Phenotypic responses to interspecies competition and commensalism in a naturally-derived microbial co-culture

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## Supplementary Methods for Data Analysis

#### About this document

This is an R Markdown document. Markdown is a simple formatting syntax for authoring HTML, PDF, and MS Word documents. For more details on using R Markdown see <http://rmarkdown.rstudio.com>.

This project makes use of many packages, especially: DESeq2 from Bioconductor <http://bioconductor.org/packages/release/bioc/html/DESeq2.html>.

The goal of this document it to provide a reproducible comprehensive overview of data analysis methods

## Library Setup:

library("checkpoint") #Part of MS R Open to make this software stack more reproducible  
library("knitr") #package for report generation; assists with R markdown formatting  
checkpoint("2017-07-01", use.knitr = T)  
  
library("dplyr") #package for manipulating data frames  
library("ggplot2") #plotting package  
library("reshape2") #package for manipulating data frames  
library("ggrepel") #add on to ggplot2 for generating labels  
library("cowplot") #add on to ggplot2 for building "publication ready plots"  
library("viridis") #color palette package  
library("broom") #tidy stats outputs  
library("kableExtra")  
  
library("DESeq2") #significance testing for RNA-seq data  
  
knitr::opts\_chunk$set(cache=TRUE)  
theme\_set(theme\_bw())  
set.seed('711')

# DESeq is installed through bioconductor  
source("https://bioconductor.org/biocLite.R")  
biocLite("DESeq2", suppressUpdates = T)

## Import data:

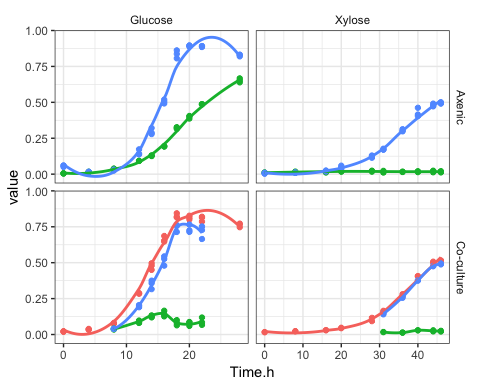
options(stringsAsFactors = FALSE)  
  
#info of treatments and replicates  
info.48 <- read.csv("../data/Condition\_Info\_HL\_48.csv")  
info.58 <- read.csv("../data/Condition\_Info\_HL\_58.csv")  
  
#growth curves and GFP-enabled FACS data  
grow <- read.csv("../data/Growth.2.csv")  
  
#gene annotation files  
an.48 <- read.csv("../data/HL\_48\_Neat.csv")  
an.58 <- read.csv("../data/HL\_58\_Neat.csv")  
  
#raw gene rollup counts  
raw.48 <- read.csv("../data/HL48\_Raw\_Counts.csv")  
raw.58 <- read.csv("../data/HL58\_Raw\_Counts.csv")  
  
meta <- read.csv("../data/Metabolites.2.csv")

## Process species-specific growth kinetics

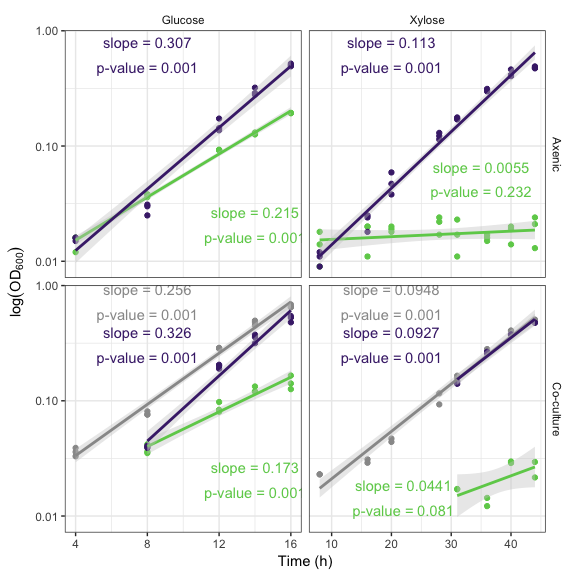
Bulk growth curves measured for each axenic and co-culture treatment the relative abundance of each species in co-culture is reported as the fraction of cells maintaining GFP these fractions were measured via FACS

