Transcriptome Demo

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2024-05-09

Load required packages

1

2

3

4

5

dnaA

dnaN recF

gyrB

lptA

```
library(ballgown)
library(RColorBrewer)
library(genefilter)
library(dplyr)
library(devtools)
#Organizes data into a dataframe that has two columns. #The first column contains the ids while the
second specifies the corresponding stage, either planktonic or biofilm.
pheno_data<-data.frame(ids = c("plank01", "plank02", "biofilm01", "biofilm02"),</pre>
                        stage = c("planktonic", "planktonic", "biofilm", "biofilm"))
#Creates a Ballgown object and check transcript number #Organizes the file path into a vector
samples.c <- paste('ballgown', pheno_data$ids, sep = '/')</pre>
bg <- ballgown(samples = samples.c, meas='all', pData = pheno_data)
bg
## ballgown instance with 5737 transcripts and 4 samples
#Filters out transcripts with a small variance. #Only transcripts with a variance larger than 1 are kept.
bg_filt = subset(bg,"rowVars(texpr(bg)) >1",genomesubset=TRUE)
bg_filt
## ballgown instance with 5163 transcripts and 4 samples
#Creates a table of transcripts
results_transcripts<- stattest(bg_filt, feature = "transcript", covariate = "stage",
getFC = TRUE, meas = "FPKM")
results_transcripts <- data.frame(geneNames=geneNames(bg_filt),
transcriptNames=transcriptNames(bg_filt), results_transcripts)
#Shows the top of the table stored in results_transcripts
head(results_transcripts)
     geneNames transcriptNames
##
                                    feature id
                                                           fс
                                                                    pval
```

gene-PA0001 transcript 1 5.247471e+01 0.3048003 0.9471885 gene-PA0002 transcript 2 1.745401e+01 0.1001167 0.9471885

gene-PA0003 transcript 3 5.229990e-01 0.8960742 0.9845954

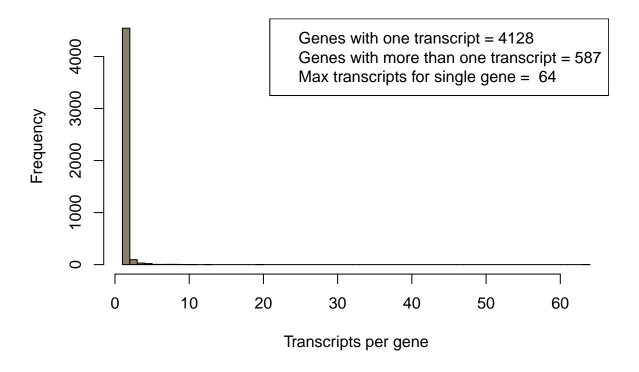
gene-PA0004 transcript 4 4.834298e+10 0.2743082 0.9471885

