class13

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Differential Section Analysis

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

Attaching package: 'S4Vectors'

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

expand.grid, I, unname

Loading required package: IRanges

Loading required package: GenomicRanges

Loading required package: GenomeInfoDb

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, 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, rowWeighted

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
Let's load our data and take a look at it.
  # Import metadata and take a peak
  colData = read.csv("GSE37704_metadata.csv", row.names=1)
  head(colData)
              condition
SRR493366 control_sirna
SRR493367 control_sirna
SRR493368 control_sirna
SRR493369
               hoxa1_kd
               hoxa1_kd
SRR493370
SRR493371
               hoxa1_kd
  # Import countdata
  countData = read.csv("GSE37704_featurecounts.csv", row.names=1)
  head(countData)
                length SRR493366 SRR493367 SRR493368 SRR493369 SRR493370
                   918
                               0
                                                   0
ENSG00000186092
                                         0
                                                              0
                                                                        0
                   718
                              0
ENSG00000279928
                                         0
                                                   0
                                                             0
                                                                        0
                              23
                                        28
                                                   29
                                                             29
ENSG00000279457
                  1982
                                                                       28
```

TNGGOOOOOTOE	000	^	^	^	^	^
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					

```
# Note we need to remove the odd first $length col
countData <- as.matrix(countData[,-1])
head(countData)</pre>
```

	SRR493366	SRR493367	SRR493368	SRR493369	SRR493370	SRR493371
ENSG00000186092	0	0	0	0	0	0
ENSG00000279928	0	0	0	0	0	0
ENSG00000279457	23	28	29	29	28	46
ENSG00000278566	0	0	0	0	0	0
ENSG00000273547	0	0	0	0	0	0
ENSG00000187634	124	123	205	207	212	258

```
# Filter count data where you have 0 read count across all samples. zerocounts <- rowSums(countData) == \frac{0}{100} head(zerocounts)
```

ENSG00000186092 ENSG00000279928 ENSG00000279457 ENSG00000278566 ENSG00000273547

TRUE TRUE FALSE TRUE TRUE

FALSE

```
newcounts <- countData[!zerocounts,]
head(newcounts)</pre>
```

	SRR493366	SRR493367	SRR493368	SRR493369	SRR493370	SRR493371
ENSG00000279457	23	28	29	29	28	46
ENSG00000187634	124	123	205	207	212	258
ENSG00000188976	1637	1831	2383	1226	1326	1504

ENSG00000187961	120	153	180	236	255	357
ENSG00000187583	24	48	65	44	48	64
ENSG00000187642	4	9	16	14	16	16

nrow(newcounts)

[1] 15975

Running DESeq2

Nice now lets setup the DESeqDataSet object required for the DESeq() function and then run the DESeq pipeline.

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

```
dds = DESeq(dds)

estimating size factors

estimating dispersions

gene-wise dispersion estimates

mean-dispersion relationship

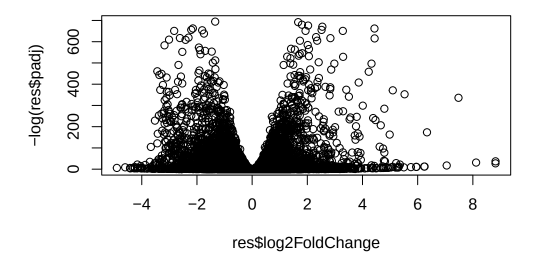
final dispersion estimates

fitting model and testing

## estimating size factors
```

estimating dispersions

```
## gene-wise dispersion estimates
  ## mean-dispersion relationship
  ## final dispersion estimates
  ## fitting model and testing
  dds
class: DESeqDataSet
dim: 15975 6
metadata(1): version
assays(4): counts mu H cooks
rownames(15975): ENSG00000279457 ENSG00000187634 ... ENSG00000276345
  ENSG00000271254
rowData names(22): baseMean baseVar ... deviance maxCooks
colnames(6): SRR493366 SRR493367 ... SRR493370 SRR493371
colData names(2): condition sizeFactor
  colData = read.csv("GSE37704_metadata.csv", row.names=1)
  res = results(dds, contrast=c("condition", "hoxa1_kd", "control_sirna"))
  summary(res)
out of 15975 with nonzero total read count
adjusted p-value < 0.1
LFC > 0 (up)
                   : 4349, 27%
LFC < 0 (down)
                  : 4396, 28%
outliers [1]
                   : 0, 0%
low counts [2]
                   : 1237, 7.7%
(mean count < 0)
[1] see 'cooksCutoff' argument of ?results
[2] see 'independentFiltering' argument of ?results
Volcano plot A plot of log2 fold change vs -log adjusted p-value.
  plot( res$log2FoldChange, -log(res$padj) )
```

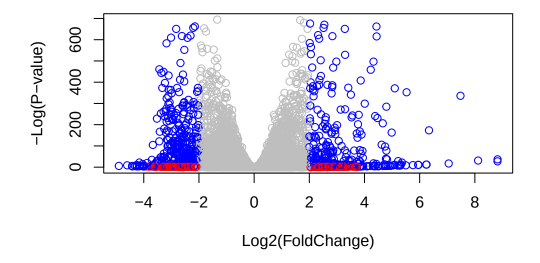