#GFP.num <- data.frame(grow$OD600\*grow$HL58.GFP.frac)  
#colnames(GFP.num) <- "HL-58"  
#parent.num <- data.frame(grow$OD600\*grow$HL48.frac)  
#colnames(parent.num) <- "HL-48"  
#grow <- cbind(grow, GFP.num, parent.num)  
#vars <- c("HL.58.GFP.frac", "parent.num") #try to pull frac values out of data frame  
#grow <- cbind(grow[!vars], GFP.num, parent.num) #try to add abs values  
t1 <- melt(grow, id.vars = c("Treatment", "Cult", "Species", "Substrate",  
 "Sample.ID", "Replicate", "Time.h", "GFP.frac", "parent.frac"))  
#long format data  
  
p.gro <- ggplot(t1, aes(x = Time.h, y = value, color = variable)) +  
 geom\_point() +  
 #scale\_y\_log10() +  
 facet\_grid(Cult ~ Substrate, scales = "free\_x") +  
 theme(strip.background = element\_blank(), legend.position = "none") +  
 geom\_smooth(se = F)  
p.gro

## `geom\_smooth()` using method = 'loess'



t1 %>% dim  
t2 <- subset(t1, Time.h > 0)  
  
t2.g <- subset(t2, Substrate=="Glucose" & Time.h < 18)  
t2.x <- subset(t2, Substrate=="Xylose" & Time.h < 45)  
t2 <- rbind(t2.g, t2.x)  
  
t2 %>% dim  
  
# Let's see if we can add slopes and p-values to these graphs  
# See https://stackoverflow.com/questions/17022553/adding-r2-on-graph-with-facets  
# Also used on https://github.com/pnnl/bernstein-2017-productivity-and-diversity-2/  
  
df <- subset(t2, Substrate=="Xylose" & Cult=="Axenic" & variable == "HL.48")  
df <- subset(t2, Substrate=="Xylose" & Cult=="Axenic" & variable == "Total.OD600")  
head(df)  
dim(df)  
  
lm\_eqn\_growth = function(df){  
  
 if(all(is.na(df$value))) return("")  
 # Super important! Return an empty string for the missing values.  
  
 m = summary(lm(log(value) ~ Time.h, df)) # Hardcoded to my data  
 m  
  
 m$coefficients  
 m$coefficients[2] # Slope of Time.h  
 m$coefficients[8] # P value of Time.h  
  
 if(m$coefficients[8] < 0.001) {  
 outputp <- "0.001"  
 }else{  
 outputp <- round(m$coefficients[8], digits = 3)  
 }  
  
 eq <- substitute(  
 atop("slope ="~slope, "p-value ="~pr), # Two lines  
 #"slope ="~slope~","~~R^2~"="~r2, # One line  
 list(slope = signif(m$coefficients[2], digits = 3),  
 pr = outputp)  
 )  
 return(as.character(as.expression(eq)))  
}  
  
lm\_eqn\_growth(df)  
  
eqns <- by(t2, INDICES = list(t2$Substrate, t2$Cult, t2$variable), lm\_eqn\_growth)  
eqns  
  
df2 <-  
 data.frame(eq = c(eqns),  
 Substrate = rep(c("Glucose", "Xylose"), 6),  
 Cult = rep(c("Axenic", "Axenic", "Co-culture","Co-culture"), 3),  
 variable = c(rep("Total.OD600", 4), rep("HL.58", 4), rep("HL.48", 4))  
 ,graphx = c(10, 10, 8, 20, 14, 35, 14, 22, 8, 20, 8, 20)  
 ,graphy = c(.1, .1, .7, .7, .02, .05, .02, .014, .6, .6, .3, .3)  
)  
  
df2  
df2 <- subset(df2, eq != "") # Remove the empty lines  
  
p.gro <- ggplot(t2, aes(x = Time.h, y = value, color = variable)) +  
 geom\_point() +  
 scale\_y\_log10() +  
 geom\_text(data = df2, aes(x = graphx, y = graphy, label = eq), parse = TRUE, show.legend = F) +  
 facet\_grid(Cult ~ Substrate, scales = "free\_x") +  
 labs(x = "Time (h)", y = expression(log(OD[600])), parse = T) +  
 scale\_color\_manual(values = c("#999999", "#6DCD59FF", "#482878")) +  
 theme(strip.background = element\_blank(), legend.position = "none")  
p.gro + geom\_smooth(method = "lm", fill = "#CCCCCC", show.legend = F)



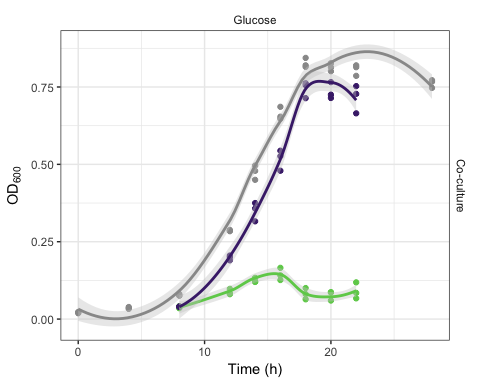
ggsave("figures/fig1-parts/fig1-growth.pdf", width = 120, units = "mm", height = 90, scale = 1.5)  
  
  
  
