Class 13: RNASeq with DESeq2

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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")</pre>
```

A wee peak

head(counts)

	SRR1039508	SRR1039509	SRR1039512	SRR1039513	SRR1039516
ENSG0000000003	723	486	904	445	1170
ENSG0000000005	0	0	0	0	0
ENSG00000000419	467	523	616	371	582
ENSG00000000457	347	258	364	237	318
ENSG00000000460	96	81	73	66	118
ENSG00000000938	0	0	1	0	2
	SRR1039517	SRR1039520	SRR1039521		
ENSG0000000003	1097	806	604		
ENSG0000000005	0	0	0		
ENSG00000000419	781	417	509		
ENSG00000000457	447	330	324		
ENSG00000000460	94	102	74		
ENSG00000000938	0	0	0		

head(metadata)

```
id dex celltype geo_id
1 SRR1039508 control N61311 GSM1275862
2 SRR1039509 treated N61311 GSM1275863
3 SRR1039512 control N052611 GSM1275866
4 SRR1039513 treated N052611 GSM1275867
5 SRR1039516 control N080611 GSM1275870
6 SRR1039517 treated N080611 GSM1275871
```

Q1. How many genes are in this dataset?

```
nrow(counts)
```

[1] 38694

Q2. How many 'control' cell lines do we have?

```
sum(metadata$dex == "control")
```

[1] 4

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

```
control treated 4 4
```

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

```
control.inds <- metadata$dex == "control"
control.counts <- counts[,control.inds]
head(control.counts)</pre>
```

	SRR1039508	SRR1039512	SRR1039516	SRR1039520
ENSG00000000003	723	904	1170	806
ENSG0000000005	0	0	0	0
ENSG00000000419	467	616	582	417
ENSG00000000457	347	364	318	330
ENSG00000000460	96	73	118	102
ENSG00000000938	0	1	2	0

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

To make the code more robust, you would use the apply function to apply the mean to the column above.

2. Find the mean per gene across all control columns.

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

- 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)
- 3. Do the same steps to find the treated.mean values.

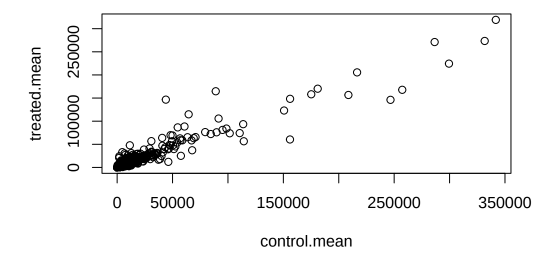
```
treated.inds <- metadata$dex == "treated"
treated.counts <- counts[,treated.inds]
head(treated.counts)</pre>
```

	SRR1039509	SRR1039513	SRR1039517	SRR1039521
ENSG0000000003	486	445	1097	604
ENSG0000000005	0	0	0	0
ENSG00000000419	523	371	781	509
ENSG00000000457	258	237	447	324
ENSG00000000460	81	66	94	74
ENSG00000000938	0	0	0	0

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

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

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.



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?

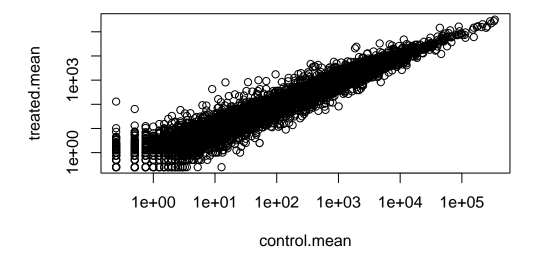
geom_(point)

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")

Warning in xy.coords(x, y, xlabel, ylabel, log): 15032 x values <= 0 omitted from logarithmic plot

Warning in xy.coords(x, y, xlabel, ylabel, log): 15281 y values <= 0 omitted from logarithmic plot



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

log2(10/10)

[1] 0

log2(20/10)

[1] 1

log2(10/20)

[1] -1

These log 2 values make the interpretation of "fold-change" a little easier and a rule-of-thumb in the filed is a log2 fold-change of +2 or -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.

```
control.mean treated.mean
                                                 log2fc
ENSG0000000003
                       900.75
                                    658.00 -0.45303916
ENSG0000000005
                         0.00
                                      0.00
                                                    NaN
ENSG00000000419
                      520.50
                                    546.00
                                            0.06900279
ENSG00000000457
                       339.75
                                    316.50 -0.10226805
ENSG00000000460
                       97.25
                                     78.75 -0.30441833
ENSG00000000938
                         0.75
                                      0.00
                                                   -Inf
```

```
# meancounts[,1:2] == 0
```

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

	${\tt control.mean}$	${\tt treated.mean}$	log2fc
ENSG0000000003	900.75	658.00	-0.45303916
ENSG00000000419	520.50	546.00	0.06900279
ENSG00000000457	339.75	316.50	-0.10226805
ENSG00000000460	97.25	78.75	-0.30441833
ENSG00000000971	5219.00	6687.50	0.35769358
ENSG0000001036	2327.00	1785.75	-0.38194109

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?

The purpose of the arr.ind rgument in the which() function call above is to return both the row and column indices (i.e. positions) where there are TRUE values. Genes (rows) and samples (columns) with 0 counts will be displayed. Any genes that have 0 counts in any sample will be ignored (focus on the row answer). Calling unique() means no row will be counted twice if it has zero entries in both samples.

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?

- 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?