gene-PA0005 transcript 5 8.951420e+00 0.2455336 0.9471885

```
## 6
                    gene-PA0006 transcript 6 3.264697e+02 0.2949859 0.9471885
#Choose a transcript to examine more closely
results_transcripts[results_transcripts$transcriptNames == "gene-PA0004", ] #I chose to examine gene-PA
##
     geneNames transcriptNames
                                    feature id
                                                         fc
                                                                  pval
                                                                             qval
## 4
                    gene-PA0004 transcript 4 48342981060 0.2743082 0.9471885
          gyrB
##This transcript is for the gene gyrB. ##The fold difference between planktonic and biofilm stages is
48342981060 ##The corrected q-value of 0.9471885 indicates that the difference is not significant.
#Filters out non-significant transcripts. #Only results with a significant p-value (less than 0.05) are kept
#The dim function gives the number of transcripts kept.
sigdiff <- results_transcripts %>% filter(pval<0.05)</pre>
dim(sigdiff)
## [1] 207
#Organizes the table by p-value and fold change (fc). #The p-values go from smallest to biggest (since
decreasing=FALSE) #The fold change is in increasing order since the negative sign in front and setting
decreasing=FALSE.
o = order(sigdiff[,"pval"], -abs(sigdiff[,"fc"]), decreasing=FALSE)
output = sigdiff[o,c("geneNames", "transcriptNames", "id", "fc", "pval", "qval")]
write.table(output, file="SigDiff.txt", sep="\t", row.names=FALSE, quote=FALSE)
head(output)
##
        geneNames transcriptNames
                                                                pval
                                                                           qval
                                      id
                                                    fc
                       gene-PA3992 4091 9.886091e+01 0.0003032315 0.9471885
## 4091
## 4958
                       gene-PA4804 4958 3.563696e-04 0.0006661432 0.9471885
## 2745
                       gene-PA2690 2745 5.783390e-02 0.0014192618 0.9471885
## 2896
                       gene-PA2832 2896 1.786570e+03 0.0023414834 0.9471885
               tpm
                       gene-PA0365 370 3.964652e-07 0.0023906201 0.9471885
## 370
## 3129
                       gene-PA3059 3129 1.687425e-03 0.0025838457 0.9471885
             pelF
#Loads gene names
bg table = texpr(bg filt, 'all')
bg_gene_names = unique(bg_table[, 9:10])
#Pulls out gene expression data and visualize
gene_expression = as.data.frame(gexpr(bg_filt))
head(gene_expression)
              FPKM.plank01 FPKM.plank02 FPKM.biofilm01 FPKM.biofilm02
##
                                                                 2.685373
## .
                   1.198359
                                0.9103059
                                                 2.526183
## MSTRG.1
                 405.892761 400.8589780
                                               232.324417
                                                               181.932617
## MSTRG.10
                  89.649139
                              78.5762100
                                                35.010487
                                                                59.757320
## MSTRG.100
                 116.443428 106.2109530
                                                92.206810
                                                                95.322479
## MSTRG.1000
                   7.833186
                                5.5019700
                                                15.717344
                                                                42.342495
## MSTRG.1001
                   6.845010
                                4.7381980
                                                38.199095
                                                                89.078876
#Renames the columns to the names specified within c("")
colnames(gene_expression) <- c("plank01", "plank02", "biofilm01", "biofilm02")</pre>
head(gene_expression)
##
                  plank01
                              plank02 biofilm01 biofilm02
```

```
## .
                1.198359
                           0.9103059
                                       2.526183
                                                  2.685373
## MSTRG.1
              405.892761 400.8589780 232.324417 181.932617
## MSTRG.10 89.649139 78.5762100 35.010487 59.757320
## MSTRG.100 116.443428 106.2109530 92.206810 95.322479
## MSTRG.1000
                7.833186
                           5.5019700 15.717344 42.342495
## MSTRG.1001
                6.845010
                           4.7381980 38.199095 89.078876
dim(gene_expression)
## [1] 4592
#Loads the transcript to gene table and determine the number of transcripts and unique genes
transcript_gene_table = indexes(bg)$t2g
head(transcript_gene_table)
    t_id
            g_id
## 1
       1 MSTRG.1
## 2
       2 MSTRG.2
## 3
       3 MSTRG.3
       4 MSTRG.3
## 4
## 5
       5 MSTRG.4
## 6
       6 MSTRG.5
length(row.names(transcript_gene_table))
## [1] 5737
length(unique(transcript_gene_table[, "g_id"]))
## [1] 4715
#Plots the number of transcripts per gene
counts=table(transcript_gene_table[, "g_id"])
c_one = length(which(counts == 1))
c_more_than_one = length(which(counts > 1))
c_{max} = max(counts)
hist(counts, breaks=50, col="bisque4", xlab="Transcripts per gene",
main="Distribution of transcript count per gene")
legend_text = c(paste("Genes with one transcript =", c_one),
paste("Genes with more than one transcript =", c_more_than_one),
paste("Max transcripts for single gene = ", c_max))
legend("topright", legend_text, lty=NULL)
```

Distribution of transcript count per gene

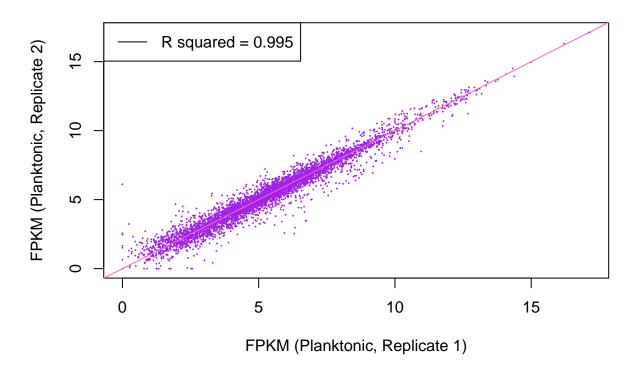


##A vast majority of the genes have a single transcript. ##Few genes have multiple transcripts ##The most transcripts for a single gene were 64.

