

Class 15 RNASeq Analysis

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#Background our data for today came from Himes et al. RNASeq analysis of the drug dexamethasone, a synthetic glucocorticoid steroid with anti-inflammatory effects (Himes et al. 2014).

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

```
head(counts)
```

```
##          SRR1039508 SRR1039509 SRR1039512 SRR1039513 SRR1039516
## ENSG000000000003      723        486       904       445      1170
## ENSG000000000005       0         0         0         0         0
## ENSG000000000419     467       523       616       371      582
## ENSG000000000457     347       258       364       237      318
## ENSG000000000460      96        81        73        66      118
## ENSG000000000938      0         0         1         0         2
##          SRR1039517 SRR1039520 SRR1039521
## ENSG000000000003    1097       806       604
## ENSG000000000005       0         0         0
## ENSG000000000419     781       417       509
## ENSG000000000457     447       330       324
## ENSG000000000460      94        102       74
## ENSG000000000938      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

control <- metadata[metadata[, "dex"]=="control",]
control.counts <- counts[, control$id]
control.mean <- rowSums( control.counts )/4
head(control.mean)

## ENSG0000000003 ENSG0000000005 ENSG00000000419 ENSG00000000457 ENSG00000000460
##         900.75          0.00        520.50        339.75        97.25
## ENSG0000000938
##         0.75

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)
control.mean <- rowSums(control.counts)/4
head(control.mean)

## ENSG0000000003 ENSG0000000005 ENSG00000000419 ENSG00000000457 ENSG00000000460
##         900.75          0.00        520.50        339.75        97.25
## ENSG0000000938
##         0.75

```

Q3. How would you make the above code in either approach more robust?

First i need to extract all the “control” columns. Then I will take the rowise mean to get the average count values for all genes in these four experiments.

```

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

##           SRR1039508 SRR1039512 SRR1039516 SRR1039520
## ENSG00000000003      723       904      1170       806
## ENSG00000000005        0         0         0         0
## ENSG00000000419      467       616      582       417
## ENSG00000000457      347       364      318       330
## ENSG00000000460       96        73      118       102
## ENSG00000000938        0         1         2         0

```

```
control.mean <- rowMeans(control.counts)
```

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

```
## SRR1039509 SRR1039513 SRR1039517 SRR1039521  
## ENSG00000000003 486 445 1097 604  
## ENSG00000000005 0 0 0 0  
## ENSG00000000419 523 371 781 509  
## ENSG00000000457 258 237 447 324  
## ENSG00000000460 81 66 94 74  
## ENSG00000000938 0 0 0 0
```

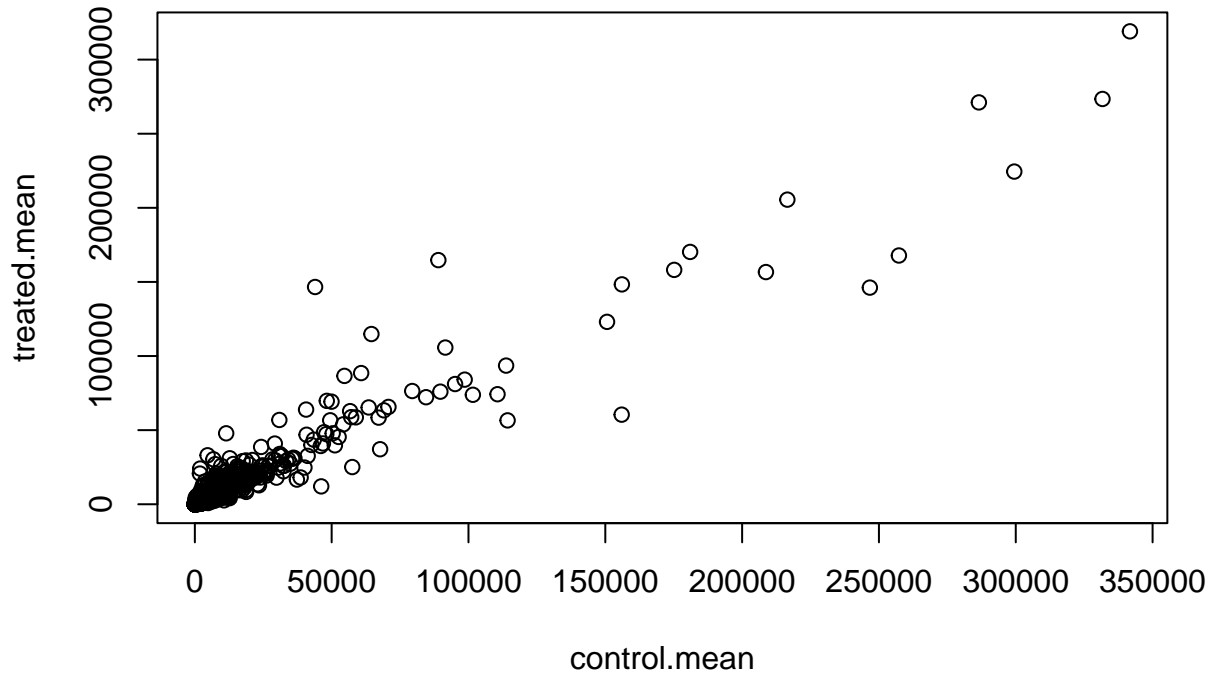
```
treated.mean <- rowMeans(treated.counts)
```