```
sum(mycounts log 2fc < -2)
```

[1] 367

Q8. Using the up.ind vector above can you determine how many up regulated genes we have at the greater than 2 fc level?

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

[1] 250

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)</pre>
```

Γ1] 367

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

I do not trust these results because fold-chain can be large without being statistically significant as well as missing the stats of whether the differences we are seeing are significant. In order to obtain the stats, we need to use the DESeq2 package.

Wow hold on we are missing the stats here. Is the difference in the mean counts significant?

Let's do this analysis the right way with stats and use the **DESeq2** package

DESeq analysis

```
#/ message: false
library(DESeq2)
Loading required package: S4Vectors
Loading required package: stats4
Loading required package: BiocGenerics
Attaching package: 'BiocGenerics'
The following objects are masked from 'package:stats':
    IQR, mad, sd, var, xtabs
The following objects are masked from 'package:base':
    anyDuplicated, aperm, append, as.data.frame, basename, cbind,
    colnames, dirname, do.call, duplicated, eval, evalq, Filter, Find,
    get, grep, grepl, intersect, is.unsorted, lapply, Map, mapply,
    match, mget, order, paste, pmax, pmax.int, pmin, pmin.int,
    Position, rank, rbind, Reduce, rownames, sapply, saveRDS, setdiff,
    table, tapply, union, unique, unsplit, which.max, which.min
Attaching package: 'S4Vectors'
The following object is masked from 'package:utils':
    findMatches
The following objects are masked from 'package:base':
    expand.grid, I, unname
```

Loading required package: IRanges

Attaching package: 'IRanges'

The following object is masked from 'package:grDevices':

windows

Loading required package: GenomicRanges

Loading required package: GenomeInfoDb

Loading required package: SummarizedExperiment

Loading required package: MatrixGenerics

Loading required package: matrixStats

Warning: package 'matrixStats' was built under R version 4.4.2

Attaching package: 'MatrixGenerics'

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

colAlls, colAnyNAs, colAnys, colAvgsPerRowSet, colCollapse, colCounts, colCummaxs, colCummins, colCumprods, colCumsums, colDiffs, colIQRDiffs, colIQRs, colLogSumExps, colMadDiffs, colMads, colMaxs, colMeans2, colMedians, colMins, colOrderStats, colProds, colQuantiles, colRanges, colRanks, colSdDiffs, colSds, colSums2, colTabulates, colVarDiffs, colVars, colWeightedMads, colWeightedMeans, colWeightedMedians, colWeightedSds, colWeightedVars, rowAlls, rowAnyNAs, rowAnys, rowAvgsPerColSet, rowCollapse, rowCounts, rowCummaxs, rowCummins, rowCumprods, rowCumsums, rowDiffs, rowIQRDiffs, rowIQRs, rowLogSumExps, rowMadDiffs, rowMads, rowMaxs, rowMeans2, rowMedians, rowMins, rowOrderStats, rowProds, rowQuantiles, rowRanges, rowRanks, rowSdDiffs, rowSds, rowSums2, rowTabulates, rowVarDiffs, rowVars, rowWeightedMads, rowWeightedMeans, rowWeightedMedians, rowWeightedSds, rowWeightedVars

```
Loading required package: Biobase
Welcome to Bioconductor
    Vignettes contain introductory material; view with
    'browseVignettes()'. To cite Bioconductor, see
    'citation("Biobase")', and for packages 'citation("pkgname")'.
Attaching package: 'Biobase'
The following object is masked from 'package:MatrixGenerics':
    rowMedians
The following objects are masked from 'package:matrixStats':
    anyMissing, rowMedians
The first function that we will use will setup the data in the way (format) DESeq wants it.
dds <- DESeqDataSetFromMatrix(countData = counts,</pre>
                        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

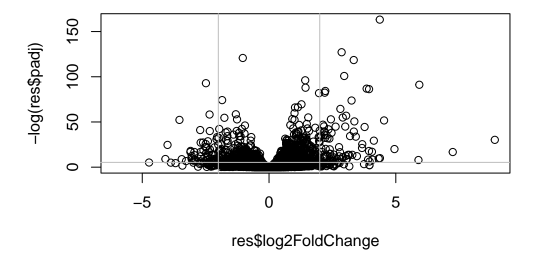
I will get the results from dds with the results() function:

