

Class 13: RNAseq Mini Project

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Background

Today we will run through a complete RNAseq analysis

The data for for hands-on session comes from GEO entry: GSE37704, which is associated with the following publication:

Trapnell C, Hendrickson DG, Sauvageau M, Goff L et al. “Differential analysis of gene regulation at transcript resolution with RNA-seq”. Nat Biotechnol 2013 Jan;31(1):46-53. PMID: 23222703

The authors report on differential analysis of lung fibroblasts in response to loss of the developmental transcription factor HOXA1. Their results and others indicate that HOXA1 is required for lung fibroblast and HeLa cell cycle progression.

Data Import

```
counts <- read.csv("/Users/wadeingersoll/Desktop/BGGN213/class13/GSE37704_featurecounts.csv")
metadata <- read.csv("/Users/wadeingersoll/Desktop/BGGN213/class13/GSE37704_metadata.csv")
```

Check correspondence of `metadata` and `counts` (i.e. that the columns in `counts` match the rows in the `metadata`).

```
metadata
```

```
      id    condition
1 SRR493366 control_sirna
2 SRR493367 control_sirna
3 SRR493368 control_sirna
4 SRR493369    hoxa1_kd
5 SRR493370    hoxa1_kd
6 SRR493371    hoxa1_kd
```

```
head(counts)
```

	length	SRR493366	SRR493367	SRR493368	SRR493369	SRR493370
ENSG00000186092	918	0	0	0	0	0
ENSG00000279928	718	0	0	0	0	0
ENSG00000279457	1982	23	28	29	29	28
ENSG00000278566	939	0	0	0	0	0
ENSG00000273547	939	0	0	0	0	0
ENSG00000187634	3214	124	123	205	207	212
						SRR493371
ENSG00000186092		0				
ENSG00000279928		0				
ENSG00000279457		46				
ENSG00000278566		0				
ENSG00000273547		0				
ENSG00000187634		258				

```
colnames(counts)
```

```
[1] "length"      "SRR493366"   "SRR493367"   "SRR493368"   "SRR493369"   "SRR493370"
[7] "SRR493371"
```

```
metadata$id
```

```
[1] "SRR493366" "SRR493367" "SRR493368" "SRR493369" "SRR493370" "SRR493371"
```

Fix to remove that first “length” column of `counts`

```
counts <- counts[,-1]
```

Also let's remove low count genes

```
tot.counts <- rowSums(counts)
head(tot.counts)
```

```
ENSG00000186092 ENSG00000279928 ENSG00000279457 ENSG00000278566 ENSG00000273547
          0           183           0           0
ENSG00000187634
          1129
```

Let's remove all zero count genes

```
zero inds <- tot.counts == 0
head(zero inds)
```

```
ENSG00000186092 ENSG00000279928 ENSG00000279457 ENSG00000278566 ENSG00000273547
      TRUE        TRUE       FALSE       TRUE       TRUE
ENSG00000187634
      FALSE
```

```
counts <- counts[!zero inds,]
```

```
all(c(T, T, F))
```

```
[1] FALSE
```

```
test_cols <- !all(colnames(counts) == metadata$id)
```

```
if( test_cols ) {
  message("Wow... there is a problem with the metadata counts setup")
}
```

Setup for DESeq

```
library(DESeq2)
```

Loading required package: S4Vectors

Loading required package: stats4

Loading required package: BiocGenerics

Loading required package: generics

Attaching package: 'generics'

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

```
as.difftime, as.factor, as.ordered, intersect, is.element, setdiff,
setequal, union
```

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, is.unsorted, lapply, Map, mapply, match, mget,
order, paste, pmax, pmax.int, pmin, pmin.int, Position, rank,
rbind, Reduce, rownames, sapply, saveRDS, table, tapply, 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
```

```
Loading required package: GenomicRanges
```

```
Loading required package: Seqinfo
```

```
Loading required package: SummarizedExperiment
```

```
Loading required package: MatrixGenerics
```

```
Loading required package: matrixStats
```

```
Attaching package: 'MatrixGenerics'
```

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

```
  colAlls, colAnyNAs, colAnyNs, 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, rowAnyNs, rowAvgsPerColSet,
  rowCollapse, rowCounts, rowCummaxs, rowCummins, rowCumprods,
  rowCumsums, rowDiffs, rowIQRDiffs, rowIQRs, rowLogSumExps,
  rowMadDiffs, rowMads, rowMaxs, rowMeans2, rowMedians, rowMins,
```