```
# Make a color vector for all genes
mycols <- rep("gray", nrow(res) )

# 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
mycols[ res$padj<0.05 & abs(res$log2FoldChange) > 2] <- "blue"

plot( res$log2FoldChange, -log(res$padj), col=mycols, xlab="Log2(FoldChange)", ylab="-Log(</pre>
```



Adding gene annotation

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

columns(org.Hs.eg.db)

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

```
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
                                                       3.040350 2.36304e-03
ENSG00000187634 183.229650
                                 0.4264571 0.1402658
ENSG00000188976 1651.188076
```

ENSG00000187642

ENSG00000237330

	padj	symbol	entrez	name
	<numeric></numeric>	<character></character>	<character></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

```
res = res[order(res$pvalue),]
write.csv(res, file="deseq_results.csv")
```

Pathway analysis

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)
  data(sigmet.idx.hs)
  # Focus on signaling and metabolic pathways only
  kegg.sets.hs = kegg.sets.hs[sigmet.idx.hs]
  # Examine the first 3 pathways
  head(kegg.sets.hs, 3)
$`hsa00232 Caffeine metabolism`
[1] "10"
           "1544" "1548" "1549" "1553" "7498" "9"
$`hsa00983 Drug metabolism - other enzymes`
 [1] "10"
               "1066"
                        "10720"
                                            "151531" "1548"
                                                               "1549"
                                                                         "1551"
                                  "10941"
               "1576"
                        "1577"
 [9] "1553"
                                  "1806"
                                            "1807"
                                                      "1890"
                                                               "221223" "2990"
[17] "3251"
               "3614"
                        "3615"
                                  "3704"
                                            "51733"
                                                      "54490"
                                                               "54575"
                                                                         "54576"
                                            "54657"
[25] "54577"
               "54578"
                        "54579"
                                  "54600"
                                                      "54658"
                                                               "54659"
                                                                         "54963"
[33] "574537" "64816"
                        "7083"
                                  "7084"
                                            "7172"
                                                      "7363"
                                                               "7364"
                                                                         "7365"
                                            "7378"
[41] "7366"
               "7367"
                        "7371"
                                  "7372"
                                                      "7498"
                                                               "79799"
                                                                         "83549"
[49] "8824"
                        "9"
                                  "978"
               "8833"
$`hsa00230 Purine metabolism`
  [1] "100"
                "10201"
                         "10606"
                                   "10621"
                                             "10622"
                                                       "10623"
                                                                "107"
                                                                          "10714"
  [9] "108"
                "10846"
                         "109"
                                   "111"
                                             "11128"
                                                       "11164"
                                                                 "112"
                                                                          "113"
 [17] "114"
                "115"
                         "122481" "122622" "124583" "132"
                                                                 "158"
                                                                          "159"
                "171568" "1716"
                                   "196883"
                                             "203"
                                                       "204"
                                                                 "205"
                                                                          "221823"
 [25] "1633"
 [33] "2272"
                "22978"
                         "23649"
                                   "246721" "25885"
                                                       "2618"
                                                                 "26289"
                                                                          "270"
 [41] "271"
                "27115"
                         "272"
                                             "2977"
                                                                          "2984"
                                   "2766"
                                                       "2982"
                                                                 "2983"
 [49] "2986"
                "2987"
                         "29922"
                                   "3000"
                                             "30833"
                                                       "30834"
                                                                "318"
                                                                          "3251"
                                                                 "4830"
 [57] "353"
                "3614"
                         "3615"
                                   "3704"
                                             "377841"
                                                      "471"
                                                                          "4831"
                "4833"
                                             "4882"
 [65] "4832"
                         "4860"
                                   "4881"
                                                       "4907"
                                                                 "50484"
                                                                          "50940"
                "51251"
                                   "5136"
                                             "5137"
                                                       "5138"
                                                                 "5139"
                                                                          "5140"
 [73] "51082"
                         "51292"
 [81] "5141"
                "5142"
                         "5143"
                                   "5144"
                                             "5145"
                                                       "5146"
                                                                "5147"
                                                                          "5148"
 [89] "5149"
                "5150"
                         "5151"
                                   "5152"
                                             "5153"
                                                       "5158"
                                                                "5167"
                                                                          "5169"
 [97] "51728"
                                                                 "54107"
                                                                          "5422"
                "5198"
                         "5236"
                                   "5313"
                                             "5315"
                                                       "53343"
[105] "5424"
                "5425"
                         "5426"
                                   "5427"
                                             "5430"
                                                       "5431"
                                                                 "5432"
                                                                          "5433"
                                   "5437"
[113] "5434"
                "5435"
                          "5436"
                                             "5438"
                                                       "5439"
                                                                 "5440"
                                                                          "5441"
[121] "5471"
                "548644" "55276"
                                   "5557"
                                             "5558"
                                                       "55703"
                                                                "55811"
                                                                          "55821"
[129] "5631"
                "5634"
                          "56655"
                                   "56953"
                                             "56985"
                                                       "57804"
                                                                 "58497"
                                                                          "6240"
                                                       "7498"
                                                                 "8382"
[137] "6241"
                "64425"
                         "646625" "654364"
                                             "661"
                                                                          "84172"
                                                                          "9060"
                                   "8622"
                                                                "8833"
[145] "84265"
                "84284"
                         "84618"
                                             "8654"
                                                       "87178"
[153] "9061"
                "93034"
                         "953"
                                   "9533"
                                             "954"
                                                       "955"
                                                                 "956"
                                                                          "957"
```

```
[161] "9583" "9615"
```