```
res <- results(dds)
head(res)</pre>
```

```
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
                                                                 pvalue
                                                         stat
                 <numeric>
                                <numeric> <numeric> <numeric> <numeric>
ENSG00000000003 747.194195
                               -0.3507030 0.168246 -2.084470 0.0371175
ENSG00000000005
                  0.000000
                                                 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
                               -1.7322890 3.493601 -0.495846 0.6200029
                  0.319167
                     padj
                <numeric>
ENSG0000000000 0.163035
ENSG00000000005
ENSG00000000419 0.176032
ENSG00000000457
                 0.961694
ENSG00000000460
                0.815849
ENSG00000000938
                       NA
```

Make a common overall results figure from this analysis. This is designed to keep our inner biologist and inner stats nerd happy.

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



```
log(0.0005)
```

[1] -7.600902

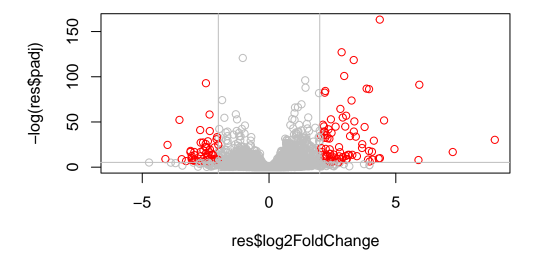
```
log(0.00000000005)
```

[1] -23.719

Add some color to this plot:

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

plot(res$log2FoldChange, -log(res$padj), col=mycols)
abline(v=c(-2,2), col="gray")
abline(h= -log(0.005), col="gray")</pre>
```



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.

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

pvalue	stat	lfcSE	log2FoldChange	baseMean	
<numeric></numeric>	<numeric></numeric>	<numeric></numeric>	<numeric></numeric>	<numeric></numeric>	
0.0371175	-2.084470	0.168246	-0.3507030	747.194195	ENSG0000000003
NA	NA	NA	NA	0.000000	ENSG0000000005
0.0414026	2.039475	0.101059	0.2061078	520.134160	ENSG00000000419
0.8658106	0.168982	0.145145	0.0245269	322.664844	ENSG00000000457
0.5669691	-0.572521	0.257007	-0.1471420	87.682625	ENSG00000000460
0.6200029	-0.495846	3.493601	-1.7322890	0.319167	ENSG00000000938
				padj	
				<numeric></numeric>	

```
ENSG00000000003 0.163035

ENSG00000000005 NA

ENSG00000000419 0.176032

ENSG00000000457 0.961694

ENSG00000000460 0.815849

ENSG000000000938 NA
```

Annotation

I need to translate our gene identifiers "ENSG0000..." into gene names that the rest of the world can understand.

To 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] "ACCNUM"
                    "ALIAS"
                                    "ENSEMBL"
                                                    "ENSEMBLPROT"
                                                                    "ENSEMBLTRANS"
 [6] "ENTREZID"
                    "ENZYME"
                                    "EVIDENCE"
                                                    "EVIDENCEALL"
                                                                   "GENENAME"
[11] "GENETYPE"
                    "GO"
                                    "GOALL"
                                                    "IPI"
                                                                   "MAP"
                                    "ONTOLOGYALL" "PATH"
[16] "OMIM"
                    "ONTOLOGY"
                                                                   "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").

^{&#}x27;select()' returned 1:many mapping between keys and columns

```
#head(res)
```

Add "GENENAME"

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

```
#head(res)
```

Add "ENTREZID"

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

```
#head(res)
```

Save our annotated results object.

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

Pathway Analysis

Now that we have our results with added annoatation 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.

```
#/ message: false
library(pathview)
```

Pathview is an open source software package distributed under GNU General Public License version 3 (GPLv3). Details of GPLv3 is available at http://www.gnu.org/licenses/gpl-3.0.html. Particullary, users are required to formally cite the original Pathview paper (not just mention it) in publications or products. For details, do citation("pathview") within R.

The pathview downloads and uses KEGG data. Non-academic uses may require a KEGG license agreement (details at http://www.kegg.jp/kegg/legal.html).

```
library(gage)
```

```
library(gageData)

data(kegg.sets.hs)

# Examine the first 2 pathways in this kegg set for humans head(kegg.sets.hs, 1)
```

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

What **gage** wants as input 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 is in this keggres object?

attributes(keggres)

\$names

[1] "greater" "less" "stats"

head(keggres\$less, 3)

```
p.val
                                      p.geomean stat.mean
hsa05332 Graft-versus-host disease 0.0004250461 -3.473346 0.0004250461
                                  0.0017820293 -3.002352 0.0017820293
hsa04940 Type I diabetes mellitus
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
                                                    29 0.0020045888
hsa05310 Asthma
                                   0.14232581
```

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 C:/Users/chloe/OneDrive/Desktop/BIMM143/class13

Info: Writing image file hsa05310.pathview.png

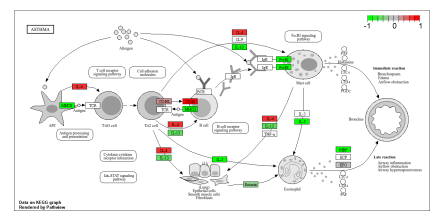


Figure 1: Asthma pathway with my DEGs