

# Skeleton Key for RNAseq analysis

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*See README.md for more detailed instructions of how to use script*

## Analysis

### libraries

```
library(edgeR)
library(yaml)
```

### Read in YAML guide

```
yamls <- yaml.load_file("de.yaml")
```

This part assigns your YMAL to a object in R. This will be used throughout the script to specify which sample types you are comparing.

```
sample1 <- yamls$sample1
sample2 <- yamls$sample2
```

```
sample1
```

```
## [1] "tf2cmbr"
```

```
sample2
```

```
## [1] "wtcmbr"
```

### Read in Data

Read in raw count data per gene.

```
counts <- read.delim("../requisiteData/sam2countsResults.tsv",row.names=1)

#check the file
head(counts)
colnames(counts)
#need to convert NA to 0 counts
counts[is.na(counts)] <- 0
```

## Subset DE experiment

Start by subsetting the particular treatments which are being compared.

```
colnames(counts)
```

```
## [1] "tf2ambr1"      "tf2ambr3"      "tf2ambr4"      "tf2ambr6"
## [5] "tf2aother1"    "tf2aother2"    "tf2aother4"    "tf2aother7"
## [9] "tf2bmr2"       "tf2bmr5"       "tf2bmr6"       "tf2bmr1"
## [13] "tf2bmr3"       "tf2bmr4"       "tf2bmr6"       "tf2cmbr1.4"
## [17] "tf2cmbr3"      "tf2cmbr6"      "tf2cmbr7"      "tf2cother2"
## [21] "tf2cother5"    "tf2cother6"    "tf2cother7"    "wtambr2"
## [25] "wtambr4"       "wtambr5"       "wtambr1"       "wtambr5"
## [29] "wtambr6"       "wtambr7"       "wtambr8"       "wtbmr2"
## [33] "wtbmr3"        "wtbmr6"        "wtbmr8"        "wtbmr1.4"
## [37] "wtbmr3"        "wtbmr5"        "wtbmr8"        "wtcmbr10"
## [41] "wtcmbr1.4.6"   "wtcmbr2"       "wtcmbr3"       "wtcmbr7"
## [45] "wtcmbr9"       "wtcother1.3.4" "wtcother2"     "wtcother6"
```

```
counts1 <- counts[,grep(sample1, colnames(counts), value = TRUE)]
count1Len <- length(colnames(counts1)) #used in to specify library group in next step.
```

```
counts2 <- counts[,grep(sample2, colnames(counts), value = TRUE)]
count2Len <- length(colnames(counts2)) #used to specify library group in next step.
```

```
counts <- cbind(counts1, counts2)
```

```
head(counts)
```

```
##           tf2cmbr1.4 tf2cmbr3 tf2cmbr6 tf2cmbr7 wtcmb10
## Solyc00g005040.2.1      0      6      8      4      0
## Solyc00g005050.2.1      1     34     17     12      5
## Solyc00g005060.1.1      0      1      0      0      1
## Solyc00g005070.1.1     23     11      8      9      5
## Solyc00g005080.1.1     22      7      8     12      0
## Solyc00g005150.1.1      1      3      0      0      0
##           wtcmb1.4.6 wtcmb2 wtcmb3 wtcmb7 wtcmb9
## Solyc00g005040.2.1      9      3      1      0      0
## Solyc00g005050.2.1     38     21     11      4      7
## Solyc00g005060.1.1      3      0      0      1      0
## Solyc00g005070.1.1     12      7      4      6      1
## Solyc00g005080.1.1      7     19     45      4      7
## Solyc00g005150.1.1      1      3      3      2      1
```

## Add column specifying library Group

Make a vector called group that will be used to make a new column named group to identify library region type.

```
group <- c(rep(sample1, count1Len), rep(sample2, count2Len))
d <- DGEList(counts=counts,group=group)
```

Check to see if the group column matches your sample name and they are appropriate.

```
d$samples
```

```
##           group lib.size norm.factors
## tf2cmbr1.4  tf2cmbr  443572          1
## tf2cmbr3    tf2cmbr 1337575          1
## tf2cmbr6    tf2cmbr  790129          1
## tf2cmbr7    tf2cmbr  832907          1
## wtcnbr10     wtcnbr  459717          1
## wtcnbr1.4.6  wtcnbr 1158809          1
## wtcnbr2      wtcnbr 1130695          1
## wtcnbr3      wtcnbr 1560130          1
## wtcnbr7      wtcnbr  374882          1
## wtcnbr9      wtcnbr  386974          1
```

