maketitle

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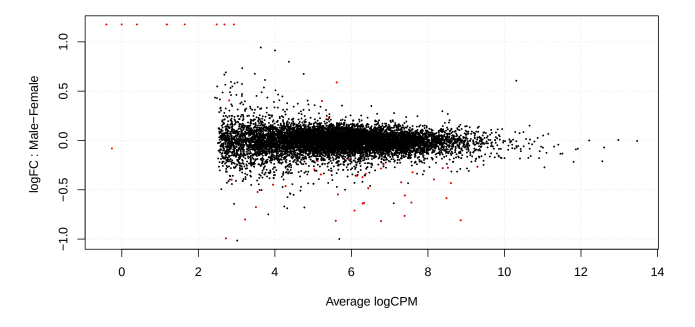
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```
library(DEGreport)
data(humanSexDEedgeR)
library(edgeR)
```

## 1 QC figures from DE analysis

We are going to do a differential expression analysis with edgeR. We have an object that is comming from the edgeR package. It countains a gene count matrix for 85 TSI HapMap individuals, and the gender information. With that, we are going to apply the 'glmFit' function to get genes differentially expressed between males and females.

```
des<-humanSexDEedgeR$design
fit <- glmFit(humanSexDEedgeR,des)
lrt <- glmLRT(fit)
tab<-cbind(lrt$table,p.adjust(lrt$table$PValue,method="BH"))
detags <- rownames(tab[tab[,5]<=0.1,])
plotSmear(humanSexDEedgeR, de.tags=detags)</pre>
```



We need to extract the experiment design data.frame where the condition is Male or Female.

```
counts<-cpm(humanSexDEedgeR,log=FALSE)
g1<-colnames(counts)[1:41]
g2<-colnames(counts)[42:85]
design<-data.frame(condition=sub("1","Male",sub("0","Female",des[,2])))</pre>
```

We are getting the chromosome information for each gene. This way we can colour genes according autosomic,X or Y chromosomes.

```
data(geneInfo)
```

Create the report. The main parameters are the column names in group1, and group2. Then, the count matrix, gene

names that are DE, p-values, fold changes and path to create the report. As optional, you can give colours for each gene, and the number of permutation.

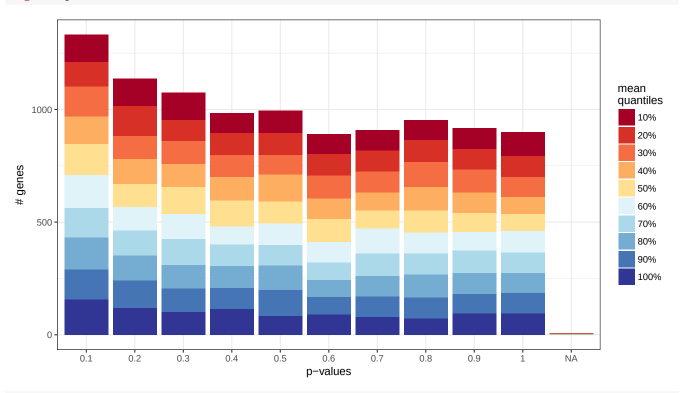
```
detag10<-detags[1:10]
pval<-tab[,4]
fc<-tab[detag10,1]</pre>
```

Run the fowlling lines if you want to visualize your expression values by condition:

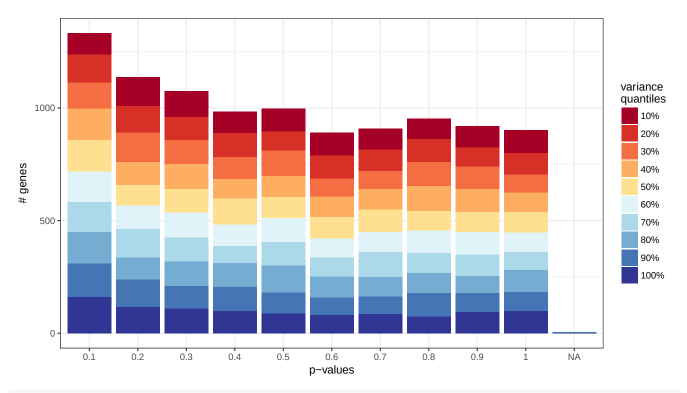
```
degObj(counts,design,"degObj.rda")
library(shiny)
runGist(9930881)
```

You can use individual functions, like degRank or degMean. This will create specific figures and tables that are included in the report.