  
#### Try it flipped!  
# p.gro <- ggplot(t2, aes(x = Time.h, y = value, color = variable)) +  
# geom\_point() +  
# scale\_y\_log10() +  
# geom\_text(data = df2, aes(x = graphx, y = graphy, label = eq), parse = TRUE, show.legend = F) +  
# facet\_grid(Substrate ~ Cult, scales = "free\_x") +  
# labs(x = "Time (h)", y = expression(log(OD[600])), parse = T) +  
# scale\_color\_manual(values = c("#999999", "#6DCD59FF", "#482878")) +  
# theme(strip.background = element\_blank(), legend.position = "none")  
# p.gro + geom\_smooth(method = "lm", fill = "#CCCCCC", show.legend = F)  
#  
# # Note how "free\_x" no longer applied because Time.h must be consistent  
# while stacked.  
# # Positions of labels also make less sense.  
# ggsave("figures/fig1-parts/fig1-growth-flipped.pdf", width = 120, units = "mm", height = 90, scale = 1.5)

Cutaway of one frame from the above figure.

Confirm this goal: Show full time course of one block of the ANOVA design, without dropping timepoints or using a log10() transform.

# Start with the full t1 subset  
t3 <- subset(t1, Substrate=="Glucose" & Cult == "Co-culture")  
  
p.gro <- ggplot(t3, aes(x = Time.h, y = value, color = variable)) +  
 geom\_point() +  
 facet\_grid(Cult ~ Substrate, scales = "free\_x") +  
 labs(x = "Time (h)", y = expression(OD[600]), parse = T) +  
 scale\_color\_manual(values = c("#999999", "#6DCD59FF", "#482878")) +  
 theme(strip.background = element\_blank(), legend.position = "none")  
p.gro + geom\_smooth(fill = "#CCCCCC", show.legend = F)

## `geom\_smooth()` using method = 'loess'

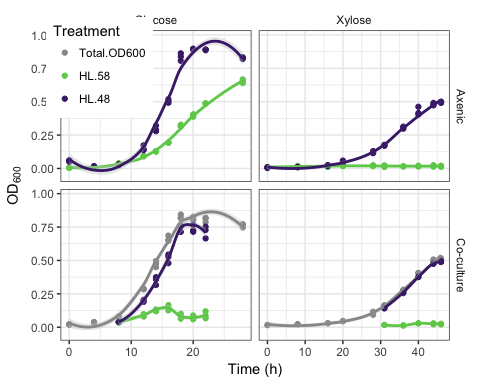


ggsave("figures/fig1-parts/fig1-growth-cutaway.pdf", width = 60, units = "mm", height = 50, scale = 1.5)

## `geom\_smooth()` using method = 'loess'

# All data, for use as a sub figure.  
# We add back in the legend, so this does not need the legend in another part of a larger figure.  
p.gro <- ggplot(t1, aes(x = Time.h, y = value, color = variable)) +  
 geom\_point() +  
 facet\_grid(Cult ~ Substrate, scales = "free\_x") +  
 labs(x = "Time (h)", y = expression(OD[600]), parse = T) +  
 scale\_color\_manual(values = c("#999999", "#6DCD59FF", "#482878"), name = "Treatment") +  
 theme(strip.background = element\_blank(), legend.position = c(.1,.88))  
p.gro + geom\_smooth(fill = "#CCCCCC", show.legend = F)

## `geom\_smooth()` using method = 'loess'



ggsave("figures/fig1-parts/fig1-growth-cutaway-full.pdf", width = 120, units = "mm", height = 90, scale = 1.5)

## `geom\_smooth()` using method = 'loess'

## Process external metabolites

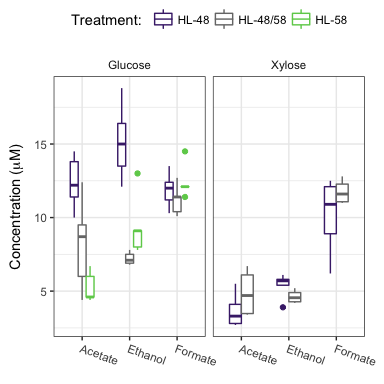
NMR metabolomics data derived from culture filtrate

Analyze and compare the concentrations of external metabolites across treatments.

