# class 14 RNA-seq analysis mini-project

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load up DESeq2 to use for today's lab and download the files we will use

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 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: GenomeInfoDb

Warning: package 'GenomeInfoDb' was built under R version 4.3.2

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, rowWeightedMedians, rowWeightedMedians, rowWeightedMedians, 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
  metaFile <- "GSE37704_metadata.csv"</pre>
  countFile <- "GSE37704_featurecounts.csv"</pre>
  # Import metadata and take a peak
  colData = read.csv(metaFile, row.names=1)
  head(colData)
              condition
SRR493366 control_sirna
SRR493367 control_sirna
SRR493368 control_sirna
SRR493369
               hoxa1_kd
               hoxa1 kd
SRR493370
               hoxa1_kd
SRR493371
  # Import countdata
  countData = read.csv(countFile, row.names=1)
  head(countData)
```

	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
	SRR4933	371				
ENSG00000186092		0				
ENSG00000279928		0				
ENSG00000279457		46				
ENSG00000278566		0				
ENSG00000273547		0				
ENSG00000187634	2	258				

Remember that we need the countData and colData files to match up so we will need to remove that odd first column in countData namely contData\$length.

Q1. Complete the code below to remove the troublesome first column from countData

```
# 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

Now let's get rid of the zero entities as we have no data for these.

Q2. Complete the code below to filter countData to exclude genes (i.e. rows) where we have 0 read count across all samples (i.e. columns).

```
# Filter count data where you have 0 read count across all samples.
countData = countData[rowSums(countData)>0, ]
head(countData)
```

	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

## ##Running DESeq2

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

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

Next, get results for the HoxA1 knockdown versus control siRNA (remember that these were labeled as "hoxa1\_kd" and "control\_sirna" in our original colData metaFile input to DESeq, you can check this above and by running resultsNames(dds) command)

```
res = results(dds, contrast=c("condition", "hoxa1_kd", "control_sirna"))
```

Q3. Call the summary() function on your results to get a sense of how many genes are up or down-regulated at the default 0.1 p-value cutoff.

```
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)

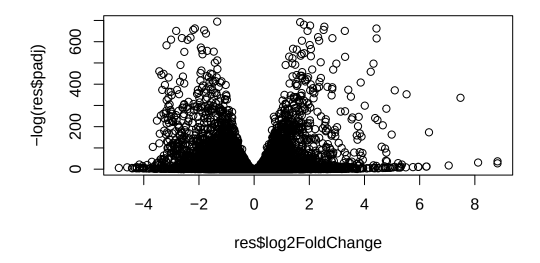
[1] see 'cooksCutoff' argument of ?results

[2] see 'independentFiltering' argument of ?results
```

#### Volcono plot

Now we will make a volcano plot, a commonly produced visualization from this type of data that we introduced last day.

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



Q4. Improve this plot by completing the below code, which adds color and axis labels

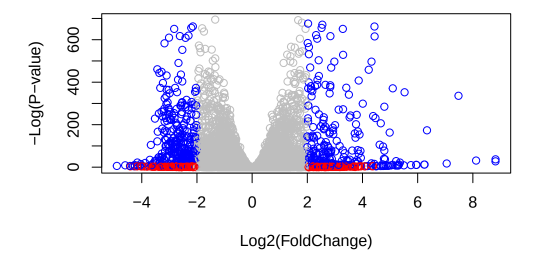
```
# Make a color vector for all genes
mycols <- rep("gray", nrow(res) )

# 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$padj<0.01) & (abs(res$log2FoldChange) > 2 )
mycols[ inds ] <- "blue"

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



## ##Adding gene annotation

Since we mapped and counted against the Ensembl annotation, our results only have information about Ensembl gene IDs. However, our pathway analysis downstream will use KEGG pathways, and genes in KEGG pathways are annotated with Entrez gene IDs.

Q5. Use the mapIDs() function multiple times to add SYMBOL, ENTREZID and GENE-NAME annotation to our results by completing the code below.

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

Loading required package: AnnotationDbi

Warning: package 'AnnotationDbi' was built under R version 4.3.2

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