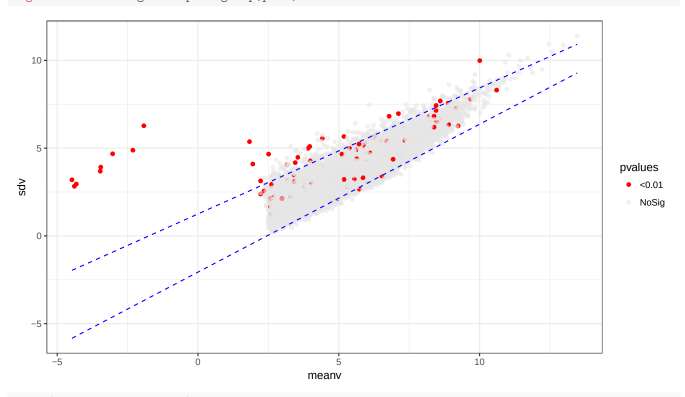
degMean(pval,counts)



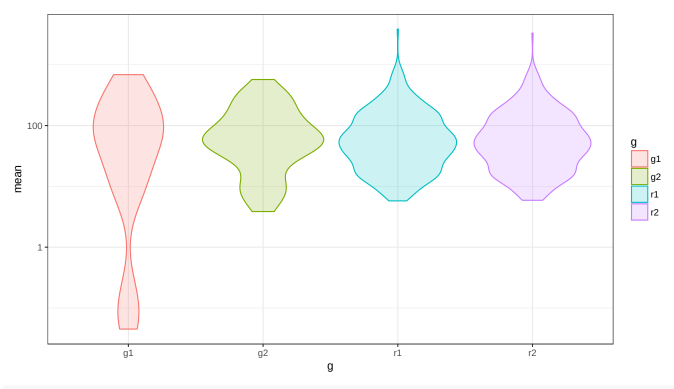
degVar(pval,counts)



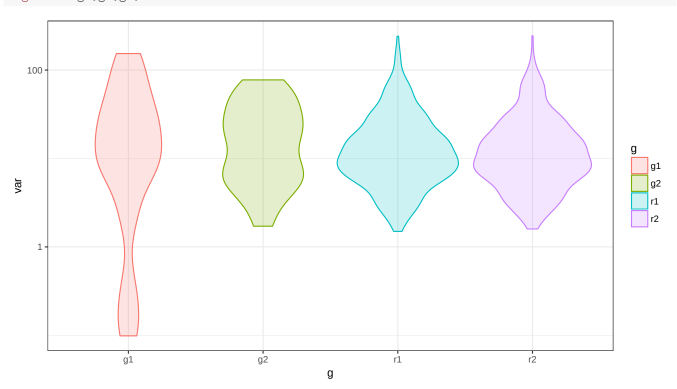
degMV(humanSexDEedgeR\$samples\$group,pval,counts)



degMB(detags,g1,g2,counts)



## degVB(detags,g1,g2,counts)

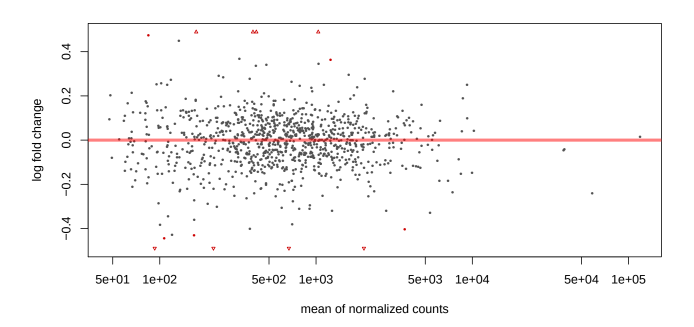


<sup>#</sup> require(rjags)
# rank<-degRank(g1,g2,counts[detag10,],fc,400,500)
# degPR(rank)</pre>

## 2 Report from DESeq2 analysis

In this section, we show how to use DESeq2 output to create a full report, including figures and tbale with top deregulated genes, GO enrichment analysis and heatmaps and PCA plots. If you set path\_results, different files will be saved there.