We will combine our meancount data for bookkeeping purposes.

```
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?

geom_point

This indicates that we need a log transformation to see details of our data!

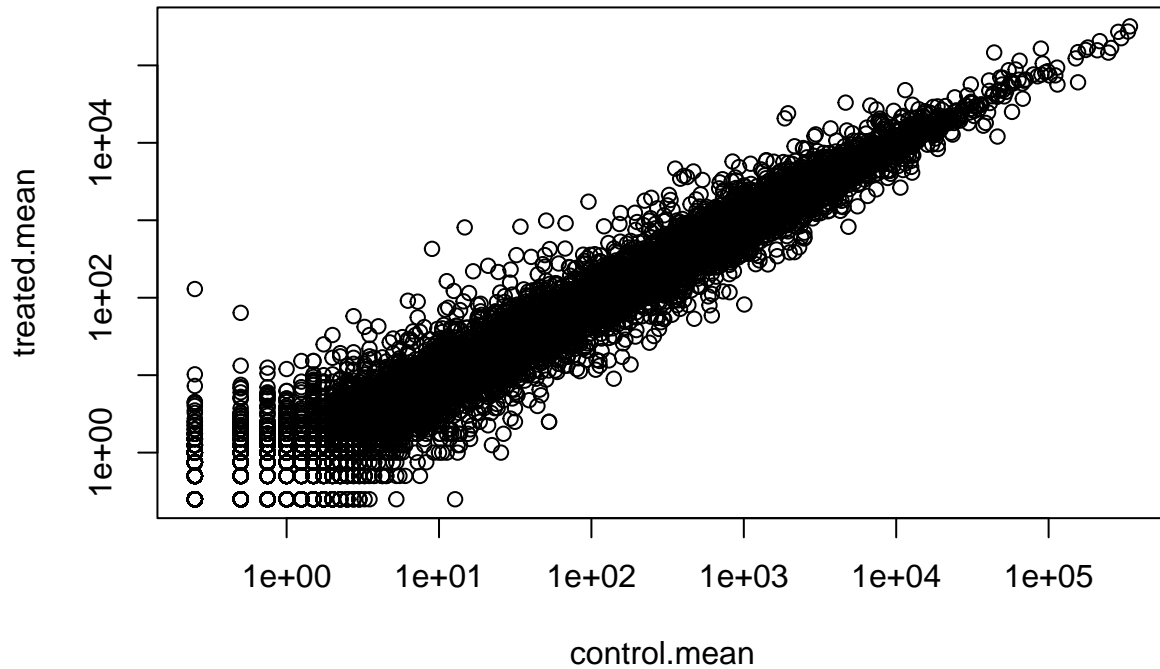
Q6. Try plotting both axes on a log scale. What is the argument to plot() that allows you to do this?

replot with Log-scale

```
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
```



often use `log2` in this field because it has nice math properties that make interpretation easier.

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

```
## [1] 0
```

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

```
## [1] 1
```

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

```
## [1] 2
```

```
log2(5/10)
```

```
## [1] -1
```

We see 0 values for no change and + values for increases and minus values for decreases. This nice property leads us to work with `log2(fold-change)` all the time in the genomics and proteomics field.

Let's add the `log2(fold-change)` values to our 'meancounts' dataframe.

```
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
## 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
```

I need to exclude the genes (i.e. rows) with zero counts because we can't say anything about these as we have no data for them!