```
[1] "ACCNUM"
                    "ALIAS"
                                    "ENSEMBL"
                                                   "ENSEMBLPROT"
                                                                  "ENSEMBLTRANS"
 [6] "ENTREZID"
                    "ENZYME"
                                   "EVIDENCE"
                                                   "EVIDENCEALL"
                                                                  "GENENAME"
                    "GO"
                                                   "IPI"
[11] "GENETYPE"
                                   "GOALL"
                                                                  "MAP"
[16] "OMIM"
                    "ONTOLOGY"
                                   "ONTOLOGYALL"
                                                  "PATH"
                                                                  "PFAM"
[21] "PMID"
                                   "REFSEO"
                                                   "SYMBOL"
                                                                  "UCSCKG"
                    "PROSITE"
[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="ENTREZID",
                      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></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></numeric>	<character> <c< td=""><td>haracter&gt;</td><td><character></character></td><td></td></c<></character>	haracter>	<character></character>	
ENSG00000279457	6.86555e-01	NA	NA	NA	
ENSG00000187634	5.15718e-03	SAMD11	148398	148398	
ENSG00000188976	1.76549e-35	NOC2L	26155	26155	
ENSG00000187961	1.13413e-07	KLHL17	339451	339451	
ENSG00000187583	9.19031e-01	PLEKHN1	84069	84069	
ENSG00000187642	4.03379e-01	PERM1	84808	84808	
ENSG00000188290	1.30538e-24	HES4	57801	57801	
ENSG00000187608	2.37452e-02	ISG15	9636	9636	
ENSG00000188157	4.21963e-16	AGRN	375790	375790	
ENSG00000237330	NA	RNF223	401934	401934	

Q6. Finally for this section let's reorder these results by adjusted p-value and save them to a CSV file in your current project directory.

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

Now lets see how pathway analysis can help us make further sense out of this ranked list of differentially expressed genes.

```
##Section 2. Pathway Analysis
```

Here we are going to use the gage package for pathway analysis. Once we have a list of enriched pathways, we're going to use the pathview package to draw pathway diagrams, shading the molecules in the pathway by their degree of up/down-regulation.

### **#KEGG** pathways

The gageData package has pre-compiled databases mapping genes to KEGG pathways and GO terms for common organisms. kegg.sets.hs is a named list of 229 elements. Each element is a character vector of member gene Entrez IDs for a single KEGG pathway.

First we need to do our one time install of these required bioconductor packages. Now we can load the packages and setup the KEGG data-sets we need.

```
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`
          "1544" "1548" "1549" "1553" "7498" "9"
[1] "10"
$`hsa00983 Drug metabolism - other enzymes`
 [1] "10"
             "1066"
                       "10720" "10941"
                                        "151531" "1548"
                                                           "1549"
                                                                    "1551"
                                "1806"
                                        "1807"
 [9] "1553"
             "1576"
                       "1577"
                                                  "1890"
                                                           "221223" "2990"
[17] "3251"
                                "3704"
             "3614"
                       "3615"
                                        "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"
                                                                 "79799"
                                                       "7498"
                                                                           "83549"
[49] "8824"
                         "9"
                                   "978"
               "8833"
$`hsa00230 Purine metabolism`
  [1] "100"
                "10201"
                          "10606"
                                                        "10623"
                                                                  "107"
                                                                            "10714"
                                    "10621"
                                              "10622"
  [9] "108"
                "10846"
                          "109"
                                    "111"
                                              "11128"
                                                        "11164"
                                                                  "112"
                                                                            "113"
 [17] "114"
                "115"
                          "122481" "122622" "124583" "132"
                                                                  "158"
                                                                            "159"
 [25] "1633"
                "171568" "1716"
                                    "196883"
                                              "203"
                                                        "204"
                                                                  "205"
                                                                            "221823"
                                                                            "270"
 [33] "2272"
                "22978"
                          "23649"
                                    "246721" "25885"
                                                        "2618"
                                                                  "26289"
 [41] "271"
                "27115"
                          "272"
                                    "2766"
                                              "2977"
                                                        "2982"
                                                                  "2983"
                                                                            "2984"
 [49] "2986"
                "2987"
                          "29922"
                                    "3000"
                                              "30833"
                                                        "30834"
                                                                  "318"
                                                                            "3251"
                          "3615"
                                    "3704"
                                              "377841"
                                                        "471"
                                                                  "4830"
                                                                            "4831"
 [57] "353"
                "3614"
 [65] "4832"
                          "4860"
                                    "4881"
                                              "4882"
                                                        "4907"
                                                                  "50484"
                                                                            "50940"
                "4833"
                                                                  "5139"
 [73] "51082"
                "51251"
                          "51292"
                                    "5136"
                                              "5137"
                                                        "5138"
                                                                            "5140"
 [81] "5141"
                "5142"
                          "5143"
                                    "5144"
                                              "5145"
                                                        "5146"
                                                                  "5147"
                                                                            "5148"
 [89] "5149"
                "5150"
                          "5151"
                                    "5152"
                                              "5153"
                                                        "5158"
                                                                  "5167"
                                                                            "5169"
 [97] "51728"
                "5198"
                          "5236"
                                    "5313"
                                              "5315"
                                                        "53343"
                                                                  "54107"
                                                                            "5422"
[105] "5424"
                "5425"
                          "5426"
                                    "5427"
                                              "5430"
                                                        "5431"
                                                                  "5432"
                                                                            "5433"
[113] "5434"
                "5435"
                          "5436"
                                    "5437"
                                              "5438"
                                                        "5439"
                                                                  "5440"
                                                                            "5441"
                                    "5557"
[121] "5471"
                "548644" "55276"
                                              "5558"
                                                        "55703"
                                                                  "55811"
                                                                            "55821"
[129] "5631"
                                    "56953"
                                                        "57804"
                                                                            "6240"
                "5634"
                          "56655"
                                              "56985"
                                                                  "58497"
[137] "6241"
                "64425"
                          "646625"
                                    "654364"
                                              "661"
                                                        "7498"
                                                                  "8382"
                                                                            "84172"
[145] "84265"
                "84284"
                          "84618"
                                    "8622"
                                              "8654"
                                                        "87178"
                                                                  "8833"
                                                                            "9060"
[153] "9061"
                "93034"
                          "953"
                                    "9533"
                                              "954"
                                                        "955"
                                                                  "956"
                                                                            "957"
[161] "9583"
                "9615"
```

