## **GO** Enrichment

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

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

#### Read in YAML guide

```
library(yaml)
yamls <- yaml.load_file("de.yml")
sample1 <- yamls$sample2
sample2 <- yamls$sample2

## [1] "tf2cmbr"
sample2

## [1] "wtcmbr"
library(goseq)
library(GO.db)</pre>
```

# Setting up the DE table for GO analysis

# File Input

Input the output from DE analysis. This is made for a list that includes only the significant genes.

```
sigOnly <- read.table(paste(sample1,"_",sample2,"_DE_sig.txt", sep=""), header = TRUE, fill = TRUE)
sigOnly$logFC <- as.numeric(as.character(sigOnly$logFC))
colnames(sigOnly)[1] <- "itag"</pre>
```

#### Subset

First I need to subset the list to up or down regulated, then add a new colum that specififys 1. This column is need to for merging.

```
upITAG <- subset(sigOnly, logFC > 0, select = c(itag))
upITAG$up <- 1

downITAG <- subset(sigOnly, logFC < 0, select = c(itag))
downITAG$down <- 1

allITAG <- subset(sigOnly, select = c(itag))
allITAG$all <- 1</pre>
```

### Merge I - with normalized ITAG length gene list

read in guide.

```
geneLength <- read.csv("../normalized_genes_length.csv")
head(geneLength)</pre>
```

```
##
                    itag length
## 1 Solyc00g005040.2.1
                            357
## 2 Solyc00g005050.2.1
                            588
## 3 Solyc00g005060.1.1
                            273
## 4 Solyc00g005070.1.1
                             81
## 5 Solyc00g005080.1.1
                            297
## 6 Solyc00g005150.1.1
                           1143
#isolate just the gene list
genes <- subset(geneLength, select = c(itag))</pre>
```

First merge each table to geneLength

```
upITAGmerge <- merge(genes, upITAG, by = "itag", all= TRUE)
downITAGmerge <- merge(genes, downITAG, by = "itag", all= TRUE)
allITAGmerge <- merge(genes, allITAG, by = "itag", all= TRUE)</pre>
```

### Merge II - Merge them all together.

```
matrixGOupdown <- merge(upITAGmerge, downITAGmerge, by = "itag", all = TRUE)
matrixGOupdownall <- merge(matrixGOupdown, allITAG, by = "itag", all = TRUE)
matrixGO <- merge(matrixGOupdownall, geneLength, by = "itag", all = TRUE)</pre>
```

## Clean Up

```
matrixGO[is.na(matrixGO)] <- 0
head(matrixGO)</pre>
```

```
##
                  itag up down all length
## 1 Solyc00g005040.2.1 0
                                 0
                                      357
## 2 Solyc00g005050.2.1 0
                                 0
                                      588
                             0
## 3 Solyc00g005060.1.1 0
                                      273
## 4 Solyc00g005070.1.1 0
                                       81
                             0
                                0
## 5 Solyc00g005080.1.1 0
                             0
                                0
                                      297
## 6 Solyc00g005150.1.1 0
                                     1143
```

This is if you want to write out the table of the GO matrix. #write.table(matrixGO, "mydata.txt", sep="\hat{v}, quote= FALSE)

### GO enrichment

The is the input of the GOslim categories. There are only two columns 1. itag and 2. go

```
pat <- matrixGO
head(pat)
##
                   itag up down all length
## 1 Solyc00g005040.2.1 0
## 2 Solyc00g005050.2.1 0
                                  0
                                       588
                              0
## 3 Solyc00g005060.1.1 0
                              0
                                  0
                                       273
## 4 Solyc00g005070.1.1 0
                                 0
                                        81
## 5 Solyc00g005080.1.1 0
                              0
                                  0
                                       297
## 6 Solyc00g005150.1.1 0
                              0
                                  0
                                      1143
cate <- read.table("../melted.GOTable.txt",header=TRUE)</pre>
head(cate)
##
                   itag
## 1 Solyc00g005000.2.1 GO:0006508
## 2 Solyc00g005040.2.1 GO:0005774
## 3 Solyc00g005050.2.1 GD:0005829
## 4 Solyc00g005080.1.1 GO:0005524
## 5 Solyc00g005130.1.1 GD:0006508
## 6 Solyc00g005150.1.1 GO:0003676
```