```
which(c(F,F,T,T))
```

```
## [1] 3 4
```

Use the **which()** functions with the arr.ind=TRUE argument to get the columns and rows where the TRUE values are (i.e. the zero counts)

```
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
## 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
## ENSG00000001036   2327.00    1785.75 -0.38194109
```

```
nrow(mycounts)
```

```
## [1] 21817
```

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?

arr.ind is used to find the zero counts within the data so they can be ignored

```
up.ind <- mycounts$log2fc > 2
down.ind <- mycounts$log2fc < (-2)
```

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)
```

```
## [1] 367
```

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

I do not trust these results because fold can be large without being significant so the results can be misleading.

#DESeq Analysis

Let's do this the right way. DESeq2 is an R package specifically for analyzing count-based NGS data like RNA-seq. It is available from Bioconductor.

```
library(DESeq2)
```

```
## Loading required package: S4Vectors
```

```
## Loading required package: stats4
```

```
## Loading required package: BiocGenerics
```

```
##
```

```
## Attaching package: 'BiocGenerics'
```

```
## The following objects are masked from 'package:dplyr':
```

```
##
```

```
##     combine, intersect, setdiff, union
```

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

```
##
```

```
##     IQR, mad, sd, var, xtabs
```

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

```
##
```

```
##     anyDuplicated, 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, setdiff, sort, table, tapply,
##     union, unique, unsplit, which.max, which.min
```

```
##
```

```
## Attaching package: 'S4Vectors'
```

```

## The following objects are masked from 'package:dplyr':
##
##     first, rename

## The following objects are masked from 'package:base':
##
##     expand.grid, I, unname

## Loading required package: IRanges

##
## Attaching package: 'IRanges'

## The following objects are masked from 'package:dplyr':
##
##     collapse, desc, slice

## Loading required package: GenomicRanges

## Loading required package: GenomeInfoDb

## Loading required package: SummarizedExperiment

## Loading required package: MatrixGenerics

## Loading required package: matrixStats

##
## Attaching package: 'matrixStats'

## The following object is masked from 'package:dplyr':
##
##     count

##
## 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

citation("DESeq2")

##
## Love, M.I., Huber, W., Anders, S. Moderated estimation of fold change
## and dispersion for RNA-seq data with DESeq2 Genome Biology 15(12):550
## (2014)
##
## A BibTeX entry for LaTeX users is
##
## @Article{,
##   title = {Moderated estimation of fold change and dispersion for RNA-seq data with DESeq2},
##   author = {Michael I. Love and Wolfgang Huber and Simon Anders},
##   year = {2014},
##   journal = {Genome Biology},
##   doi = {10.1186/s13059-014-0550-8},
##   volume = {15},
##   issue = {12},
##   pages = {550},
## }

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

dds

```



```

## ENSG00000283116      NA
## ENSG00000283119      NA
## ENSG00000283120      NA
## ENSG00000283123      NA

res <- results(dds)
res

## log2 fold change (MLE): dex treated vs control
## Wald test p-value: dex treated vs control
## DataFrame with 38694 rows and 6 columns
##           baseMean log2FoldChange      lfcSE      stat     pvalue
## <numeric>      <numeric> <numeric> <numeric> <numeric>
## ENSG00000000003  747.1942   -0.3507030  0.168246 -2.084470 0.0371175
## ENSG00000000005   0.0000      NA        NA        NA        NA
## ENSG00000000419  520.1342   0.2061078  0.101059  2.039475 0.0414026
## ENSG00000000457  322.6648   0.0245269  0.145145  0.168982 0.8658106
## ENSG00000000460   87.6826   -0.1471420  0.257007 -0.572521 0.5669691
## ...
##           ...       ...       ...       ...       ...
## ENSG00000283115  0.000000      NA        NA        NA        NA
## ENSG00000283116  0.000000      NA        NA        NA        NA
## ENSG00000283119  0.000000      NA        NA        NA        NA
## ENSG00000283120  0.974916   -0.668258  1.69456  -0.394354 0.693319
## ENSG00000283123  0.000000      NA        NA        NA        NA
##           padj
## <numeric>
## ENSG00000000003  0.163035
## ENSG00000000005   NA
## ENSG00000000419  0.176032
## ENSG00000000457  0.961694
## ENSG00000000460  0.815849
## ...
##           ...
## ENSG00000283115   NA
## ENSG00000283116   NA
## ENSG00000283119   NA
## ENSG00000283120   NA
## ENSG00000283123   NA

summary(res)

##
## out of 25258 with nonzero total read count
## adjusted p-value < 0.1
## LFC > 0 (up)      : 1563, 6.2%
## LFC < 0 (down)    : 1188, 4.7%
## outliers [1]       : 142, 0.56%
## low counts [2]     : 9971, 39%
## (mean count < 10)
## [1] see 'cooksCutoff' argument of ?results
## [2] see 'independentFiltering' argument of ?results