summary(meta)  
str(meta)  
  
met <- filter(meta, Metabolite != "DSS-d6 (Chemical Shape Indicator)")  
met <- filter(met, Experiment.Date != "Control March 2017")  
met <- filter(met, Metabolite != "Glucose")  
met <- filter(met, Metabolite != "Xylose")  
met$Microbe <- factor(met$Microbe, levels = c("HL-48", "HL-48/58", "HL-58"))

g.met <- ggplot(met, aes(Metabolite, Conc.uM, color = Microbe))  
g.met +  
 geom\_boxplot(width = .5) +  
 labs(y = expression(paste("Concentration (", mu, M, ")")), color = "Treatment: ", x = "") +  
 facet\_wrap(~Substrate, ncol = 2) +  
 scale\_color\_manual(values = c("#482878", "#777777", "#6DCD59FF")) +  
 theme(strip.background = element\_blank(), axis.text.x = element\_text(angle = -20, hjust=0)  
 ,legend.position = "top", legend.direction = "horizontal"  
 ,plot.margin = margin(0, 1, 0, .5, "lines")  
 #,legend.position = "none", plot.margin = unit(c(0, 0, 0, .5), "lines")  
 )

## Warning: Removed 3 rows containing non-finite values (stat\_boxplot).



# wide; full width of growth curve  
ggsave("figures/fig1-parts/fig1-mets-wide1.pdf", width = 120, units = "mm", height = 60, scale = 1.3)

## Warning: Removed 3 rows containing non-finite values (stat\_boxplot).

# square; half width of growth curve  
ggsave("figures/fig1-parts/fig1-mets.pdf", width = 60, units = "mm", height = 50, scale = 1.4)

## Warning: Removed 3 rows containing non-finite values (stat\_boxplot).

# Try it flipped (matches flipped growth graph)  
# g.met +  
# geom\_boxplot(width = .5) +  
# labs(y = expression(paste("Concentration (", mu, M, ")")), color = "Treatment", x = "") +  
# facet\_wrap(~Substrate, ncol = 1, strip.position = "right") +  
# scale\_color\_manual(values = c("#482878", "#777777", "#6DCD59FF")) +  
# theme(strip.background = element\_blank(), axis.text.x = element\_text(angle = -20, hjust=0))  
#  
# ggsave("figures/fig1-parts/fig1-mets-flipped.pdf", width = 60, units = "mm", height = 50, scale = 1.4)

We need to do t-test to establish differences between ethanol and acetate abundances between treatments. This can be output as a supplementary table or kable in the markdown. You will see some notes in the Results text corresponding to this as well.

met %>% head

## Treatment Sample.Number Experiment.Date Replicate Metabolite Conc.uM X  
## 1 HL-48 G N1 Main Jan 2017 1 Acetate 11.4 G  
## 2 HL-48 G N1 Main Jan 2017 1 Ethanol 18.8 G  
## 3 HL-48 G N1 Main Jan 2017 1 Formate 11.2 G  
## 4 HL-48 G N2 Main Jan 2017 2 Acetate 10.0 G  
## 5 HL-48 G N2 Main Jan 2017 2 Ethanol 15.0 G  
## 6 HL-48 G N2 Main Jan 2017 2 Formate 12.4 G  
## Substrate Cult Microbe  
## 1 Glucose Axenic HL-48  
## 2 Glucose Axenic HL-48  
## 3 Glucose Axenic HL-48  
## 4 Glucose Axenic HL-48  
## 5 Glucose Axenic HL-48  
## 6 Glucose Axenic HL-48

# Thanks to the magic of dplyr, broom, and default stats, we can do this:  
met %>% group\_by(Substrate, Metabolite) %>%  
 do(tidy(pairwise.t.test(.$Conc.uM, .$Microbe, p.adj = "holm"))) %>%  
 kable() %>% kable\_styling(full\_width = F)

## Currently generic markdown table using pandoc is not supported.

|  |  |  |  |  |
| --- | --- | --- | --- | --- |
| Substrate | Metabolite | group1 | group2 | p.value |
| Glucose | Acetate | HL-48/58 | HL-48 | 0.0200808 |
| Glucose | Acetate | HL-58 | HL-48 | 0.0006665 |
| Glucose | Acetate | HL-58 | HL-48/58 | 0.0529371 |
| Glucose | Ethanol | HL-48/58 | HL-48 | 0.0000932 |
| Glucose | Ethanol | HL-58 | HL-48 | 0.0010511 |
| Glucose | Ethanol | HL-58 | HL-48/58 | 0.1014061 |
| Glucose | Formate | HL-48/58 | HL-48 | 0.7321162 |
| Glucose | Formate | HL-58 | HL-48 | 0.7321162 |
| Glucose | Formate | HL-58 | HL-48/58 | 0.3372185 |
| Xylose | Acetate | HL-48/58 | HL-48 | 0.2453340 |
| Xylose | Ethanol | HL-48/58 | HL-48 | 0.1623205 |
| Xylose | Formate | HL-48/58 | HL-48 | 0.2733803 |

# Note we are using the 'holm' correction because it's 'uniformly more powerful' then 'bonf'.  
# If we want to switch to Bonferroni because it's popular or we want CIs, that's ok too.  
  