#Creates a plot of how similar the two planktonic replicates are to one another.

```
x = gene_expression[,"plank01"]
y = gene_expression[,"plank02"]
min_nonzero=1
plot(x=log2(x+min_nonzero), y=log2(y+min_nonzero), pch=16, col="purple", cex=0.25,
xlab="FPKM (Planktonic, Replicate 1)", ylab="FPKM (Planktonic, Replicate 2)",
main="Comparison of expression values for a pair of replicates")
abline(a=0,b=1, col = "hotpink")
rs=cor(x,y)^2
legend("topleft", paste("R squared = ", round(rs, digits=3), sep=""), lwd=1, col="black")
```

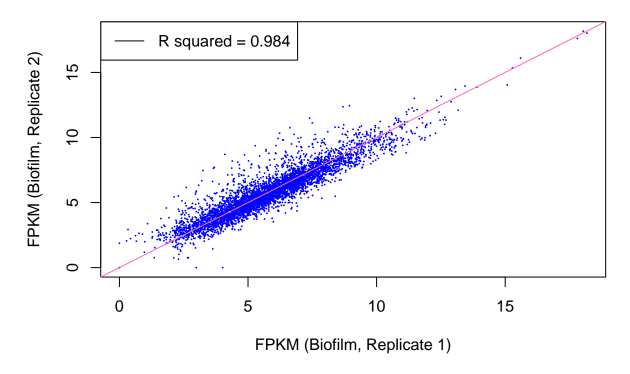
Comparison of expression values for a pair of replicates



#Creates a plot of how similar the two biofilm replicates are to one another.

```
x = gene_expression[,"biofilm01"]
y = gene_expression[,"biofilm02"]
min_nonzero=1
plot(x=log2(x+min_nonzero), y=log2(y+min_nonzero), pch=16, col="blue", cex=0.25,
xlab="FPKM (Biofilm, Replicate 1)", ylab="FPKM (Biofilm, Replicate 2)",
main="Comparison of expression values for a pair of replicates")
abline(a=0,b=1, col = "hotpink")
rs=cor(x,y)^2
legend("topleft", paste("R squared = ", round(rs, digits=3), sep=""), lwd=1, col="black")
```

Comparison of expression values for a pair of replicates

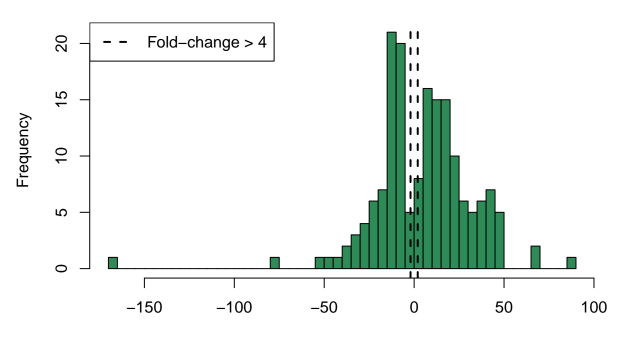


##Both replicates within each condition were similar. ##If both replicates within a single condition are similar, this means we can reliable compare between conditions (planktonic and biofilm)

#Creates plot of differential gene expression between the conditions

```
results_genes = stattest(bg_filt, feature="gene", covariate="stage", getFC=TRUE, meas="FPKM")
results_genes = merge(results_genes,bg_gene_names,by.x=c("id"),by.y=c("gene_id"))
sig=which(results_genes$pval<0.05)
results_genes[,"de"] = log2(results_genes[,"fc"])
hist(results_genes[sig,"de"], breaks=50, col="seagreen",
xlab="log2(Fold change) Planktonic vs Biofilm",
main="Distribution of differential expression values")
abline(v=-2, col="black", lwd=2, lty=2)
abline(v=2, col="black", lwd=2, lty=2)
legend("topleft", "Fold-change > 4", lwd=2, lty=2)
```