```

```
res05 <- results(dds, alpha=0.05)
summary(res05)

## 
## out of 25258 with nonzero total read count
## adjusted p-value < 0.05
## LFC > 0 (up)      : 1236, 4.9%
## LFC < 0 (down)    : 933, 3.7%
## outliers [1]       : 142, 0.56%
## low counts [2]     : 9033, 36%
## (mean count < 6)
## [1] see 'cooksCutoff' argument of ?results
## [2] see 'independentFiltering' argument of ?results
```

##Save our results

write out whole results dataset (including genes that don't change significantly)

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

focus in on those genes with a small p-value (i.e. show a significant change)

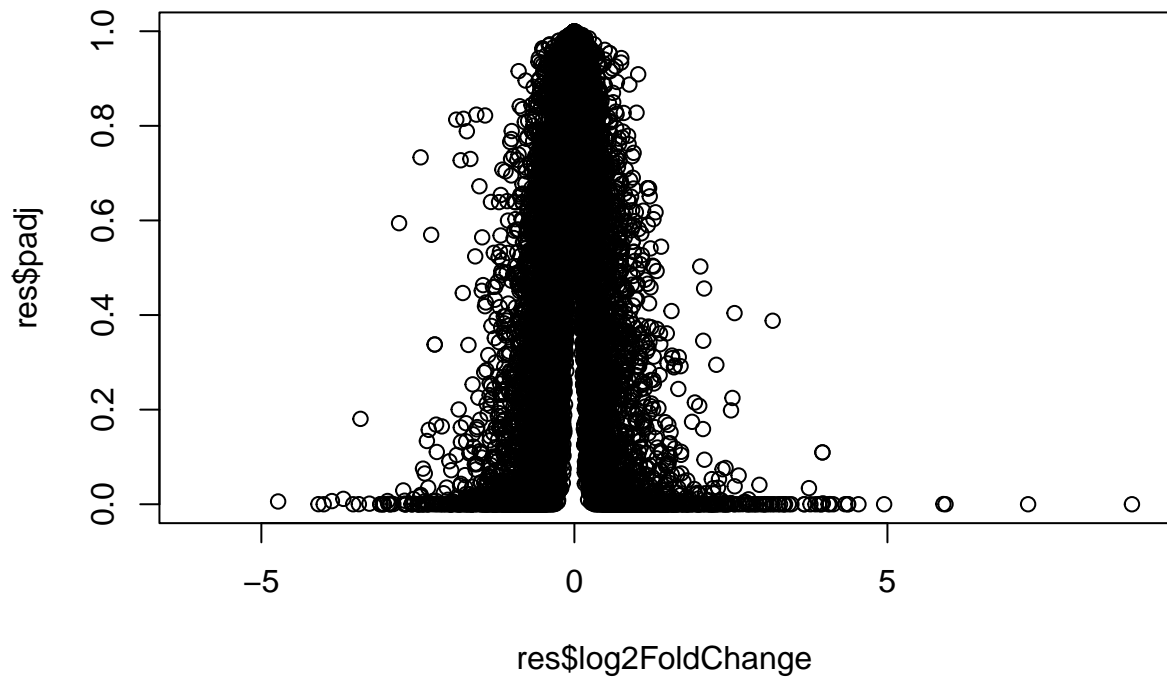
```
res05 <- results(dds, alpha=0.05)
summary(res05)
```

```
## 
## out of 25258 with nonzero total read count
## adjusted p-value < 0.05
## LFC > 0 (up)      : 1236, 4.9%
## LFC < 0 (down)    : 933, 3.7%
## outliers [1]       : 142, 0.56%
## low counts [2]     : 9033, 36%
## (mean count < 6)
## [1] see 'cooksCutoff' argument of ?results
## [2] see 'independentFiltering' argument of ?results
```

##Volcano plots Let's make a commonly produced visualization from this data, namely a so-called Volcano plot. These summary figures are frequently used to highlight the proportion of genes that are both significantly regulated and display a high fold change.