### Subseting for GO analysis

Specify the column you are interested in pat\$all refers to all the DE gene regardless if they are up or down regulated. If you want to specify down regulated, specify pat\$down. I am going to put this into a loop, where each time the loop goes thought it will perform GO enrichment on all three types of lists of significant genes and them write them to a table.

```
sigType <- c("up", "down", "all")
for(type in sigType) {
   genes = as.integer(pat[,type])
   names(genes) = pat$itag
   table(genes)
length(genes)

pwf = nullp(genes,bias.data=pat$length)

GO.wall = goseq(pwf,gene2cat = cate)
head(GO.wall)

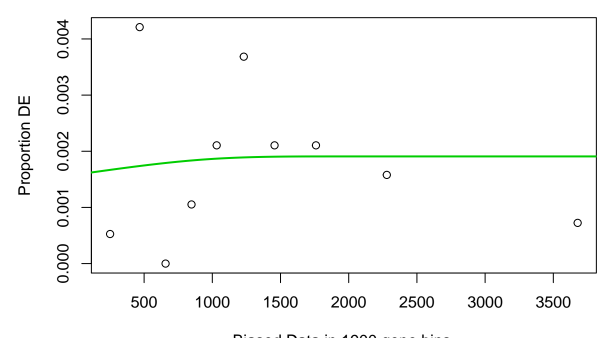
#This is going to correct for multiple testing. You can specify the p-value cut-off of GO categories y
enriched.GO = GO.wall$category[p.adjust(GO.wall$over_represented_pvalue, method = "BH") < 0.05]
enriched.GO</pre>
```

```
my.GO <- as.character(enriched.GO)
my.GO.table <- Term(my.GO)
my.GO.table
t <- as.matrix(my.GO.table)

print(type) #this is for the knitr document
print(t) #this is for the knitr document
write.table(t, file=paste(sample1,"_",sample2,"DE1_sigonly_",type,"_GO.txt", sep=""))
}</pre>
```

```
## Using manually entered categories.
## For 2940 genes we could not find any cat
```

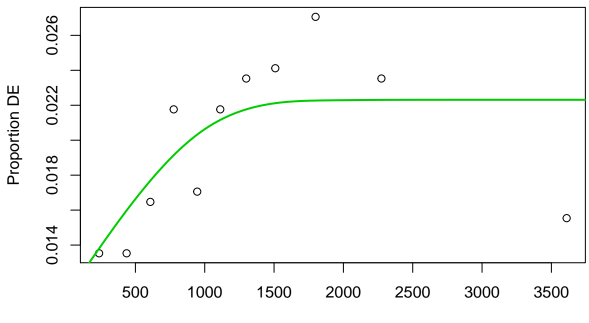
- ## For 2940 genes, we could not find any categories. These genes will be excluded.
- ## To force their use, please run with use\_genes\_without\_cat=T (see documentation).
- ## This was the default behavior for version 1.15.1 and earlier.
- ## Calculating the p-values...



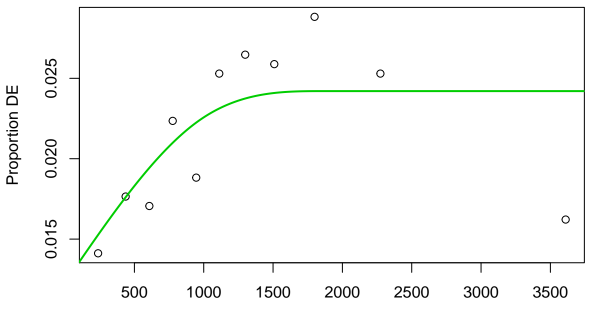
Biased Data in 1900 gene bins.

```
## [1] "up"
## [,1]

## Using manually entered categories.
## For 2940 genes, we could not find any categories. These genes will be excluded.
## To force their use, please run with use_genes_without_cat=T (see documentation).
## This was the default behavior for version 1.15.1 and earlier.
## Calculating the p-values...
```



Biased Data in 1700 gene bins.



Biased Data in 1700 gene bins.