```
rowOrderStats, rowProds, rowQuantiles, rowRanges, rowRanks,  
rowSdDiff, rowSds, rowSums2, rowTabulates, rowVarDiff, 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
```

```
library(DESeq2)  
dds <- DESeqDataSetFromMatrix(countData = counts,  
                                colData = metadata,  
                                design = ~condition)
```

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

Run DESeq

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

```
estimating size factors
```

```
estimating dispersions  
gene-wise dispersion estimates  
mean-dispersion relationship  
final dispersion estimates  
fitting model and testing
```

Get results

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

```
head(res)
```

```
log2 fold change (MLE): condition hoxa1 kd vs control sirna  
Wald test p-value: condition hoxa1 kd vs control sirna  
DataFrame with 6 rows and 6 columns  
  baseMean log2FoldChange      lfcSE       stat      pvalue  
  <numeric>      <numeric> <numeric>      <numeric>    <numeric>  
ENSG00000279457  29.9136     0.1792571  0.3248216   0.551863 5.81042e-01  
ENSG00000187634 183.2296     0.4264571  0.1402658   3.040350 2.36304e-03  
ENSG00000188976 1651.1881    -0.6927205 0.0548465 -12.630158 1.43990e-36  
ENSG00000187961 209.6379     0.7297556  0.1318599   5.534326 3.12428e-08  
ENSG00000187583 47.2551      0.0405765  0.2718928   0.149237 8.81366e-01  
ENSG00000187642 11.9798      0.5428105  0.5215598   1.040744 2.97994e-01  
  padj  
  <numeric>  
ENSG00000279457 6.86555e-01  
ENSG00000187634 5.15718e-03  
ENSG00000188976 1.76549e-35  
ENSG00000187961 1.13413e-07  
ENSG00000187583 9.19031e-01  
ENSG00000187642 4.03379e-01
```

Add annotation

```
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"
```

```
res$symbol = mapIds(org.Hs.eg.db,
                     keys=row.names(res),
                     keytype="ENSEMBL",
                     column="SYMBOL",
                     multiVals="first")
```

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

```
res$entrez = mapIds(org.Hs.eg.db,
                     keys=row.names(res),
                     keytype="ENSEMBL",
                     column="ENTREZID",
                     multiVals="first")
```

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

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

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

```
head(res, 10)
```

```
log2 fold change (MLE): condition hoxa1 kd vs control sirna
Wald test p-value: condition hoxa1 kd vs control sirna
DataFrame with 10 rows and 9 columns
  baseMean log2FoldChange      lfcSE      stat     pvalue
  <numeric>      <numeric> <numeric>      <numeric>    <numeric>
ENSG00000279457   29.913579   0.1792571  0.3248216   0.551863 5.81042e-01
ENSG00000187634  183.229650   0.4264571  0.1402658   3.040350 2.36304e-03
ENSG00000188976 1651.188076  -0.6927205  0.0548465 -12.630158 1.43990e-36
ENSG00000187961  209.637938   0.7297556  0.1318599   5.534326 3.12428e-08
ENSG00000187583  47.255123   0.0405765  0.2718928   0.149237 8.81366e-01
ENSG00000187642  11.979750   0.5428105  0.5215598   1.040744 2.97994e-01
ENSG00000188290  108.922128   2.0570638  0.1969053   10.446970 1.51282e-25
ENSG00000187608  350.716868   0.2573837  0.1027266   2.505522 1.22271e-02
ENSG00000188157  9128.439422  0.3899088  0.0467163   8.346304 7.04321e-17
ENSG00000237330   0.158192   0.7859552  4.0804729   0.192614 8.47261e-01
  padj      symbol      entrez             name
  <numeric> <character> <character>           <character>
ENSG00000279457 6.86555e-01        NA        NA            NA
ENSG00000187634 5.15718e-03       SAMD11    148398 sterile alpha motif ..
ENSG00000188976 1.76549e-35       NOC2L     26155 NOC2 like nucleolar ..
ENSG00000187961 1.13413e-07      KLHL17    339451 kelch like family me..
ENSG00000187583 9.19031e-01      PLEKHN1   84069 pleckstrin homology ..
ENSG00000187642 4.03379e-01      PERM1     84808 PPARGC1 and ESRR ind..
ENSG00000188290 1.30538e-24       HES4     57801 hes family bHLH tran..
ENSG00000187608 2.37452e-02      ISG15     9636 ISG15 ubiquitin like..
ENSG00000188157 4.21963e-16      AGRN     375790          agrin
ENSG00000237330          NA      RNF223    401934 ring finger protein ..
```

Visualize results