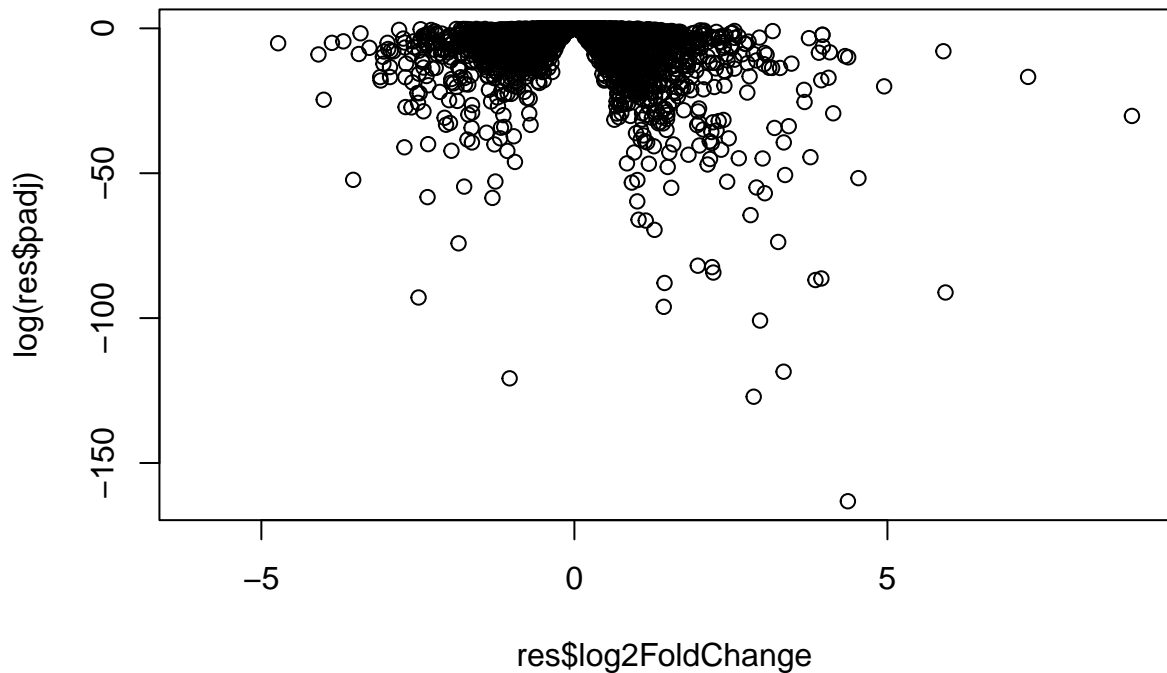
This is not useful

```
plot(res$log2FoldChange, res$padj)
```