we used the mapIDs() function above to obtain Entrez gene IDs (stored in res\$entrez) and we have the fold change results from DESeq2 analysis (stored in res\$log2FoldChange).

3.201955 -2.313738 -2.059631 -1.888019 -1.649792

2317

```
foldchanges = res$log2FoldChange
names(foldchanges) = res$entrez
head(foldchanges)
1266 54855 1465 51232 2034
```

Now, let's run the gage pathway analysis.

-2.422719

```
# 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	p.val
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
hsa04110	Cell cycle	0.001448312	121 8	.995727e-06
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

Each keggres\$less and keggres\$greater object is data matrix with gene sets as rows sorted by p-value. 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/trinitylee/Documents/BIMM143/class 14

Info: Writing image file hsa04110.pathview.png

You can play with the other input arguments to pathview() to change the display in various ways including generating a PDF graph. For example:

```
# A different PDF based output of the same data pathview(gene.data=foldchanges, pathway.id="hsa04110", kegg.native=FALSE)
```

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

Warning: reconcile groups sharing member nodes!

```
[,1] [,2]
[1,] "9" "300"
[2,] "9" "306"
```

Info: Working in directory /Users/trinitylee/Documents/BIMM143/class 14

Info: Writing image file hsa04110.pathview.pdf

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 pathview() function. We'll use these KEGG pathway IDs for pathview plotting below.

```
## 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/trinitylee/Documents/BIMM143/class 14

Info: Writing image file hsa04640.pathview.png

```
'select()' returned 1:1 mapping between keys and columns
Info: Working in directory /Users/trinitylee/Documents/BIMM143/class 14
Info: Writing image file hsa04630.pathview.png
'select()' returned 1:1 mapping between keys and columns
Info: Working in directory /Users/trinitylee/Documents/BIMM143/class 14
Info: Writing image file hsa00140.pathview.png
'select()' returned 1:1 mapping between keys and columns
Info: Working in directory /Users/trinitylee/Documents/BIMM143/class 14
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/trinitylee/Documents/BIMM143/class 14
Info: Writing image file hsa04330.pathview.png
Q7. Can you do the same procedure as above to plot the pathview figures for the top 5
down-reguled pathways?
  ## Focus on top 5 upregulated pathways here for demo purposes only
  keggrespathways <- rownames(keggres$less)[1:5]</pre>
  # Extract the 8 character long IDs part of each string
  keggresids = substr(keggrespathways, start=1, stop=8)
```

[1] "hsa04110" "hsa03030" "hsa03013" "hsa03440" "hsa04114"

keggresids

```
pathview(gene.data=foldchanges, pathway.id=keggresids, species="hsa")
'select()' returned 1:1 mapping between keys and columns
Info: Working in directory /Users/trinitylee/Documents/BIMM143/class 14
Info: Writing image file hsa04110.pathview.png
'select()' returned 1:1 mapping between keys and columns
Info: Working in directory /Users/trinitylee/Documents/BIMM143/class 14
Info: Writing image file hsa03030.pathview.png
'select()' returned 1:1 mapping between keys and columns
Info: Working in directory /Users/trinitylee/Documents/BIMM143/class 14
Info: Writing image file hsa03013.pathview.png
'select()' returned 1:1 mapping between keys and columns
Info: Working in directory /Users/trinitylee/Documents/BIMM143/class 14
Info: Writing image file hsa03440.pathview.png
'select()' returned 1:1 mapping between keys and columns
Info: Working in directory /Users/trinitylee/Documents/BIMM143/class 14
Info: Writing image file hsa04114.pathview.png
```

