Class 14: RNASeq Mini Project

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Run a complete RNASeq analysis workflow from counts to enriched genesets...

Data import

```
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

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,
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

counts <- read.csv("GSE37704_featurecounts.csv", row.names = 1)
metadata <- read.csv("GSE37704_metadata.csv")</pre>

Data Exploration

head(counts, 3)

	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
	SRR4933	371				
ENSG00000186092		0				
ENSG00000279928		0				
ENSG00000279457		46				

```
head(metadata)
         id
                condition
1 SRR493366 control_sirna
2 SRR493367 control_sirna
3 SRR493368 control_sirna
4 SRR493369
                 hoxa1_kd
                 hoxa1_kd
5 SRR493370
6 SRR493371
                 hoxa1_kd
  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
SRR493371
               hoxa1_kd
Check if the colData and countData match up
  metadata$id
[1] "SRR493366" "SRR493367" "SRR493368" "SRR493369" "SRR493370" "SRR493371"
  #RM first column
  countdata <- counts[, -1]</pre>
```

Now check if the metadata and count

all(colnames(countdata) == metadata\$id)

[1] TRUE

We need to remove all the zero count genes.

```
head(countdata, 3)
```

	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

To identify these zero count genes we can sum across the rows and check if the sum is more than zero.

```
to.keep.inds <- rowSums(countdata) > 0
nonzerocounts <- countdata[to.keep.inds,]</pre>
```

DESeq setup and analysis

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

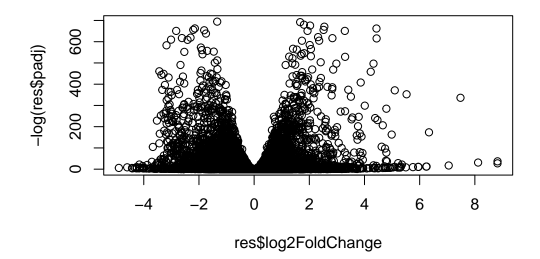
```
dds <- DESeq(dds)
using pre-existing size factors
estimating dispersions
found already estimated dispersions, replacing these
gene-wise dispersion estimates
mean-dispersion relationship
final dispersion estimates
fitting model and testing
  dds
class: DESeqDataSet
dim: 15975 6
metadata(1): version
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(3): id condition sizeFactor
```

Result extraction

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

```
Wald test p-value: condition hoxal 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
                               0.5428105 0.5215598 1.040744 2.97994e-01
ENSG00000187642
                  11.9798
                       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
  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
  plot( res$log2FoldChange, -log(res$padj) )
```

log2 fold change (MLE): condition hoxa1 kd vs control sirna



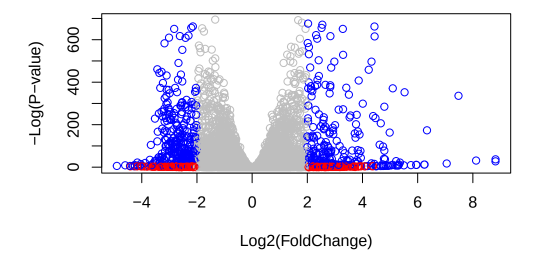
```
# Make a color vector for all genes
mycols <- rep("gray", nrow(res) )

# 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>
```



Gene annotation

```
library("AnnotationDbi")
  library("org.Hs.eg.db")
  columns(org.Hs.eg.db)
 [1] "ACCNUM"
                                    "ENSEMBL"
                     "ALIAS"
                                                    "ENSEMBLPROT"
                                                                    "ENSEMBLTRANS"
 [6] "ENTREZID"
                                    "EVIDENCE"
                     "ENZYME"
                                                    "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>
                  29.913579
                                 0.1792571 0.3248216
ENSG00000279457
ENSG00000187634 183.229650
                                 0.4264571 0.1402658
```

ENSG00000237330	0.158192	0.7859	552 4.0804729	0.192614 8.47261e-01
	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(gage)
```

```
library(gageData)
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.

