Skeleton Key for RNAseq analysis

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About

sample2

This is the script used to perform differential gene expression analysis using edgeR. The tissue came from p4 leaves of *Solanum lycopersicum* using Laser Capture Microdissection.

Key to Samples

```
genotype: either wildtype of tf2 region: A. tip B. early emmerging leaflet C. base type: MBR = Marginal Blastozone Region, other = the rachis or midvein region
```

Run this first chunk before rendering knittedn document

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

```
library(edgeR)
library(yaml)
### Set Working Directory
rstudioapi::getActiveDocumentContext
## function ()
## {
       context <- callFun("getActiveDocumentContext")</pre>
##
##
       context$selection <- as.document_selection(context$selection)</pre>
##
       structure(context, class = "document_context")
## }
## <environment: namespace:rstudioapi>
setwd(dirname(rstudioapi::getActiveDocumentContext()$path))
### Read in YAML guide
## This reads in the information in the `de.yml` file which has the two names of the samples you are in
yamls <- yaml.load_file("de.yml")</pre>
## This part assigns your YMAL to a object in R. This will be used throughout the script to specify wh
sample1 <- yamls$sample1
sample2 <- yamls$sample2</pre>
sample1
## [1] "tf2ambr"
```

```
## [1] "tf2aother"
```

To make report

Run the render() function below and everything will be run with report at end.

```
library(rmarkdown)
render("skeletonDE_sept2017.Rmd", "pdf_document", output_file = paste(sample1,"_",sample2,"_","DE.pdf",
Samples:
print(sample1)
## [1] "tf2ambr"
print(sample2)
## [1] "tf2aother"
```

Analysis (Acutal start)

Read in Data

Read in raw count data per gene.

```
counts <- read.delim("../../data/Ciera_coveragebed_counts.txt", row.names = 1)

colnames(counts)
#need to convert NA to 0 counts
counts[is.na(counts)] <- 0

## Get rid of low count libraries "wtbother1.4", "wtbmbr8", "tf2ambr3"
counts <- counts[,-c(36,35,2)]</pre>
```

Subset DE expirement

Start by subsetting the particular treatments which are being compared. This might need to be modified depending on the naming of your samples. In my case each sample is named by sample and rep number, so the script is identifying any sample with the sample name given in the de.yml file.

```
colnames(counts)
```

```
##
    [1] "tf2ambr1"
                         "tf2ambr4"
                                          "tf2ambr6"
                                                           "tf2aother1"
                                          "tf2aother7"
                                                           "tf2bmbr2"
##
    [5] "tf2aother2"
                         "tf2aother4"
   [9] "tf2bmbr5"
                         "tf2bmbr6"
                                          "tf2bother1"
                                                           "tf2bother3"
## [13] "tf2bother4"
                                                           "tf2cmbr3"
                         "tf2bother6"
                                          "tf2cmbr1.4"
## [17] "tf2cmbr6"
                         "tf2cmbr7"
                                          "tf2cother2"
                                                           "tf2cother5"
                                                           "wtambr4"
## [21] "tf2cother6"
                         "tf2cother7"
                                          "wtambr2"
## [25] "wtambr5"
                         "wtaother1"
                                          "wtaother5"
                                                           "wtaother6"
                                                           "wtbmbr3"
## [29] "wtaother7"
                         "wtaother8"
                                          "wtbmbr2"
## [33] "wtbmbr6"
                         "wtbother3"
                                          "wtbother5"
                                                           "wtbother8"
## [37] "wtcmbr1.4.6"
                         "wtcmbr10"
                                          "wtcmbr2"
                                                           "wtcmbr3"
## [41] "wtcmbr7"
                         "wtcmbr9"
                                          "wtcother1.3.4" "wtcother2"
## [45] "wtcother6"
```

```
counts1 <- counts[,grep(sample1, colnames(counts), value = TRUE)]
count1Len <- length(colnames(counts1)) #used 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)</pre>
```

```
tf2ambr1 tf2ambr4 tf2ambr6 tf2aother1 tf2aother2 tf2aother4
##
## Solyc00g069887
                          3
                                    4
                                            37
                                                         4
                                                                     2
                                                                                20
## Solyc00g009145
                          4
                                   59
                                            11
                                                        78
                                                                     0
                                                                               181
## Solyc00g021530
                          4
                                    6
                                             3
                                                         2
                                                                     1
                                                                                 5
## Solyc00g023020
                          0
                                    4
                                             1
                                                         2
                                                                     0
                                                                                 5
## Solyc00g024690
                                    5
                                                         2
                                                                     0
                                                                                 4
                          1
                                             1
## Solyc00g042147
                          0
                                             0
                                                         0
                                                                     0
                                                                                 1
##
                   tf2aother7
## Solyc00g069887
                            4
## Solyc00g009145
                            9
## Solyc00g021530
                            1
## Solyc00g023020
                            0
## Solyc00g024690
                            0
## Solyc00g042147
                            0
```