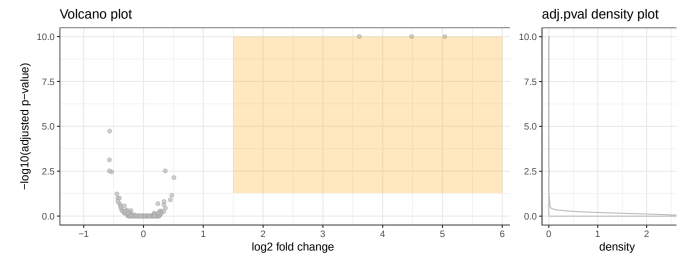
```
data(humanSexDEedgeR)
library(DESeq2)
idx <- c(1:10, 75:85)
dse <- DESeqDataSetFromMatrix(humanSexDEedgeR$counts[1:1000, idx],</pre>
humanSexDEedgeR$samples[idx,], design=~group)
dse <- DESeq(dse)</pre>
res <- degResults(dds=dse, name="test", org=NULL,</pre>
do_go=FALSE, group="group", xs="group", path_results = NULL)
## ## Comparison: test {.tabset}
##
##
    <br>out of 1000 with nonzero total read count<br/>djusted p-value < 0.1<br/>br>LFC > 0 (up)
##
                                                                                                     : 6, 0.6%
##
##
## Differential expression file at: test_de.csv
##
## Normalized counts matrix file at: test_log2_counts.csv
##
## ### MA plot plot
```



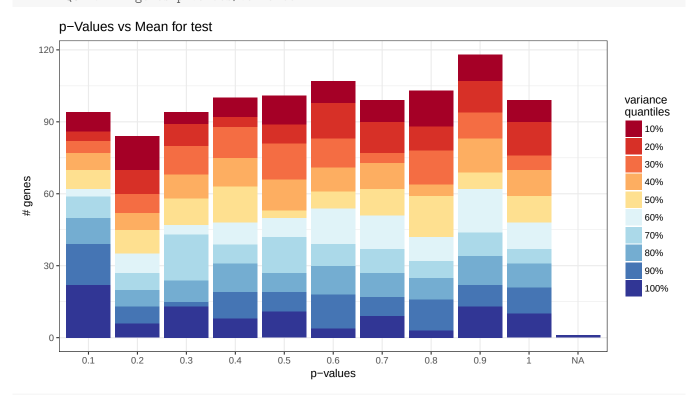
```
##
##
## Volcano plot
```



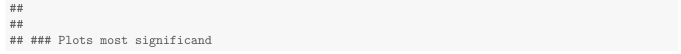


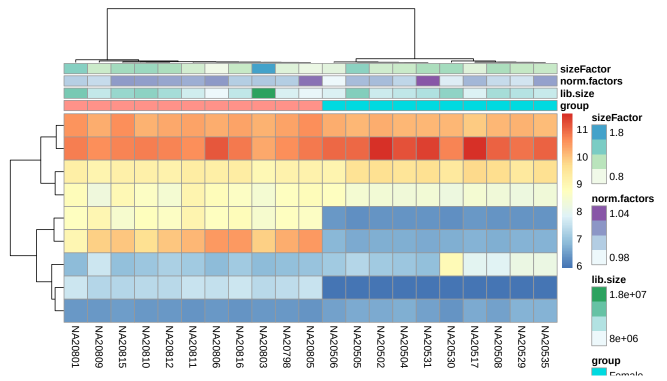


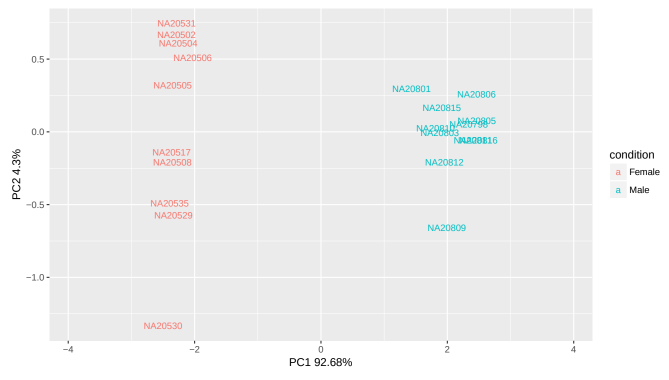
##
##
## QC for DE genes p-values/variance



##
##
## Most significand, FDR< 0.05 and log2FC > 0.1 : 9







## ##

```
##
## Plot top 9 genes
    ENSG00000012817
                                      ENSG00000067646
                                                                          ENSG00000067048
                                                    £ 14 ± 1
                                                              treatment
  5.0
                                                                       count
                                                                                                     Female
  2.5
    ENSG00000005889
                                       ENSG00000006757
                                                                           ENSG00000005020
                                                                        10.8
 10.0
                                     7.0
                                                              treatment
count
                                                                                                     Female
                                                                        10.2
  9.0
                                     6.2
                                                                                        Male
   ENSG00000073282
                                       ENSG00000005302
                                                                          ENSG00000003400
                                     12.0
                                                              treatment
count
                                                                Female
                                     10.5