## Differential expression using edgeR

Make sure there is full understanding on each edgeR command being used. The manual is amazing so read it *before* running the DE analysis below [edgeR manual](#).

```
cpm.d <- cpm(d) #counts per mutant
d <- d[rowSums(cpm.d>5)>=3,] #This might be a line to adjust. It is removing genes with low counts.
d <- estimateCommonDisp(d,verbose=T)
```

```
## Disp = 0.3524 , BCV = 0.5936
```

```
d <- calcNormFactors(d)
d <- estimateCommonDisp(d)

DEtest <- exactTest(d,pair=c(sample1,sample2))
head(DEtest$table)
```

```
##           logFC logCPM  PValue
## Solyc00g005050.2.1 -0.01411  4.243 1.00000
## Solyc00g005070.1.1 -1.53465  4.024 0.01710
## Solyc00g005080.1.1 -0.59400  4.296 0.28619
## Solyc00g005440.1.1  0.38877  4.832 0.53700
## Solyc00g005840.2.1  0.36635  4.835 0.51212
## Solyc00g005880.1.1 -1.50325  3.183 0.03331
```

```
results <- topTags(DEtest, n=Inf)
head(results)
```

```
## Comparison of groups: wtcnbr-tf2cmbr
##           logFC logCPM  PValue      FDR
## Solyc02g023990.2.1 -5.920  6.429 1.417e-18 2.223e-14
## Solyc11g013430.1.1 -7.053  5.063 1.393e-16 1.093e-12
## Solyc01g056770.1.1 -6.343  5.055 1.634e-15 8.418e-12
## Solyc06g069460.1.1 -5.717  5.087 2.146e-15 8.418e-12
## Solyc07g044980.2.1 -4.743  7.744 7.227e-15 2.267e-11
## Solyc01g098190.2.1 -4.638  5.937 8.269e-14 2.162e-10
```

```
dim(results$table)
```

```
## [1] 15687      4
```

```
sum(results$table$FDR<.05) # How many are DE genes?
```

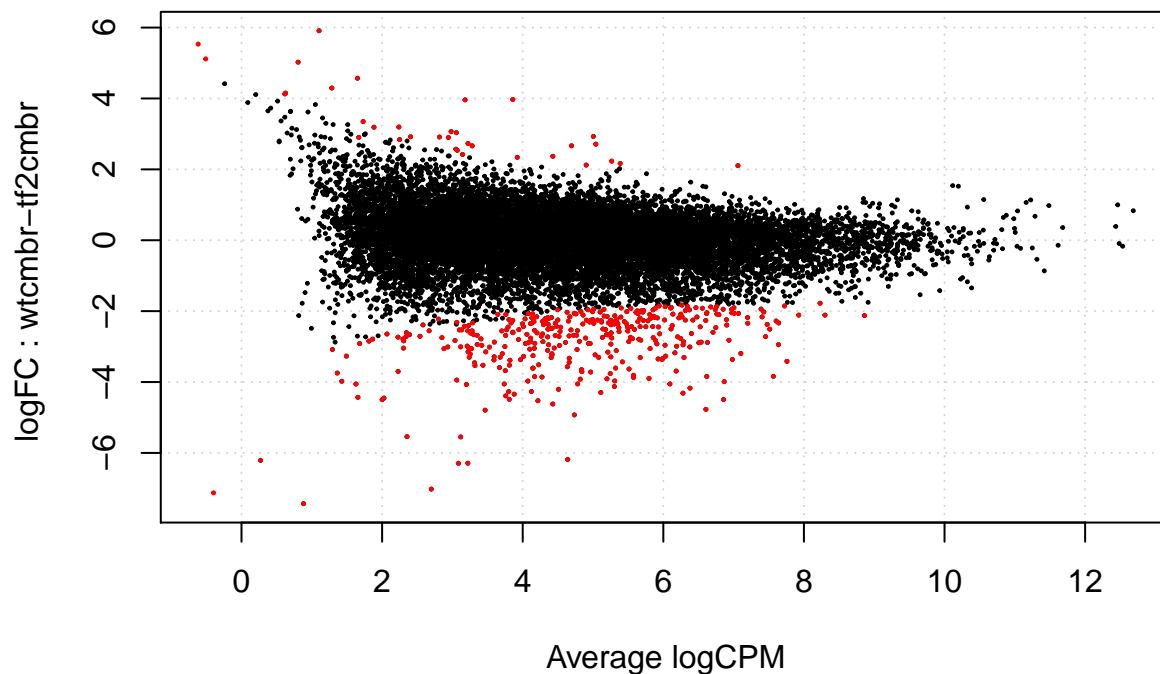
```
## [1] 401
```

```
summary(decideTestsDGE(DEtest,p.value=.05))
```

```
##      [,1]  
## -1    367  
##  0   15286  
##  1      34
```

```
sig.genes <- rownames(results$table[results$table$FDR<0.05,]) # outputs just significant gene names
```

```
plotSmea(d,de.tags=sig.genes)
```