```
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"
                                             "4882"
 [65] "4832"
                "4833"
                         "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"
Now setup the fold-change "importance" vector required for gage()
  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
  # 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
hsa03013 RNA transport
```

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

p.val

```
pathview(gene.data=foldchanges, pathway.id="hsa04110")
```

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

Info: Working in directory /Users/jaimychung/Downloads/R studio/class 14

Info: Writing image file hsa04110.pathview.png

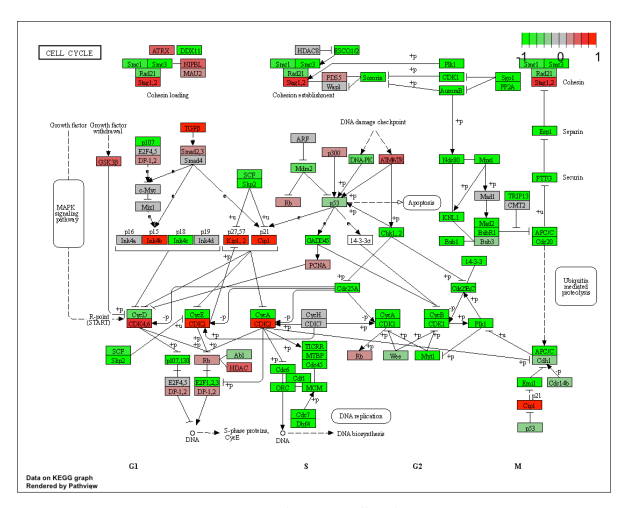


Figure 1: hsa04110 cell cycle

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/jaimychung/Downloads/R studio/class 14
Info: Writing image file hsa04110.pathview.pdf
  ## Focus on top 5 upregulated pathways here for demo purposes only
  keggrespathways <- rownames(keggres$greater)[1:5]</pre>
  # Extract the 8 character long IDs part of each string
  keggresids = substr(keggrespathways, start=1, stop=8)
  keggresids
[1] "hsa04640" "hsa04630" "hsa00140" "hsa04142" "hsa04330"
  pathview(gene.data=foldchanges, pathway.id=keggresids, species="hsa")
'select()' returned 1:1 mapping between keys and columns
Info: Working in directory /Users/jaimychung/Downloads/R studio/class 14
Info: Writing image file hsa04640.pathview.png
'select()' returned 1:1 mapping between keys and columns
Info: Working in directory /Users/jaimychung/Downloads/R studio/class 14
Info: Writing image file hsa04630.pathview.png
'select()' returned 1:1 mapping between keys and columns
```

```
Info: Working in directory /Users/jaimychung/Downloads/R studio/class 14
Info: Writing image file hsa00140.pathview.png
'select()' returned 1:1 mapping between keys and columns
Info: Working in directory /Users/jaimychung/Downloads/R studio/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/jaimychung/Downloads/R studio/class 14
Info: Writing image file hsa04330.pathview.png
  ## Focus on top 5 upregulated pathways here for demo purposes only
  keggrespathwaysless <- rownames(keggres$less)[1:5]</pre>
  # Extract the 8 character long IDs part of each string
  keggresidsless = substr(keggrespathwaysless, start=1, stop=8)
  keggresidsless
[1] "hsa04110" "hsa03030" "hsa03013" "hsa03440" "hsa04114"
  pathview(gene.data=foldchanges, pathway.id=keggresidsless, species="hsa")
'select()' returned 1:1 mapping between keys and columns
Info: Working in directory /Users/jaimychung/Downloads/R studio/class 14
Info: Writing image file hsa04110.pathview.png
'select()' returned 1:1 mapping between keys and columns
```