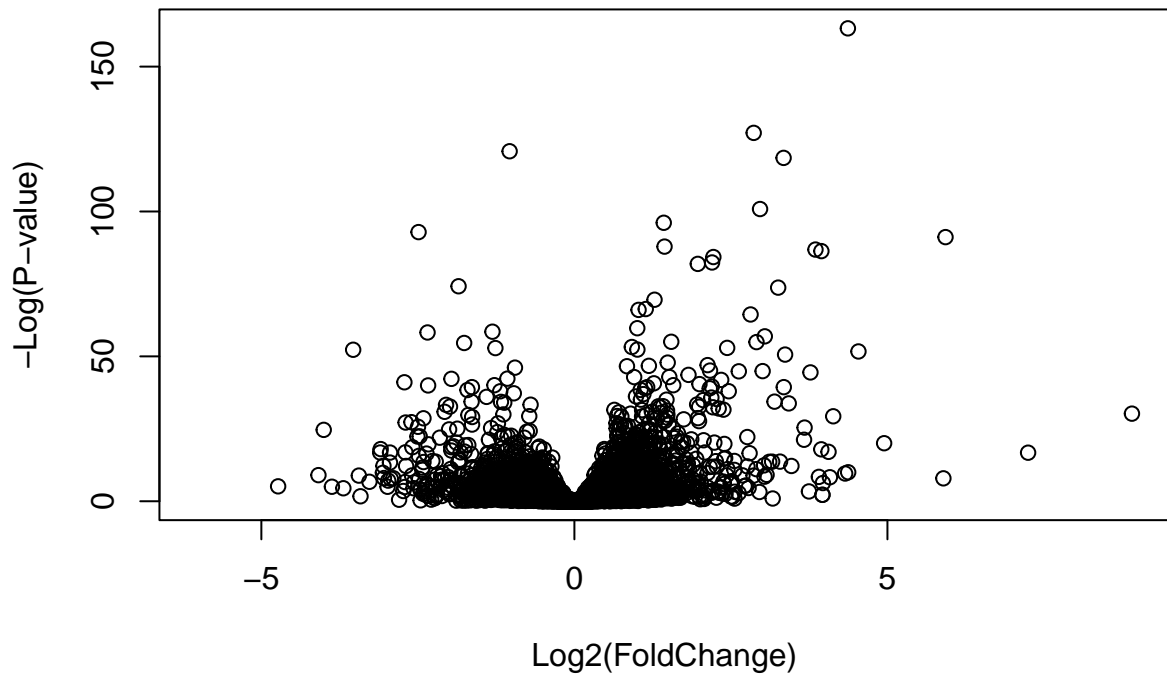


we can flip this

```
plot( res$log2FoldChange, log(res$padj))
```

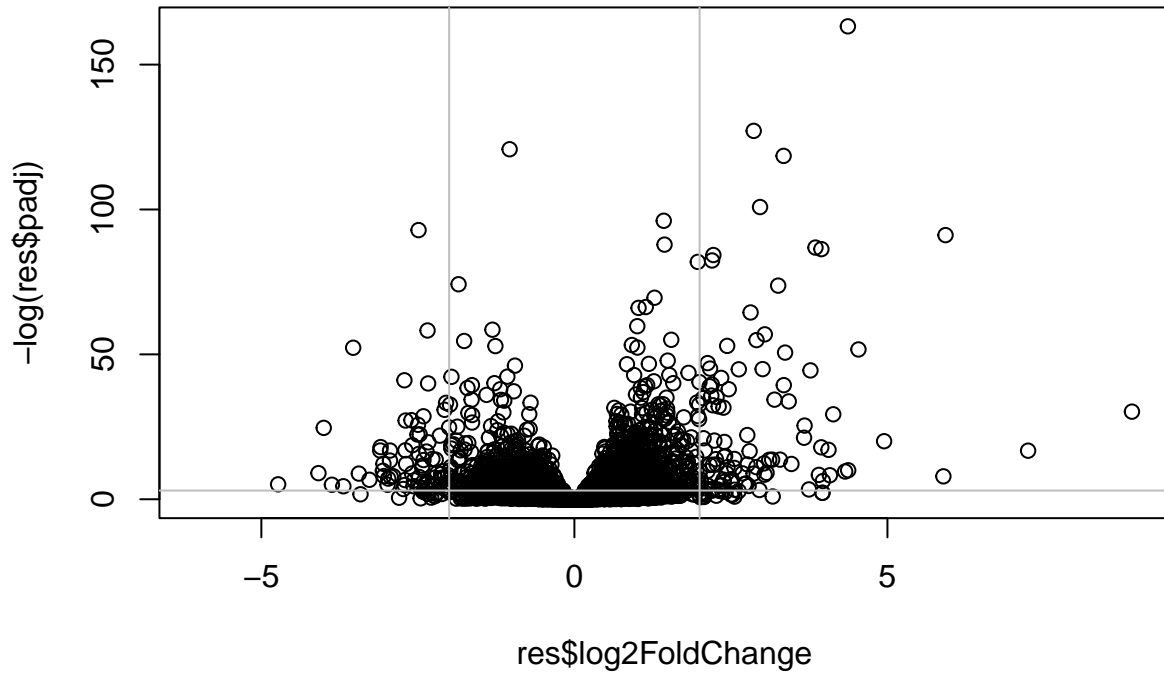


```
plot( res$log2FoldChange, -log(res$padj),
      xlab="Log2(FoldChange)",
      ylab="-Log(P-value)")
```



finally let's add some color to this plot to draw attention to the genes (i.e. points) we care about - that is those with large fold-change and low pvalues (i.e. high $-\log(\text{pvalues})$)

