): dex treated vs control  
Wald test p-value: dex treated vs control  
DataFrame with 6 rows and 6 columns  
  baseMean log2FoldChange      lfcSE      stat     pvalue  
  <numeric>      <numeric> <numeric> <numeric> <numeric>  
ENSG000000000003 747.194195    -0.3507030  0.168246 -2.084470  0.0371175  
ENSG000000000005  0.000000      NA        NA        NA        NA  
ENSG00000000419   520.134160    0.2061078  0.101059  2.039475  0.0414026  
ENSG00000000457   322.664844    0.0245269  0.145145  0.168982  0.8658106  
ENSG00000000460   87.682625    -0.1471420  0.257007 -0.572521  0.5669691  
ENSG00000000938   0.319167    -1.7322890  3.493601 -0.495846  0.6200029  
  padj  
  <numeric>  
ENSG000000000003  0.163035  
ENSG000000000005  NA
```

```

ENSG000000000419  0.176032
ENSG000000000457  0.961694
ENSG000000000460  0.815849
ENSG000000000938      NA

```

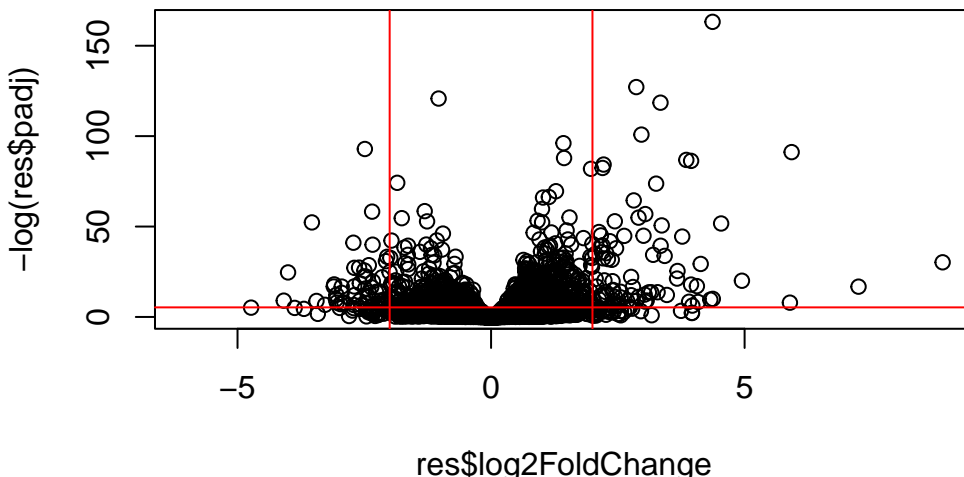
Make a common overall results figure from this analysis. This is designed to keep our inner biologist and inner stats nerd happy because it plots fold-change vs P-value.

padj: p-value that has been corrected over multiple trials; stricter

```

# log is used to better the visualization of the p-value (more negative
# corresponds to smaller p-value)
# - is used to flip the plot (now more positive values correspond to
# lower p-value)
plot(res$log2FoldChange, -log(res$padj))
abline(v = c(-2,2), col = "red")
abline(h = -log(0.005), col = "red")

```



Add some color to this plot

```

mycols <- rep("gray", nrow(res))
mycols[res$log2FoldChange > 2 | res$log2FoldChange < -2] <- "blue"
mycols[res$padj > 0.005 & res$log2FoldChange > 2] <- "red"