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)</pre>
```

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

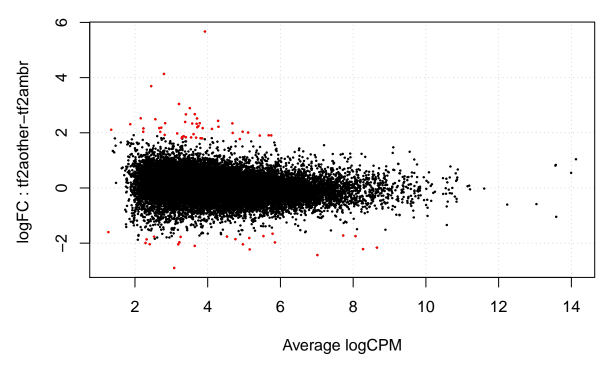
d\$samples

```
##
                  group lib.size norm.factors
                tf2ambr 6930595
## tf2ambr1
## tf2ambr4
                tf2ambr 8370884
                                            1
## tf2ambr6
                tf2ambr 5418262
                                            1
## tf2aother1 tf2aother 6091106
                                            1
## tf2aother2 tf2aother 4378292
                                            1
## tf2aother4 tf2aother 4749874
                                            1
## tf2aother7 tf2aother 1291681
                                            1
```

Differential expression using edgeR

Make sure there is full understanding of each edgeR command being used. The manual is amazing so read it before running the DE analysis below edgeR manual. There are many options and they must be set to be appropriate for your analysis.

```
cpm.d <- cpm(d) #counts per million</pre>
d \leftarrow 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)</pre>
## Disp = 0.16139 , BCV = 0.4017
d <- calcNormFactors(d)</pre>
d <- estimateCommonDisp(d)</pre>
DEtest <- exactTest(d, pair = c(sample1, sample2))</pre>
head(DEtest$table)
##
                        logFC
                                logCPM
                                              PValue
## Solyc00g009145 2.1727118 3.283642 1.584417e-05
## Solyc00g107055 -0.8660841 5.322385 5.848731e-02
## Solyc00g143770 -0.3685400 2.706547 5.882683e-01
## Solyc00g183555 0.5924603 4.897258 1.998687e-01
## Solyc00g212265 -0.0264010 4.923861 9.165576e-01
## Solyc00g005050 -0.6927157 3.403202 1.401791e-01
results <- topTags(DEtest, n = Inf)
dim(results$table)
## [1] 22820
sum(results$table$FDR < .05) # How many are DE genes?</pre>
## [1] 70
summary(decideTestsDGE(DEtest,p.value = .05))
##
      [,1]
## -1
## 0 22750
## 1
sig.genes <- rownames(results$table[results$table$FDR < 0.05,]) # outputs just significant gene names
plotSmear(d,de.tags = sig.genes)
```



Subset all the genes with a significant FDR score less than .05.

```
results.sig <- subset(results$table, results$table$FDR < 0.05)
dim(results.sig)</pre>
```

```
## [1] 70 4
```

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

## Remove the trailing
annotation$ITAG <- gsub("^(.*)[.].*", "\\1", annotation$ITAG)
annotation$ITAG <- gsub("^(.*)[.].*", "\\1", annotation$ITAG)

#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 only

#Making all table
results$table$ITAG <- rownames(results$table)
results.all.annotated <- merge(results$table, annotation,by = "ITAG")</pre>
```

Write table with results.

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 outp	ut files														