# let's pull mean values too  
#met %>% group\_by(Substrate, Metabolite, Microbe) %>% summarise(mean.Conc.uM = mean(Conc.uM, na.rm = T))  
#met %>% group\_by(Substrate, Metabolite, Microbe) %>% summarise(med.Conc.uM = median(Conc.uM, na.rm = T))  
  
met %>% group\_by(Substrate, Metabolite, Microbe) %>% summarise(med.Conc.uM = median(Conc.uM, na.rm = T)) %>%  
 filter(Substrate == "Xylose", Metabolite == "Ethanol")

## # A tibble: 2 x 4  
## # Groups: Substrate, Metabolite [1]  
## Substrate Metabolite Microbe med.Conc.uM  
## <chr> <chr> <fctr> <dbl>  
## 1 Xylose Ethanol HL-48 5.70  
## 2 Xylose Ethanol HL-48/58 4.55

5.7-4.55

## [1] 1.15

(5.7-4.55)/5.7

## [1] 0.2017544

met %>% group\_by(Substrate, Metabolite, Microbe) %>% summarise(mean.Conc.uM = mean(Conc.uM, na.rm = T)) %>%  
 filter(Substrate == "Xylose", Metabolite == "Ethanol")

## # A tibble: 2 x 4  
## # Groups: Substrate, Metabolite [1]  
## Substrate Metabolite Microbe mean.Conc.uM  
## <chr> <chr> <fctr> <dbl>  
## 1 Xylose Ethanol HL-48 5.380  
## 2 Xylose Ethanol HL-48/58 4.625

5.380-4.625

## [1] 0.755

(5.380-4.625)/5.380

## [1] 0.1403346

# Line 156  
met %>% group\_by(Substrate, Metabolite, Microbe) %>% summarise(mean.Conc.uM = mean(Conc.uM, na.rm = T)) %>%  
 filter(Substrate == "Glucose", Metabolite == "Ethanol")

## # A tibble: 3 x 4  
## # Groups: Substrate, Metabolite [1]  
## Substrate Metabolite Microbe mean.Conc.uM  
## <chr> <chr> <fctr> <dbl>  
## 1 Glucose Ethanol HL-48 15.16  
## 2 Glucose Ethanol HL-48/58 7.22  
## 3 Glucose Ethanol HL-58 9.40

15.16/7.22 # subtract 1 to get 'percent increase'

## [1] 2.099723

# line 174  
met %>% group\_by(Substrate, Metabolite, Microbe) %>% summarise(mean.Conc.uM = mean(Conc.uM, na.rm = T)) %>%  
 filter(Substrate == "Glucose", Metabolite == "Ethanol")

## # A tibble: 3 x 4  
## # Groups: Substrate, Metabolite [1]  
## Substrate Metabolite Microbe mean.Conc.uM  
## <chr> <chr> <fctr> <dbl>  
## 1 Glucose Ethanol HL-48 15.16  
## 2 Glucose Ethanol HL-48/58 7.22  
## 3 Glucose Ethanol HL-58 9.40

(7.22/9.40) # relative

## [1] 0.7680851

(7.22- 9.40) / 9.4 # subtract 1 to get 'percent increase'

## [1] -0.2319149

## Process RNAseq data

Volcano plots!

#### Normalize RNAseq data to RPKM

RNA seq data processing for Halomonas HL-48; generate normalized counts in RPKM

dfcountData <- data.frame(raw.48, row.names = 1)  
dfcolData <- data.frame(info.48, row.names = 1)  
  
dds <- DESeqDataSetFromMatrix(countData = dfcountData, colData = dfcolData, design = ~condition)

## Warning in DESeqDataSet(se, design = design, ignoreRank): some variables in  
## design formula are characters, converting to factors

dds.48 <- DESeq(dds)

## estimating size factors

## estimating dispersions

## gene-wise dispersion estimates

## mean-dispersion relationship

## -- note: fitType='parametric', but the dispersion trend was not well captured by the  
## function: y = a/x + b, and a local regression fit was automatically substituted.  
## specify fitType='local' or 'mean' to avoid this message next time.

## final dispersion estimates

## fitting model and testing

notAllZero <- (rowSums(counts(dds.48)) > 0)  
vsd <- varianceStabilizingTransformation(dds.48)

## -- note: fitType='parametric', but the dispersion trend was not well captured by the  
## function: y = a/x + b, and a local regression fit was automatically substituted.  
## specify fitType='local' or 'mean' to avoid this message next time.