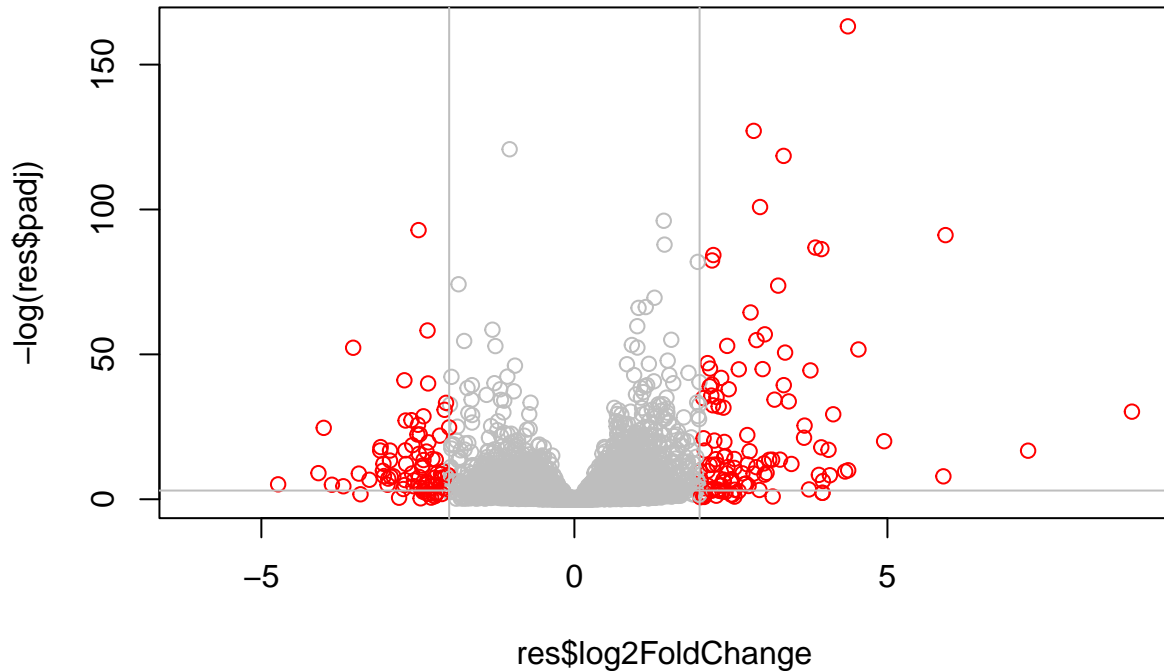
```
plot( res$log2FoldChange, -log(res$padj))
abline(v=c(-2, +2), col="gray")
abline(h=-log(0.05), col="gray")
```



add some color to the points:

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

plot( res$log2FoldChange, -log(res$padj), col=mycols)
abline(v=c(-2, +2), col="gray")
abline(h=-log(0.05), col="gray")
```



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

Add annotation data for our genes
 for this we need two biconductor packages - BiocManager::install("AnnotationDbi") - BiocManager::install("org.Hs.eg.db")

```
library("AnnotationDbi")
```

```
## Warning: package 'AnnotationDbi' was built under R version 4.1.2
```

```
##  

## Attaching package: 'AnnotationDbi'
```

```
## The following object is masked from 'package:dplyr':  

##  

##     select
```

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

```
##
```

Let's have a look at what is in the '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"
```

We will use the ‘mapIDs’ function to translate between identifiers from different databases.

```
res$symbol <- mapIds(org.Hs.eg.db,
                      keys=row.names(res), # Our genenames
                      keytype="ENSEMBL",      # The format of our genenames
                      column="SYMBOL",       # The new format we want to add
                      multiVals="first")
```

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

We need ENTREZ ids for pathway analysis with KEGG

```
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"
```

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

```
res$entrez <- mapIds(org.Hs.eg.db,
                      keys=row.names(res), # Our genenames
                      keytype="ENSEMBL",      # The format of our genenames
                      column="ENTREZID",       # The new format we want to add
                      multiVals="first")
```

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

```
res$uniprot <- mapIds(org.Hs.eg.db,
                      keys=row.names(res),
                      column="UNIPROT",
                      keytype="ENSEMBL",
                      multiVals="first")
```

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

```

res$genename <- mapIds(org.Hs.eg.db,
                      keys=row.names(res),
                      column="GENENAME",
                      keytype="ENSEMBL",
                      multiVals="first")

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

head(res)

## log2 fold change (MLE): dex treated vs control
## Wald test p-value: dex treated vs control
## DataFrame with 6 rows and 10 columns
##           baseMean log2FoldChange      lfcSE      stat     pvalue
##           <numeric>      <numeric> <numeric> <numeric>
## ENSG00000000003 747.194195     -0.3507030  0.168246 -2.084470 0.0371175
## ENSG00000000005  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      symbol      entrez      uniprot
##           <numeric> <character> <character> <character>
## ENSG00000000003 0.163035     TSPAN6      7105 AOA024RCIO
## ENSG00000000005   NA         TNMD       64102 Q9H2S6
## ENSG00000000419 0.176032     DPM1       8813 060762
## ENSG00000000457 0.961694     SCYL3      57147 Q8IZE3
## ENSG00000000460 0.815849     C1orf112    55732 AOA024R922
## ENSG00000000938   NA         FGR        2268 P09769
##           genename
##           <character>
## ENSG00000000003      tetraspanin 6
## ENSG00000000005      tenomodulin
## ENSG00000000419      dolichyl-phosphate m..
## ENSG00000000457      SCY1 like pseudokina..
## ENSG00000000460      chromosome 1 open re..
## ENSG00000000938      FGR proto-oncogene, ..

