### **GO** Enrichment

```
## Libraries
library(tidyr)
library(goseq)
library(GO.db)
library(yaml)
library(rmarkdown)
## Read in YAML guide
### 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()$path))
## Read in sample names from yaml
yamls <- yaml.load_file("de.yml")</pre>
sample1 <- yamls$sample1
sample2 <- yamls$sample2</pre>
sample1
## [1] "wtambr"
sample2
## [1] "wtbmbr"
## Render
render("GO_basedOnSkeletonGO.Rmd", "pdf_document", output_file = paste(sample1,"_",sample2,"_","GO.pdf"
```

# 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("../../../requisiteData/data_06Sept2017/", sample1,"_",sample2,"_DE_sig.
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("../../../requisiteData/normalized_genes_length.csv")

## remove trailing numbers in ITAG
geneLength$itag <- gsub("^(.*)[.].*", "\\1", geneLength$itag)
geneLength$itag <- gsub("^(.*)[.].*", "\\1", geneLength$itag)

#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</pre>
head(matrixGO)
##
              itag up down all length
## 1 Solyc00g005040 0
                        0
                          0
                                357
## 2 Solyc00g005050 0
                                588
                        0
                          0
## 3 Solyc00g005060 0 0 0
                                273
## 4 Solyc00g005070 0 0
                                81
## 5 Solyc00g005080 0
                        0 0
                                297
## 6 Solyc00g005150 0 0
                          0 1143
## This is if you want to write out the table of the GO matrix.
\# write.table(matrixGO, "mydata.txt", sep="\t", 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
                           0
                               0
                                    357
                     0
## 2 Solyc00g005050
                                    588
                           0
## 3 Solyc00g005060
                           0
                               0
                                    273
                     0
## 4 Solyc00g005070
                     0
                           0
                               0
                                     81
## 5 Solyc00g005080
                           0
                               0
                                    297
## 6 Solyc00g005150
                           0
                               0
                                   1143
## New GO table
cate <- read.table("../../../requisiteData/ITAG3.2_protein_go.tsv")</pre>
colnames(cate) <- c("itag", "go")</pre>
summary(cate$itag)
## Solyc01g111990.3.1 Solyc02g079630.2.1 Solyc02g071260.3.1
##
  Solyc03g083440.3.1 Solyc03g097290.3.1 Solyc10g044670.2.1
##
##
##
   Solyc11g065920.2.1
                      Solyc11g071610.2.1 Solyc11g071620.3.1
##
  Solyc12g008890.2.1
                      Solyc01g009235.1.1 Solyc01g059870.4.1
##
  Solyc01g080460.3.1 Solyc01g088170.4.1 Solyc01g112290.3.1
##
##
##
  Solyc04g014210.3.1 Solyc04g016430.3.1 Solyc04g076620.3.1
##
  Solyc04g080820.2.1 Solyc05g053410.3.1 Solyc06g019170.3.1
##
##
  Solyc07g008880.3.1 Solyc08g043170.3.1 Solyc09g011930.3.1
##
##
##
  Solyc09g015240.1.1 Solyc10g017990.2.1 Solyc11g010310.2.1
##
  Solyc11g040180.2.1 Solyc11g068830.2.1 Solyc11g071580.2.1
##
##
##
   Solyc11g071600.2.1 Solyc11g072140.2.1 Solyc12g008900.2.1
##
  Solyc01g080810.3.1 Solyc01g088200.3.1 Solyc01g090710.3.1
##
##
  Solyc01g102410.3.1 Solyc01g103960.3.1 Solyc01g109540.3.1
##
##
  Solyc02g063490.3.1 Solyc02g067930.3.1 Solyc02g068490.3.1
##
##
  Solyc02g093300.3.1 Solyc03g118640.3.1 Solyc04g054890.3.1
##
##
## Solyc05g009220.3.1 Solyc05g014720.3.1 Solyc07g017990.3.1
##
  Solyc07g045480.3.1 Solyc07g062650.3.1 Solyc07g063770.3.1
##
##
```

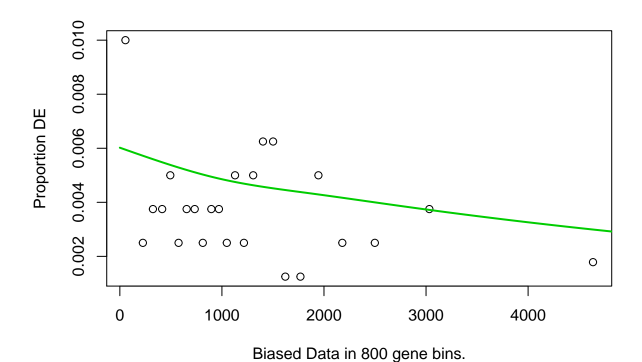
```
## Solyc07g064810.3.1 Solyc08g007420.3.1 Solyc08g061920.2.1
##
   Solyc08g061930.3.1 Solyc08g078390.3.1 Solyc08g078400.3.1
##
##
##
   Solyc08g078850.3.1 Solyc09g014710.3.1 Solyc09g014720.2.1
##
                                                            6
   Solyc09g014730.3.1 Solyc09g014740.3.1 Solyc09g074990.3.1
##
##
##
   Solyc09g090140.3.1 Solyc10g006710.3.1 Solyc10g079470.3.1
##
##
   Solyc10g079870.2.1 Solyc11g005630.1.1 Solyc11g012140.2.1
##
   Solyc11g013810.2.1 Solyc11g065930.2.1 Solyc12g007170.2.1
##
##
  Solyc12g014180.2.1 Solyc12g019110.2.1 Solyc12g056940.2.1
##
##
  Solyc00g026860.1.1 Solyc00g042130.2.1 Solyc00g055960.1.1
##
##
  Solyc01g006190.3.1 Solyc01g006520.3.1 Solyc01g008330.3.1
##
##
##
   Solyc01g073730.3.1 Solyc01g074010.3.1 Solyc01g088150.3.1
##
  Solyc01g088160.3.1 Solyc01g088230.3.1 Solyc01g088310.3.1
##
##
                                                            5
   Solyc01g091480.3.1 Solyc01g094500.3.1 Solyc01g094835.1.1
##
##
   Solyc01g096020.3.1 Solyc01g096900.3.1 Solyc01g099620.3.1
##
##
                                                            5
   Solyc01g102370.3.1 Solyc01g106480.3.1 Solyc01g106770.3.1
##
##
                                                            5
##
   Solyc02g022930.3.1 Solyc02g038740.3.1 Solyc02g062430.3.1
##
##
              (Other)
                31698
##
## remove the trailing num in itag id
cate$itag <- gsub("^(.*)[.].*", "\\1", cate$itag)</pre>
cate$itag <- gsub("^(.*)[.].*", "\\1", cate$itag)
cate <- separate(data = cate, col = go, into = c("go1", "go2", "go4", "go5", "go6", "go7", "go8", "go9"
## Warning: Too few values at 32311 locations: 1, 2, 3, 4, 5, 6, 7, 8, 9, 10,
## 11, 12, 13, 14, 15, 16, 17, 18, 19, 20, ...
cate <- gather(cate, itag, go1:go9, factor_key = TRUE)</pre>
colnames(cate)[3] <- "go"</pre>
## First remove rows with NA in go
cate <- cate[complete.cases(cate), ]</pre>
## Now every go term and itag pair is represented only once, so we can get rid of itag.1 column
cate \leftarrow cate[,-2]
head(cate)
               itag
## 1 Solyc00g005000 GD:0004190
```