HL48.norm.count <- data.frame(assay(vsd[notAllZero,]))  
HL48.norm.count <- add\_rownames(HL48.norm.count, "GeneID")

## Warning: Deprecated, use tibble::rownames\_to\_column() instead.

HL48.norm.count <- merge(an.48, HL48.norm.count, by = "GeneID")  
  
write.table(HL48.norm.count, file="RNA\_seq\_outputs/HL48\_Norm\_Exp\_Values\_annotated.csv", quote=FALSE, sep=",", row.names=FALSE, col.names=TRUE)  
kable(head(HL48.norm.count))

|  |  |  |  |  |  |  |  |  |  |  |  |  |  |  |  |  |  |  |  |  |  |  |
| --- | --- | --- | --- | --- | --- | --- | --- | --- | --- | --- | --- | --- | --- | --- | --- | --- | --- | --- | --- | --- | --- | --- |
| GeneID | Acc\_num | Main\_Role | Subrole | Product | Gene | T1\_HL48G | T2\_HL48G | T4\_HL48G | T5\_HL48G | T11\_HL48G | T12\_HL48G | T13\_HL48G | T14\_HL48G | T15\_HL48G | T16\_HL48X | T17\_HL48X | T18\_HL48X | T19\_HL48X | T21\_HL48X | T22\_HL48X | T23\_HL48X | T24\_HL48X |
| hotlake\_ucc\_124538 | CY41DRAFT\_0125 | Nucleic acid metabolism | DNA replication\_ recombination\_ and repair | DNA repair protein radc |  | -5.7848388 | -6.3143024 | -4.4814389 | -4.5377806 | -4.5265101 | -4.8708085 | -5.510563 | -4.6542463 | -5.3449012 | -6.0842219 | -3.9634773 | -6.5836009 | -5.8550017 | -5.5382288 | -5.2826004 | -5.4470966 | -5.0667479 |
| hotlake\_ucc\_124539 | CY41DRAFT\_0132 | Nucleic acid metabolism | DNA replication\_ recombination\_ and repair | Nucleotidyltransferase/DNA polymerase involved in DNA repair |  | 4.9963436 | 4.7277389 | 4.7600992 | 6.3692662 | 7.2207782 | 5.0379955 | 6.861618 | 6.0918492 | 5.2610346 | 9.1327328 | 9.3193559 | 9.1138284 | 9.3900502 | 9.1654333 | 9.0195327 | 9.1943398 | 4.6304036 |
| hotlake\_ucc\_124540 | CY41DRAFT\_0143 | Mobile and extrachromosomal element functions | Selfish genetic elements | IS3 family transposase |  | 1.8254382 | 1.9322351 | 0.8025498 | 3.1816689 | 3.8016526 | 3.3107449 | 3.334614 | 3.0542134 | 1.7957520 | 0.5970179 | -0.2134157 | 0.0705615 | 0.0969528 | 0.9398863 | 0.6832929 | 0.9688571 | 2.4263460 |
| hotlake\_ucc\_124541 | CY41DRAFT\_0145 |  |  | transcriptional regulator-like protein |  | 0.0961810 | 0.0142558 | -1.6310959 | 0.9014977 | -0.7069261 | 0.3712687 | 1.719177 | 0.2093218 | -0.2096073 | 1.5702723 | 0.9160743 | 1.9851819 | 0.7020047 | 2.3341311 | 1.7835076 | 1.1339884 | 0.6228251 |
| hotlake\_ucc\_124542 | CY41DRAFT\_0180 |  |  | hypothetical protein |  | 0.0081833 | -0.1290377 | -0.5916545 | -1.1555539 | -1.8462097 | 0.5030828 | -1.057093 | -2.0216193 | 1.2537645 | -2.0979835 | -1.3978755 | -1.2436046 | -3.7383752 | -1.7227944 | -1.8882386 | -1.7831942 | 1.7534734 |
| hotlake\_ucc\_124543 | CY41DRAFT\_0217 |  |  | transposase |  | -3.2836200 | -2.3520823 | -0.8251749 | -2.6913921 | -1.0130258 | -2.0514015 | -2.805127 | 0.4255340 | -1.2107125 | -3.6241258 | -4.7173176 | -2.2823046 | -2.8908062 | -1.9026651 | -4.4514642 | -3.1411675 | -2.9460297 |

# HL48.norm.count %>% head  
# # So this simple df includes gene annotations and is normalized...  
# dds.48 %>% assay() %>% head  
# # ...while this is the full object, without normalization.  
colData(dds.48)$condition %>% table

## .  
## HL\_48\_58\_G HL\_48\_58\_X HL\_48\_G HL\_48\_X   
## 5 4 4 4

General function for DESeq contrasts.

deseq\_diff <- function(df, contrasts, annotations){  
 # perform deseq contrast  
 r <- results(df, contrasts)  
  
 # Convert to table and relablel row names to GeneID  
 r <- add\_rownames(data.frame(r), "GeneID")  
  
 # merge in annotations from file  
 r.an <- merge(annotations, r, by = "GeneID")  
  
 return(r.an)  
}

generate differential expression output for HL-48 glucose competition treatment HL-48 glucose axenic vs. HL-58/HL-48 glucose co-culture