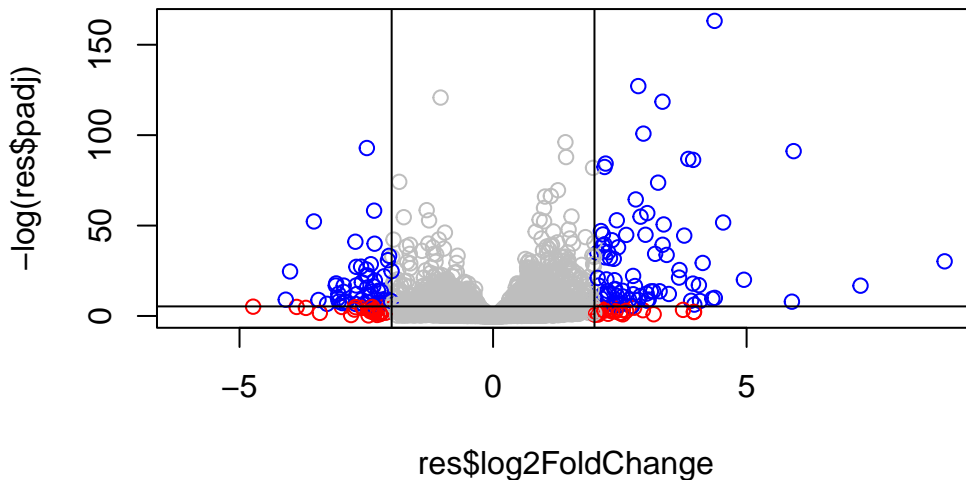
```

```

mycols[res$padj > 0.005 & res$log2FoldChange < -2] <- "red"

plot(res$log2FoldChange, -log(res$padj), col = mycols)
abline(v = c(-2,2), col = "black")
abline(h = -log(0.005), col = "black")

```



I want to save my results to date out to disc

```
write.csv(res, file = "myresults.csv")
```

We will pick this up next day and add **annotation** (i.e. what are these genes of interest) and do pathway analysis (what biology) are they known to be involved with.

I need to translate our gene identifiers “ENSG000...” into gene names that the rest of the world can understand.

To do this “annotation,” I will use the **AnnotationDbi** package. I can install this with **BiocManager::install()**

```
library(AnnotationDbi)
library(org.Hs.eg.db)

columns(org.Hs.eg.db)
```

```
[1] "ACNUM"          "ALIAS"           "ENSEMBL"         "ENSEMLPROT"      "ENSEMLTRANS"
[6] "ENTREZID"       "ENZYME"          "EVIDENCE"        "EVIDENCEALL"    "GENENAME"
[11] "GENETYPE"       "GO"               "GOALL"           "IPI"             "MAP"
[16] "OMIM"           "ONTOLOGY"        "ONTOLOGYALL"    "PATH"            "PFAM"
[21] "PMID"           "PROSITE"          "REFSEQ"          "SYMBOL"          "UCSCKG"
[26] "UNIPROT"
```

I will use the `mapIds()` function to “map” my identifiers to those from different databases. I will go between “ENSEMBL” and “SYMBOL” (and then after “GENENAME”).

```
res$symbol <- mapIds(org.Hs.eg.db,
                      keys = rownames(res),
                      keytype = "ENSEMBL",
                      column = "SYMBOL")
```

```
'select()' returned 1:many mapping between keys and columns
```

```
# head(res)
```

Q11. Run the `mapIds()` function two more times to add the Entrez ID and UniProt accession and GENENAME as new columns called `res$entrez`, `res$uniprot`, and `res$genename`.

Add “GENENAME”

```
res$genename <- mapIds(org.Hs.eg.db,
                       keys = rownames(res),
                       keytype = "ENSEMBL",
                       column = "GENENAME")
```

```
'select()' returned 1:many mapping between keys and columns
```

```
head(res$genename)
```

```

ENSG000000000003
"tetraspanin 6"
ENSG000000000005
"tenomodulin"
ENSG000000000419
"dolichyl-phosphate mannosyltransferase subunit 1, catalytic"
ENSG000000000457
"SCY1 like pseudokinase 3"
ENSG000000000460
"FIGNL1 interacting regulator of recombination and mitosis"
ENSG000000000938
"FGR proto-oncogene, Src family tyrosine kinase"

```

And “ENTREZID”

```

res$entrez <- mapIds(org.Hs.eg.db,
                      keys = rownames(res),
                      keytype = "ENSEMBL",
                      column = "ENTREZID")

```

```
'select()' returned 1:many mapping between keys and columns
```

```
# head(res$entrez)
# res
```

```

res$uniprot <- mapIds(org.Hs.eg.db,
                      keys = rownames(res),
                      keytype = "ENSEMBL",
                      column = "UNIPROT")

```

```
'select()' returned 1:many mapping between keys and columns
```

```
# head(res$uniprot)
```

save our annotated results object.

```
write.csv(res, file = "results_annotated.csv")
```

Pathway Analysis

Now that we have our results with added annotations we can do some pathway mapping.

Let's use the **gage** package to look for KEGG pathways in our results (genes of interest). I will also use the **pathview** package to draw little pathway figures.