```
library(ggplot2)

# Make a color vector for all genes
mycols <- rep("gray", nrow(res) )

# Color red the genes with absolute fold change above 2
mycols[ abs(res$log2FoldChange) > 2 ] <- "red"
```

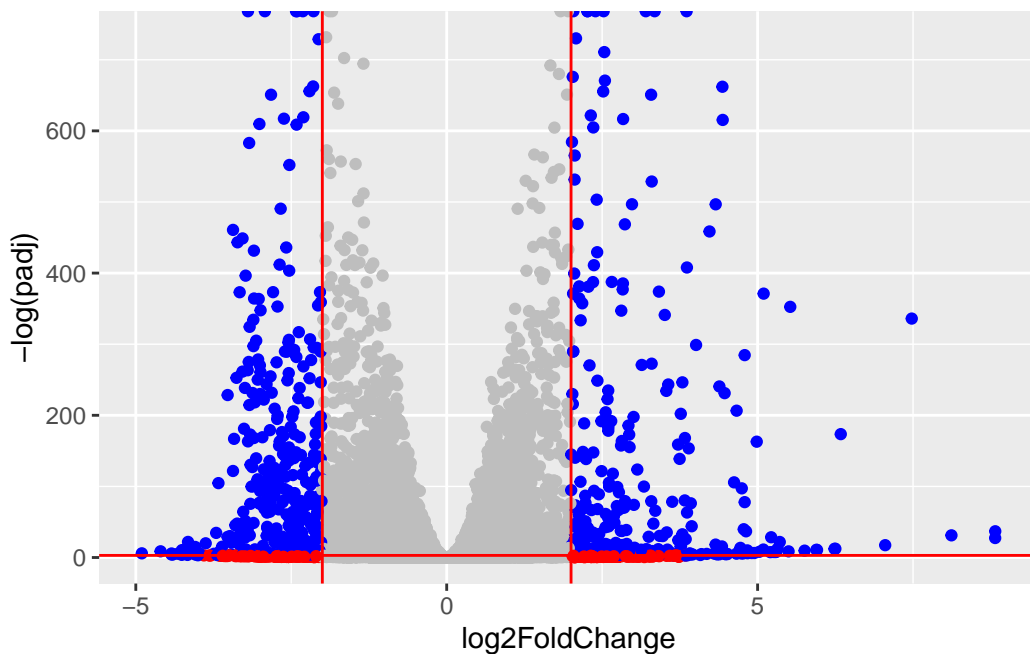
```

# Color blue those with adjusted p-value less than 0.01
# and absolute fold change more than 2
inds <- (res$padj < .05) & (abs(res$log2FoldChange) > 2 )
mycols[ inds ] <- "blue"

ggplot(res) + aes(log2FoldChange, -log(padj)) +
  geom_point(col=mycols) +
  geom_vline(xintercept = c(-2,2), col='red') +
  geom_hline(yintercept = -log(0.05), col='red')

```

Warning: Removed 1237 rows containing missing values or values outside the scale range
(`geom_point()`).



Pathway analysis

```
library(gage)
```

```
library(gageData)

data(kegg.sets.hs)
data(sigmet.idx.hs)
```

For **gage** we want a named vector of importance

GO Analysis

Let's try GO Analysis and compare to KEGG analysis

```
data(go.sets.hs)
data(go.subs.hs)

# # Focus on Biological Process subset of GO
# gobpsets = go.sets.hs[go.subs.hs$BP]
#
# gobpres = gage(foldchanges, gsets=gobpsets, same.dir=TRUE)
#
# lapply(gobpres, head)
```

Reactome

Some folks really like Reactome online (i.e. their webpage viewer) rather than the R package of the same name (available from bioconductor).

To use the viewer we want to upload our set of gene symbols for the genes we want to focus on (here those with a P-value below 0.05)

```
sig_genes <- res[res$padj <= 0.05 & !is.na(res$padj), "symbol"]
print(paste("Total number of significant genes:", length(sig_genes)))
```

```
[1] "Total number of significant genes: 8147"
```

```
head(sig_genes)
```

```
ENSG00000187634 ENSG00000188976 ENSG00000187961 ENSG00000188290 ENSG00000187608
      "SAMD11"          "NOC2L"          "KLHL17"          "HES4"          "ISG15"
ENSG00000188157
      "AGRN"
```

Save results

```
#save(res, file="my_results.RData")
```