##
##
##
##
   ### Top DE genes
##
##
## \begin{tabular}{||r|r|r|r|r|r}
     & baseMean & log2FoldChange & lfcSE & stat & pvalue & padj & absMaxLog2FC\\
##
## ENSG00000012817 & 414.71514 & 5.0383666 & 0.1280628 & 39.342935 & 0.0000000 & 0.0000000 & 5.0383666
## \hline
## ENSG00000067646 & 171.03009 & 4.4844515 & 0.1304760 & 34.369939 & 0.0000000 & 0.0000000 & 4.4844515\\
## ENSG00000067048 & 1032.64158 & 3.6088039 & 0.1418177 & 25.446777 & 0.0000000 & 0.0000000 & 3.6088039\\
  \hline
## ENSG00000005889 & 670.30349 & -0.5654992 & 0.1050602 & -5.382623 & 0.0000001 & 0.0000184 & 0.5654992\\
## ENSG00000006757 & 92.82614 & -0.5687093 & 0.1228764 & -4.628304 & 0.0000037 & 0.0007373 & 0.5687093\\
## \hline
## ENSG00000005020 & 1238.12238 & 0.3638754 & 0.0855898 & 4.251386 & 0.0000212 & 0.0030350 & 0.3638754
## ENSG00000073282 & 220.50620 & -0.5703537 & 0.1336781 & -4.266619 & 0.0000198 & 0.0030350 & 0.5703537\\
## \hline
## ENSG00000005302 & 2024.85620 & -0.5313921 & 0.1267987 & -4.190831 & 0.0000278 & 0.0034742 & 0.5313921\\
  \hline
```

## ENSG00000003400 & 394.82940 & 0.5090003 & 0.1273944 & 3.995469 & 0.0000646 & 0.0071740 & 0.5090003

```
## \hline
## ENSG00000069702 & 106.95162 & -0.4442940 & 0.1291537 & -3.440042 & 0.0005816 & 0.0581623 & 0.4442940\\
## \hline
## ENSG00000010278 & 84.64206 & 0.4736610 & 0.1406137 & 3.368527 & 0.0007557 & 0.0687009 & 0.4736610\\
## ENSG00000023171 & 165.99513 & -0.4299847 & 0.1332187 & -3.227661 & 0.0012481 & 0.0999395 & 0.4299847\\
## \hline
## ENSG00000072501 & 3694.13874 & -0.4022772 & 0.1250802 & -3.216153 & 0.0012992 & 0.0999395 & 0.4022772\\
## \hline
## ENSG00000059377 & 132.55552 & 0.4494421 & 0.1435071 & 3.131846 & 0.0017371 & 0.1240792 & 0.4494421\\
## \hline
## ENSG00000068079 & 1039.04270 & 0.3447312 & 0.1138155 & 3.028860 & 0.0024548 & 0.1534241 & 0.3447312
## ENSG0000070018 & 119.93653 & -0.4286051 & 0.1408198 & -3.043642 & 0.0023373 & 0.1534241 & 0.4286051\\
## \hline
## ENSG00000008277 & 378.03916 & -0.4001595 & 0.1376207 & -2.907699 & 0.0036410 & 0.2022771 & 0.4001595\\
## \hline
## ENSG00000012963 & 1836.75128 & 0.2397646 & 0.0821682 & 2.917972 & 0.0035232 & 0.2022771 & 0.2397646\\
## \hline
## ENSG00000005059 & 481.54751 & 0.3411158 & 0.1192221 & 2.861179 & 0.0042207 & 0.2221415 & 0.3411158\\
## ENSG00000038427 & 100.44612 & -0.3833305 & 0.1384660 & -2.768408 & 0.0056331 & 0.2682422 & 0.3833305\\
## \hline
## \end{tabular}
```

## 3 Detect patterns of expression

In this section, we show how to detect pattern of expression. Mainly useful when data is a time course experiment. degPatterns needs a expression matrix, the design experiment and the column used to group samples.