```
library(pathview)
library(gage)
library(gageData)

data(kegg.sets.hs)
head(kegg.sets.hs, 1)
```

```
$`hsa00232 Caffeine metabolism`
[1] "10"   "1544" "1548" "1549" "1553" "7498" "9"
```

What **gage** wants as inputs is not my big table/data.frame of results. It just wants a “vector of importance”. For RNASeq data like we have this is our log2FC values...

```
foldchanges = res$log2FoldChange
names(foldchanges) = res$entrez
head(foldchanges)
```

```
7105      64102      8813      57147      55732      2268
-0.35070302      NA  0.20610777  0.02452695 -0.14714205 -1.73228897
```

Now, let's run the gage pathway analysis.

```
# Get the results
keggres = gage(foldchanges, gsets=kegg.sets.hs)
# What's in this keggres object?
attributes(keggres)
```

```
$names
[1] "greater" "less"    "stats"

head(keggres$less, 3)
```

	p.geomean	stat.mean	p.val
hsa05332 Graft-versus-host disease	0.0004250461	-3.473346	0.0004250461
hsa04940 Type I diabetes mellitus	0.0017820293	-3.002352	0.0017820293
hsa05310 Asthma	0.0020045888	-3.009050	0.0020045888
	q.val	set.size	exp1
hsa05332 Graft-versus-host disease	0.09053483	40	0.0004250461
hsa04940 Type I diabetes mellitus	0.14232581	42	0.0017820293
hsa05310 Asthma	0.14232581	29	0.0020045888

Let's use the **pathview** package to look at one of these highlighted KEGG pathways with our genes highlighted. "hsa05310 Asthma"

```
pathview(gene.data=foldchanges, pathway.id="hsa05310")
```

```
'select()' returned 1:1 mapping between keys and columns
```

```
Info: Working in directory /Users/nayounsoo/Desktop/BIMM 143/Lab 13/Lab 13
```

```
Info: Writing image file hsa05310.pathview.png
```

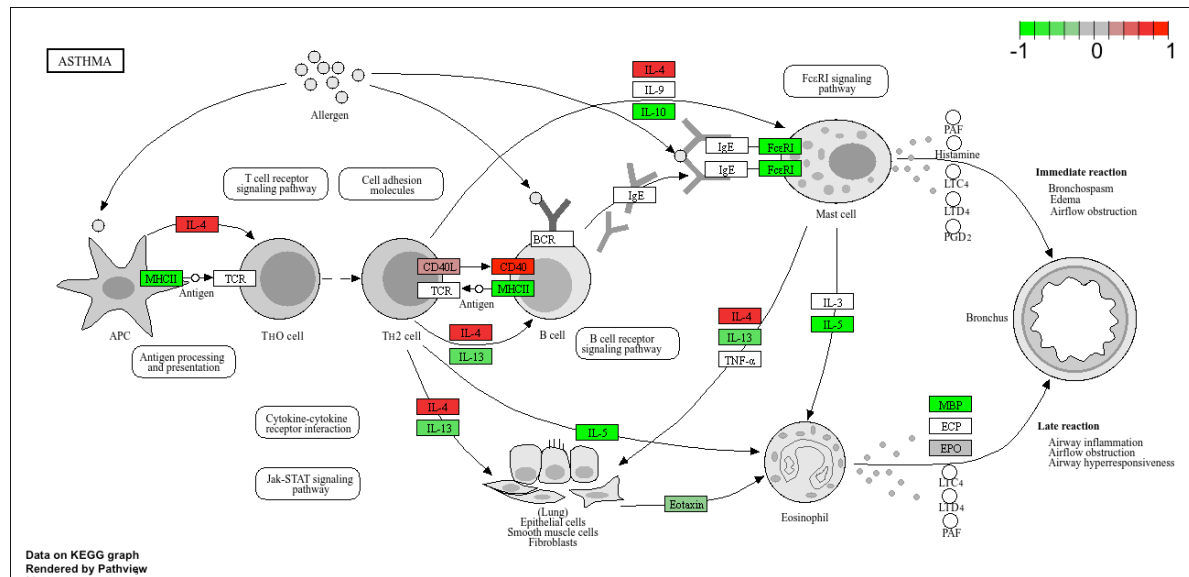


Figure 1: Asthma pathway with my DEGs