```
## 2 Solyc00g005285 G0:0008168
## 3 Solyc00g005285 G0:0008171
## 4 Solyc00g005440 G0:0003723
## 5 Solyc00g005460 G0:0003723
## 6 Solyc00g005840 G0:0045454
```

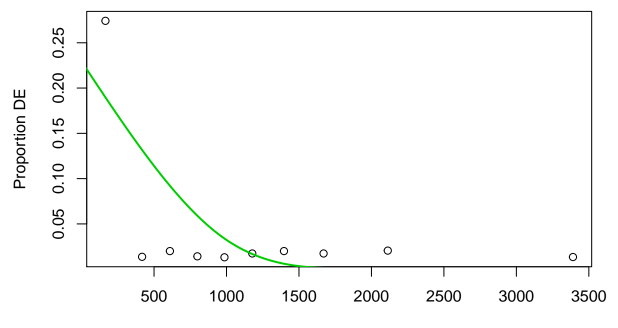
# Subseting for GO analysis

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")</pre>
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
  my.GO <- as.character(enriched.GO)
  my.GO.table <- Term(my.GO)</pre>
  mv.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 = ""))
write.table(GO.wall, file = paste(sample1,"_",sample2,"DE1_sigValues_",type,"_GO.txt", sep = ""))
}
## Using manually entered categories.
## For 7168 genes, we could not find any categories. These genes will be excluded.
## To force their use, please run with use_genes_without_cat=TRUE (see documentation).
## This was the default behavior for version 1.15.1 and earlier.
## Calculating the p-values...
## 'select()' returned 1:1 mapping between keys and columns
```

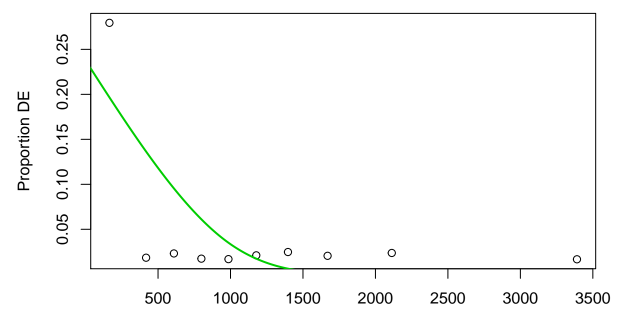


- ## Using manually entered categories.
- ## For 7168 genes, we could not find any categories. These genes will be excluded.
- ## To force their use, please run with use\_genes\_without\_cat=TRUE (see documentation).
- $\mbox{\tt \#\#}$  This was the default behavior for version 1.15.1 and earlier.
- ## Calculating the p-values...
- ## 'select()' returned 1:1 mapping between keys and columns



Biased Data in 1900 gene bins.

```
## [1] "down"
##
              [,1]
## GO:0004672 "protein kinase activity"
## GO:0006468 "protein phosphorylation"
## GO:0006508 "proteolysis"
## GO:0005524 "ATP binding"
## GO:0055085 "transmembrane transport"
## GO:0015299 "solute:proton antiporter activity"
## GO:0006812 "cation transport"
## GO:0004177 "aminopeptidase activity"
## GO:0008152 "metabolic process"
## Using manually entered categories.
## For 7168 genes, we could not find any categories. These genes will be excluded.
## To force their use, please run with use_genes_without_cat=TRUE (see documentation).
## This was the default behavior for version 1.15.1 and earlier.
## Calculating the p-values...
## 'select()' returned 1:1 mapping between keys and columns
```



Biased Data in 1900 gene bins.