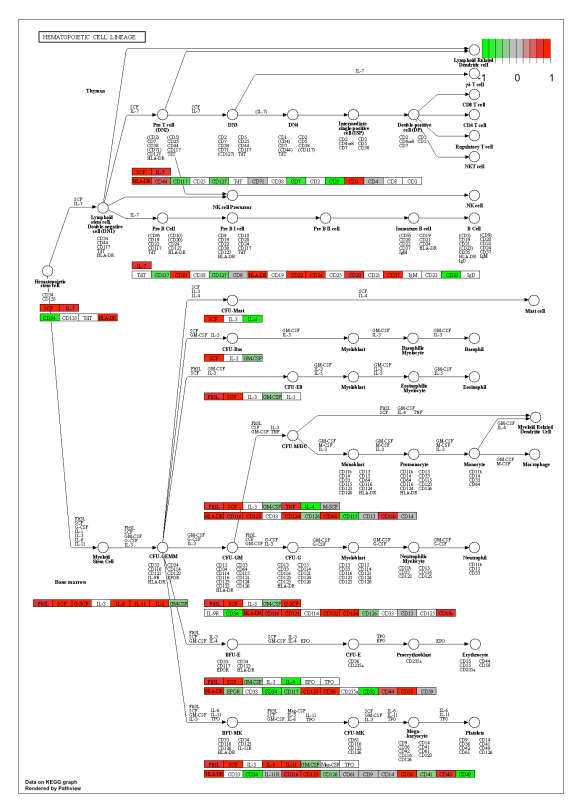


Figure 2: The KEGG cell-cycle pathway

```
Info: Working in directory /Users/jaimychung/Downloads/R studio/class 14

Info: Writing image file hsa03030.pathview.png

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

Info: Working in directory /Users/jaimychung/Downloads/R studio/class 14

Info: Writing image file hsa03013.pathview.png

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

Info: Working in directory /Users/jaimychung/Downloads/R studio/class 14

Info: Writing image file hsa03440.pathview.png

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

Info: Working in directory /Users/jaimychung/Downloads/R studio/class 14

Info: Working in directory /Users/jaimychung/Downloads/R studio/class 14

Info: Writing image file hsa04114.pathview.png
```

Using Gene Ontology (GO)

```
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)
```

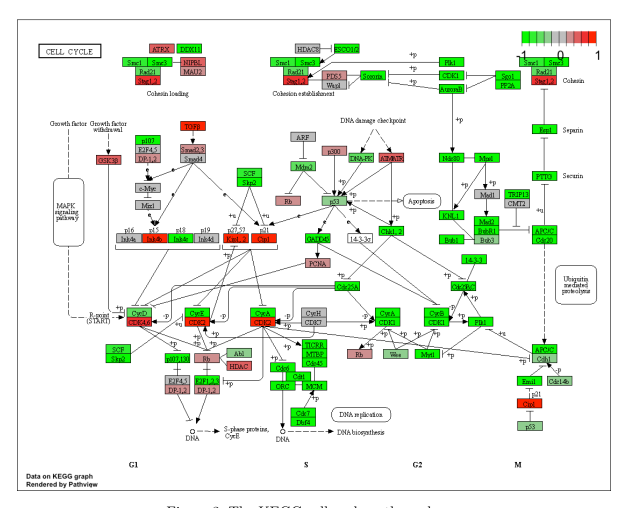


Figure 3: The KEGG cell-cycle pathway lesser $\,$

\$greater

	=
	4205 8.519724e-05
1.396681e-04 3.65	3886 1.396681e-04
1.432451e-04 3.64	3242 1.432451e-04
1.925222e-04 3.56	5432 1.925222e-04
5.932837e-04 3.26	1376 5.932837e-04
5.953254e-04 3.25	3665 5.953254e-04
q.val set.size	exp1
0.1952430 113	8.519724e-05
0.1952430 339	1.396681e-04
0.1952430 424	1.432451e-04
0.1968058 426	1.925222e-04
0.3566193 257	5.932837e-04
0.3566193 391	5.953254e-04
p.geomean stat.m	ean p.val
1.536227e-15 -8.063	910 1.536227e-15
4.286961e-15 -7.939	217 4.286961e-15
4.286961e-15 -7.939	217 4.286961e-15
1.169934e-14 -7.797	496 1.169934e-14
2.028624e-11 -6.878	340 2.028624e-11
1.729553e-10 -6.695	966 1.729553e-10
q.val set.si	ze exp1
5.843127e-12 3	76 1.536227e-15
5.843127e-12 3	52 4.286961e-15
5.843127e-12 3	52 4.286961e-15
1.195965e-11 3	62 1.169934e-14
1.659009e-08	42 2.028624e-11
1.178690e-07	84 1.729553e-10
stat.mean exp1	
3.824205 3.824205	
a 3.653886 3.653886	
3.643242 3.643242	
3.565432 3.565432	
3.261376 3.261376	
3.253665 3.253665	
r	n 1.396681e-04 3.65 1.432451e-04 3.64 1.925222e-04 3.56 5.932837e-04 3.26 5.953254e-04 3.25 q.val set.size 0.1952430 113 n 0.1952430 424 0.1968058 426 0.3566193 257 0.3566193 391 p.geomean stat.m 1.536227e-15 -8.063 4.286961e-15 -7.939 4.286961e-15 -7.939 1.169934e-14 -7.797 2.028624e-11 -6.878 1.729553e-10 -6.695 q.val set.si: 5.843127e-12 3 5.843127e-12 3 5.843127e-12 3 5.843127e-12 3 1.195965e-11 3 1.659009e-08 1 1.178690e-07 stat.mean exp1 3.824205 3.824205 n 3.653886 3.653886 3.643242 3.643242 3.565432 3.565432 3.261376 3.261376

Reactome Analysis

```
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"

write.table(sig_genes, file="significant_genes.txt", row.names=FALSE, col.names=FALSE, quenes)</pre>
```