##Section 3. Gene Ontology (GO) We can also do a similar procedure with gene ontology. Similar to above, go.sets.hs has all GO terms. go.subs.hs is a named list containing indexes for the BP, CC, and MF ontologies. Let's focus on BP (a.k.a Biological Process) here.

```
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)
$greater
                                             p.geomean stat.mean
                                                                        p.val
GO:0007156 homophilic cell adhesion
                                          8.519724e-05 3.824205 8.519724e-05
GO:0002009 morphogenesis of an epithelium 1.396681e-04 3.653886 1.396681e-04
GO:0048729 tissue morphogenesis
                                          1.432451e-04 3.643242 1.432451e-04
GD:0007610 behavior
                                          1.925222e-04 3.565432 1.925222e-04
GO:0060562 epithelial tube morphogenesis 5.932837e-04 3.261376 5.932837e-04
GO:0035295 tube development
                                          5.953254e-04 3.253665 5.953254e-04
                                              q.val set.size
                                                                     exp1
                                                         113 8.519724e-05
GO:0007156 homophilic cell adhesion
                                          0.1952430
GO:0002009 morphogenesis of an epithelium 0.1952430
                                                         339 1.396681e-04
GO:0048729 tissue morphogenesis
                                          0.1952430
                                                         424 1.432451e-04
GO:0007610 behavior
                                                         426 1.925222e-04
                                          0.1968058
GO:0060562 epithelial tube morphogenesis 0.3566193
                                                         257 5.932837e-04
GO:0035295 tube development
                                                         391 5.953254e-04
                                          0.3566193
$less
                                            p.geomean stat.mean
                                                                       p.val
GO:0048285 organelle fission
                                         1.536227e-15 -8.063910 1.536227e-15
GO:0000280 nuclear division
                                         4.286961e-15 -7.939217 4.286961e-15
GO:0007067 mitosis
                                         4.286961e-15 -7.939217 4.286961e-15
GO:0000087 M phase of mitotic cell cycle 1.169934e-14 -7.797496 1.169934e-14
GO:0007059 chromosome segregation
                                         2.028624e-11 -6.878340 2.028624e-11
GO:0000236 mitotic prometaphase
                                         1.729553e-10 -6.695966 1.729553e-10
                                                q.val set.size
                                                                       exp1
GO:0048285 organelle fission
                                         5.843127e-12
                                                           376 1.536227e-15
GO:0000280 nuclear division
                                         5.843127e-12
                                                           352 4.286961e-15
GD:0007067 mitosis
                                                           352 4.286961e-15
                                         5.843127e-12
GO:0000087 M phase of mitotic cell cycle 1.195965e-11
                                                           362 1.169934e-14
GO:0007059 chromosome segregation
                                                           142 2.028624e-11
                                        1.659009e-08
```

1.178690e-07

84 1.729553e-10

GO:0000236 mitotic prometaphase

#### \$stats

##Section 4. Reactome Analysis

Reactome is database consisting of biological molecules and their relation to pathways and processes First, Using R, output the list of significant genes at the 0.05 level as a plain text file:

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

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

```
write.table(sig_genes, file="significant_genes.txt", row.names=FALSE, col.names=FALSE, quo
```

Q8: What pathway has the most significant "Entities p-value"? Do the most significant pathways listed match your previous KEGG results? What factors could cause differences between the two methods? cell cycle mitotic has the most significant "Entities p-value" which matches our KEGG results of M phase having the most significant. Both methods use different databases that could include different amounts of data and coverage of pathways.

```
##Section 5. GO online (OPTIONAL)
```

Gene Set Gene Ontology (GO) Enrichment is a method to determine over-represented or under-represented GO terms for a given set of genes. To perform Gene Set GO Enrichment online go to the website "http://www.geneontology.org/page/go-enrichment-analysis". Paste your significant gene list from section 4. Then, select "biological process" and "homo sapiens", and click submit.

Q9: What pathway has the most significant "Entities p-value"? Do the most significant pathways listed match your previous KEGG results? What factors could cause differences between the two methods?

no significant p value results are shown from the GO online database. This could be due to the incorrect entering of information or the fact that the online database not have enough coverage of the genes to provide any information.