HL48.diff.comp <- deseq\_diff(dds.48, c("condition","HL\_48\_58\_G","HL\_48\_G"), annotations = an.48)  
  
write.table(HL48.diff.comp, file="RNA\_seq\_outputs/HL48.diff.comp.csv", quote=FALSE, sep=",", row.names=FALSE, col.names=TRUE)  
kable(head(HL48.diff.comp))

|  |  |  |  |  |  |  |  |  |  |  |  |
| --- | --- | --- | --- | --- | --- | --- | --- | --- | --- | --- | --- |
| GeneID | Acc\_num | Main\_Role | Subrole | Product | Gene | baseMean | log2FoldChange | lfcSE | stat | pvalue | padj |
| hotlake\_ucc\_124538 | CY41DRAFT\_0125 | Nucleic acid metabolism | DNA replication\_ recombination\_ and repair | DNA repair protein radc |  | 9.761608 | 0.2435938 | 0.4007812 | 0.6077975 | 0.5433218 | 0.8554307 |
| hotlake\_ucc\_124539 | CY41DRAFT\_0132 | Nucleic acid metabolism | DNA replication\_ recombination\_ and repair | Nucleotidyltransferase/DNA polymerase involved in DNA repair |  | 800.375384 | 0.3960024 | 0.3193219 | 1.2401357 | 0.2149252 | 0.6491502 |
| hotlake\_ucc\_124540 | CY41DRAFT\_0143 | Mobile and extrachromosomal element functions | Selfish genetic elements | IS3 family transposase |  | 138.723430 | 0.4380131 | 0.2171244 | 2.0173366 | 0.0436604 | 0.3363009 |
| hotlake\_ucc\_124541 | CY41DRAFT\_0145 |  |  | transcriptional regulator-like protein |  | 99.795097 | 0.1635539 | 0.2409007 | 0.6789266 | 0.4971844 | 0.8371530 |
| hotlake\_ucc\_124542 | CY41DRAFT\_0180 |  |  | hypothetical protein |  | 60.185987 | 0.0298161 | 0.3886594 | 0.0767153 | 0.9388500 | 0.9854406 |
| hotlake\_ucc\_124543 | CY41DRAFT\_0217 |  |  | transposase |  | 36.230425 | 0.5619926 | 0.3528957 | 1.5925176 | 0.1112685 | 0.5081949 |

generate differential expression output for HL-48 xylose commensalism treatment HL-48 xylose axenic vs. HL-58/HL-48 xylose co-culture

HL48.diff.cmns <- deseq\_diff(dds.48, c("condition","HL\_48\_58\_X","HL\_48\_X"), an.48)  
  
write.table(HL48.diff.cmns, file="RNA\_seq\_outputs/HL48.diff.cmns.csv", quote=FALSE, sep=",", row.names=FALSE, col.names=TRUE)  
kable(head(HL48.diff.cmns))

|  |  |  |  |  |  |  |  |  |  |  |  |
| --- | --- | --- | --- | --- | --- | --- | --- | --- | --- | --- | --- |
| GeneID | Acc\_num | Main\_Role | Subrole | Product | Gene | baseMean | log2FoldChange | lfcSE | stat | pvalue | padj |
| hotlake\_ucc\_124538 | CY41DRAFT\_0125 | Nucleic acid metabolism | DNA replication\_ recombination\_ and repair | DNA repair protein radc |  | 9.761608 | 0.0039493 | 0.4815792 | 0.0082007 | 0.9934569 | 0.9963762 |
| hotlake\_ucc\_124539 | CY41DRAFT\_0132 | Nucleic acid metabolism | DNA replication\_ recombination\_ and repair | Nucleotidyltransferase/DNA polymerase involved in DNA repair |  | 800.375384 | -0.3664611 | 0.3343952 | -1.0958924 | 0.2731259 | 0.5124676 |
| hotlake\_ucc\_124540 | CY41DRAFT\_0143 | Mobile and extrachromosomal element functions | Selfish genetic elements | IS3 family transposase |  | 138.723430 | 0.5323011 | 0.2499474 | 2.1296527 | 0.0332003 | 0.1327357 |
| hotlake\_ucc\_124541 | CY41DRAFT\_0145 |  |  | transcriptional regulator-like protein |  | 99.795097 | 0.1050630 | 0.2529892 | 0.4152866 | 0.6779321 | 0.8371175 |
| hotlake\_ucc\_124542 | CY41DRAFT\_0180 |  |  | hypothetical protein |  | 60.185987 | 0.7602655 | 0.4311079 | 1.7635158 | 0.0778135 | 0.2392591 |
| hotlake\_ucc\_124543 | CY41DRAFT\_0217 |  |  | transposase |  | 36.230425 | 0.2275172 | 0.4146091 | 0.5487511 | 0.5831763 | 0.7759768 |

generate differential expression output for HL-48 competition over commensalism HL-58/HL-48 xylose co-culture (cmns) vs. HL-58/HL-48 glucose co-culture (comp)