Distribution of differential expression values



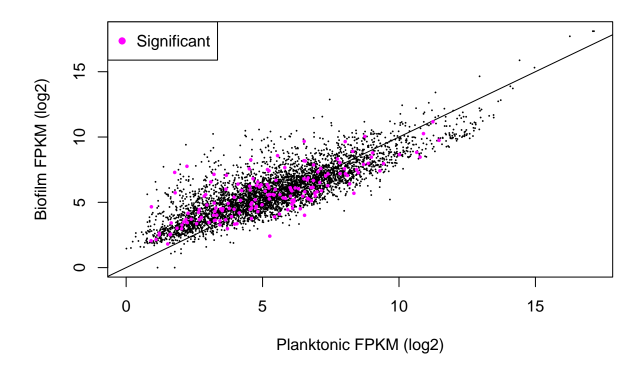
log2(Fold change) Planktonic vs Biofilm

##The differential gene expression plot is bimodal, indicating a difference in gene expression between both conditions. #Many genes had a fold-change greater than 4, suggesting a large difference in expression base on the condition (planktonic or biofilm)

#Plots total gene expression highlighting differentially expressed genes

```
gene_expression[,"plank"]=apply(gene_expression[,c(1:2)], 1, mean)
gene_expression[,"biofilm"]=apply(gene_expression[,c(3:4)], 1, mean)
x=log2(gene_expression[,"plank"]+min_nonzero)
y=log2(gene_expression[,"biofilm"]+min_nonzero)
plot(x=x, y=y, pch=16, cex=0.25, xlab="Planktonic FPKM (log2)", ylab="Biofilm FPKM (log2)",
main="Planktonic vs Biofilm FPKMs")
abline(a=0, b=1)
xsig=x[sig]
ysig=y[sig]
points(x=xsig, y=ysig, col="magenta", pch=16, cex=0.5)
legend("topleft", "Significant", col="magenta", pch=16)
```

Planktonic vs Biofilm FPKMs

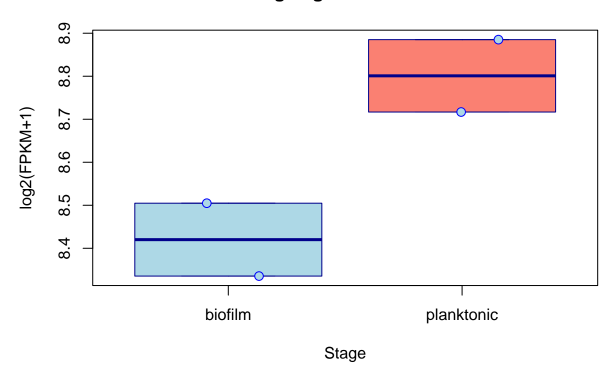


```
\#Makes a table of FPKM values
fpkm = texpr(bg_filt,meas="FPKM")
#Chooses a gene to determine individual expression
ballgown::transcriptNames(bg_filt)[10]
##
## "gene-PA0010"
ballgown::geneNames(bg_filt)[10]
##
      10
## "tag"
#Transforms to log2
transformed_fpkm <- log2(fpkm[2, ] + 1)</pre>
#Makes sure values are properly coded as numbers
numeric_stages <- as.numeric(factor(pheno_data$stage))</pre>
jittered_stages <- jitter(numeric_stages)</pre>
#Plots expression of individual gene
boxplot(transformed_fpkm ~ pheno_data$stage,
        main=paste(ballgown::geneNames(bg_filt)[10], ':', ballgown::transcriptNames(bg_filt)[10]),
        xlab="Stage",
```

```
ylab="log2(FPKM+1)",
    col=c("lightblue", "salmon"),
    border="darkblue")

points(transformed_fpkm ~ jittered_stages,
    pch=21, col="blue", bg="lightblue", cex=1.2)
```

tag: gene-PA0010



##gene-PA0010 had a higher expression in the planktonic than in the biofilm stage.