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

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

```
1266 54855 1465 51232 2034 2317 -2.422719 3.201955 -2.313738 -2.059631 -1.888019 -1.649792
```

Now, let's run the gage pathway analysis.

```
# Get the results
keggres = gage(foldchanges, gsets=kegg.sets.hs)
attributes(keggres)
```

\$names

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

```
# Look at the first few down (less) pathways
head(keggres$less)
```

```
p.geomean stat.mean
hsa04110 Cell cycle
                                      8.995727e-06 -4.378644 8.995727e-06
hsa03030 DNA replication
                                      9.424076e-05 -3.951803 9.424076e-05
hsa03013 RNA transport
                                      1.375901e-03 -3.028500 1.375901e-03
hsa03440 Homologous recombination
                                      3.066756e-03 -2.852899 3.066756e-03
hsa04114 Oocyte meiosis
                                      3.784520e-03 -2.698128 3.784520e-03
hsa00010 Glycolysis / Gluconeogenesis 8.961413e-03 -2.405398 8.961413e-03
                                            q.val set.size
                                                                   exp1
                                      0.001448312
                                                       121 8.995727e-06
hsa04110 Cell cycle
hsa03030 DNA replication
                                      0.007586381
                                                        36 9.424076e-05
hsa03013 RNA transport
                                      0.073840037
                                                       144 1.375901e-03
hsa03440 Homologous recombination
                                      0.121861535
                                                       28 3.066756e-03
hsa04114 Oocyte meiosis
                                      0.121861535
                                                       102 3.784520e-03
hsa00010 Glycolysis / Gluconeogenesis 0.212222694
                                                       53 8.961413e-03
```

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=foldchanges, pathway.id="hsa04110")
'select()' returned 1:1 mapping between keys and columns
Info: Working in directory /Users/anntruong/class13
Info: Writing image file hsa04110.pathview.png
```

Now, let's process our results a bit more to automagically pull out the top 5 upregulated pathways, then further process that just to get the pathway IDs needed by the pathwiew() function.

```
## Focus on top 5 upregulated pathways here for demo purposes only
keggrespathways <- rownames(keggres$greater)[1:5]

# Extract the 8 character long IDs part of each string
keggresids = substr(keggrespathways, start=1, stop=8)
keggresids</pre>
```

[1] "hsa04640" "hsa04630" "hsa00140" "hsa04142" "hsa04330"

Finally, lets pass these IDs in keggresids to the pathview() function to draw plots for all the top 5 pathways.

```
pathview(gene.data=foldchanges, pathway.id=keggresids, species="hsa")
'select()' returned 1:1 mapping between keys and columns
Info: Working in directory /Users/anntruong/class13
Info: Writing image file hsa04640.pathview.png
'select()' returned 1:1 mapping between keys and columns
Info: Working in directory /Users/anntruong/class13
```

Info: Writing image file hsa04630.pathview.png

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

Info: Working in directory /Users/anntruong/class13

Info: Writing image file hsa00140.pathview.png

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

Info: Working in directory /Users/anntruong/class13

Info: Writing image file hsa04142.pathview.png

Info: some node width is different from others, and hence adjusted!

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

Info: Working in directory /Users/anntruong/class13

Info: Writing image file hsa04330.pathview.png

Can you do the same procedure as above to plot the pathview figures for the top 5 down-reguled pathways? Yes, we can just manipulate the code chunk from rownames(keggres\$greater) to rownames(keggres\$lesser).