Subset by all the genes with a significant FDR score.

```
results.sig <- subset(results$table, results$table$FDR < 0.05)
```

```
dim(results.sig)
```

What are the genes that are misexpressed? For this we need to add some annotation.

Essentially we are merging two annotations files to 1.) only sig genes 2.) all genes

```

annotation1<- read.delim("../requisiteData/ITAG2.3_all_Arabidopsis_ITAG_annotations.tsv", header=FALSE)
colnames(annotation1) <- c("ITAG", "SGN_annotation")
annotation2<- read.delim ("../requisiteData/ITAG2.3_all_Arabidopsis_annotated.tsv")
annotation <- merge(annotation1,annotation2, by = "ITAG")
head(annotation)

```

```

##          ITAG
## 1 Solyc00g005000.2.1
## 2 Solyc00g005040.2.1
## 3 Solyc00g005050.2.1
## 4 Solyc00g005080.1.1
## 5 Solyc00g005900.1.1
## 6 Solyc00g006490.2.1
##
## 1          Aspartic proteinase nepenthesin I (AHRD V1 ***-
## 2          Potassium channel (AHRD V1 ***- DOEM91_9ROSI
## 3
## 4
## 5 Oxygen-evolving enhancer protein 1, chloroplastic (AHRD V1 ***- PSBO_SOLTU); contains Interpro dom
## 6 Serine/threonine-protein phosphatase 6 regulatory subunit 3 (AHRD V1 ***- SAPS3_HUMAN); contain
##      AGI symbol
## 1 AT3G20015   <NA>
## 2 AT5G46240   KAT1
## 3 AT5G11680   <NA>
## 4 ATCG01280 YCF2.2
## 5 AT5G66570 MSP-1
## 6 AT1G07990   <NA>
##
## 1
## 2
## 3
## 4
## 5
## 6 SIT4 phosphatase-associated family protein; similar to SIT4 phosphatase-associated family protein
## X..identity alignment.length e.value bit.score percent.query.align
## 1      63.76          447 7e-148      520          89.94
## 2      66.02          103 2e-37       150          85.71
## 3      76.96          204 1e-88       322          98.98
## 4      91.25           80 2e-38       153          79.80
## 5      69.62           79 4e-26       112          78.79
## 6      61.92          856 0e+00       979          99.77

```

```

#Making the only significant gene table
results.sig$ITAG <- rownames(results.sig) #change row.names to ITAG for merging
results.sig.annotated <- merge(results.sig,annotation,by = "ITAG", all.x=TRUE) #This is merging to only

#Making all table

results$table$ITAG <- rownames(results$table)
results.all.annotated <- merge(results$table, annotation,by = "ITAG")

```

Write table with results.

```
write.table(results.all.annotated, file=paste(sample1,"_",sample2,"_", "DE_all.txt", sep=""), sep="\t", row
write.table(results.sig.annotated, file=paste(sample1,"_",sample2,"_", "DE_sig.txt", sep=""), sep="\t", row
```

Now run the script below for a full knitr report of what was run and leave this report in the folder that the analysis was done with output files.

```
library(rmarkdown)
render("skeletonDE.Rmd", "pdf_document", output_file = paste(sample1,"_",sample2,"_", "DE.pdf", sep=""))
```