```

Let's make another volcano plot with some gene labels for this we can use the **EnhancedVolcano** package

```

library(EnhancedVolcano)

## Loading required package: ggplot2

## Loading required package: ggrepel

## Registered S3 methods overwritten by 'ggalt':
##   method             from
##   grid.draw.absoluteGrob  ggplot2
##   grobHeight.absoluteGrob ggplot2
##   grobWidth.absoluteGrob ggplot2
##   grobX.absoluteGrob    ggplot2
##   grobY.absoluteGrob    ggplot2

```

```

x <- as.data.frame(res)

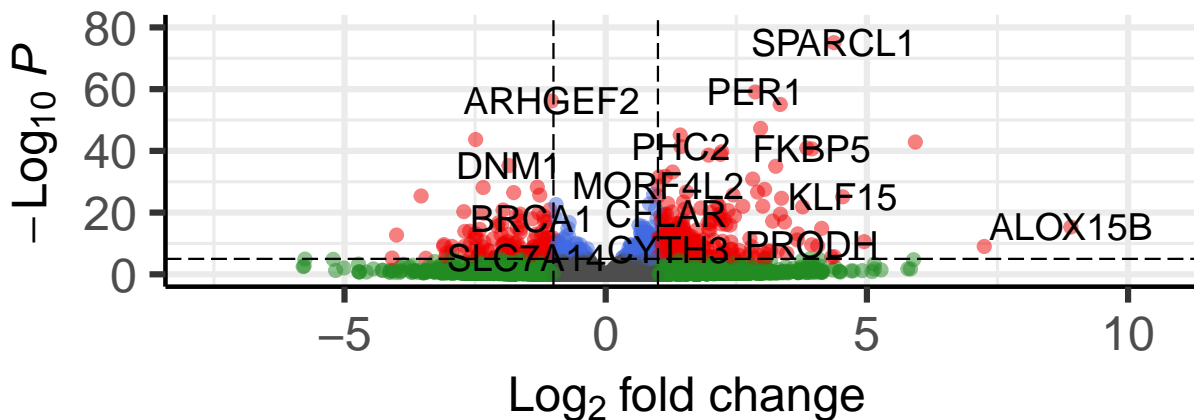
EnhancedVolcano(x,
  lab = x$symbol,
  x = 'log2FoldChange',
  y = 'pvalue')

```

Volcano plot

EnhancedVolcano

● NS ● Log₂ FC ● p-value ● p-value and log₂ FC



```
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)

head(kegg.sets.hs, 2)

## $`hsa00232 Caffeine metabolism`
## [1] "10"   "1544" "1548" "1549" "1553" "7498" "9"
##
## $`hsa00983 Drug metabolism - other enzymes`
## [1] "10"     "1066"   "10720"  "10941"  "151531" "1548"   "1549"   "1551"
## [9] "1553"   "1576"   "1577"   "1806"   "1807"   "1890"   "221223" "2990"
## [17] "3251"   "3614"   "3615"   "3704"   "51733"  "54490"  "54575"  "54576"
## [25] "54577"  "54578"  "54579"  "54600"  "54657"  "54658"  "54659"  "54963"
## [33] "574537" "64816"  "7083"   "7084"   "7172"   "7363"   "7364"   "7365"
## [41] "7366"   "7367"   "7371"   "7372"   "7378"   "7498"   "79799" "83549"
## [49] "8824"   "8833"   "9"      "978"

```

The main `gage()` function requires a named vector of fold changes, where the names of the values are the Entrez gene IDs.

```

foldchange <- res$log2FoldChange
names(foldchange) <- res$entrez
head(foldchange)

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

```

```

# Get the results
keggres = gage(foldchange, gsets=kegg.sets.hs)

```

```

attributes(keggres)

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

```

This separates out results by “greater” and “less” i.e. those that are up regulated and those that are down regulated

```

# Look at the first three down (less) pathways
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

```

Now, let's try out the 'pathview()' function from the **pathview** package to make a pathway plot with our RNA-Seq expression results shown in color.

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

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

```
## Info: Working in directory /Users/anelvaldez/Desktop/Desktop/bimm143/bimm143_github2/class 15
```

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