# New plot comparing treatments  
HL48.diff.coculture <- deseq\_diff(dds.48, c("condition","HL\_48\_58\_G", "HL\_48\_58\_X"), an.48)  
  
write.table(HL48.diff.coculture, file="RNA\_seq\_outputs/HL48.diff.coculture.csv", quote=FALSE, sep=",", row.names=FALSE, col.names=TRUE)  
kable(head(HL48.diff.coculture))

|  |  |  |  |  |  |  |  |  |  |  |  |
| --- | --- | --- | --- | --- | --- | --- | --- | --- | --- | --- | --- |
| GeneID | Acc\_num | Main\_Role | Subrole | Product | Gene | baseMean | log2FoldChange | lfcSE | stat | pvalue | padj |
| hotlake\_ucc\_124538 | CY41DRAFT\_0125 | Nucleic acid metabolism | DNA replication\_ recombination\_ and repair | DNA repair protein radc |  | 9.761608 | 0.2443369 | 0.3846645 | 0.6351949 | 0.5253013 | 0.6083656 |
| hotlake\_ucc\_124539 | CY41DRAFT\_0132 | Nucleic acid metabolism | DNA replication\_ recombination\_ and repair | Nucleotidyltransferase/DNA polymerase involved in DNA repair |  | 800.375384 | -1.0727827 | 0.3176367 | -3.3773892 | 0.0007318 | 0.0019697 |
| hotlake\_ucc\_124540 | CY41DRAFT\_0143 | Mobile and extrachromosomal element functions | Selfish genetic elements | IS3 family transposase |  | 138.723430 | 0.7620211 | 0.2151973 | 3.5410352 | 0.0003986 | 0.0011479 |
| hotlake\_ucc\_124541 | CY41DRAFT\_0145 |  |  | transcriptional regulator-like protein |  | 99.795097 | -0.4921529 | 0.2333701 | -2.1088944 | 0.0349537 | 0.0614616 |
| hotlake\_ucc\_124542 | CY41DRAFT\_0180 |  |  | hypothetical protein |  | 60.185987 | 0.0664750 | 0.3848789 | 0.1727166 | 0.8628742 | 0.8995962 |
| hotlake\_ucc\_124543 | CY41DRAFT\_0217 |  |  | transposase |  | 36.230425 | 0.9260233 | 0.3489880 | 2.6534531 | 0.0079673 | 0.0170592 |

RNA seq data processing for Marinobacter HL-58; generate normalized counts in RPKM

dfcountData <- data.frame(raw.58, row.names = 1)  
dfcolData <- data.frame(info.58, row.names = 1)  
  
dds.58 <- DESeqDataSetFromMatrix(countData = dfcountData, colData = dfcolData, design = ~condition)

## Warning in DESeqDataSet(se, design = design, ignoreRank): some variables in  
## design formula are characters, converting to factors

dds.58 <- DESeq(dds.58)

## estimating size factors

## estimating dispersions

## gene-wise dispersion estimates

## mean-dispersion relationship

## -- note: fitType='parametric', but the dispersion trend was not well captured by the  
## function: y = a/x + b, and a local regression fit was automatically substituted.  
## specify fitType='local' or 'mean' to avoid this message next time.

## final dispersion estimates

## fitting model and testing

notAllZero <- (rowSums(counts(dds.58)) > 0)  
vsd <- varianceStabilizingTransformation(dds.58)

## -- note: fitType='parametric', but the dispersion trend was not well captured by the  
## function: y = a/x + b, and a local regression fit was automatically substituted.  
## specify fitType='local' or 'mean' to avoid this message next time.

HL58.norm.count <- data.frame(assay(vsd[notAllZero,]))  
HL58.norm.count <- add\_rownames(HL58.norm.count, "GeneID")

## Warning: Deprecated, use tibble::rownames\_to\_column() instead.

HL58.norm.count <- merge(an.58, HL58.norm.count, by = "GeneID")  
  
write.table(HL58.norm.count, file="RNA\_seq\_outputs/HL58\_Norm\_Exp\_Values\_annotated.csv", quote=FALSE, sep=",", row.names=FALSE, col.names=TRUE)  
kable(head(HL58.norm.count))

|  |  |  |  |  |  |  |  |  |  |  |  |  |  |  |  |  |  |  |  |
| --- | --- | --- | --- | --- | --- | --- | --- | --- | --- | --- | --- | --- | --- | --- | --- | --- | --- | --- | --- |
| GeneID | Acc\_num | Main\_Role | Subrole | Product | Gene | T6\_HL58G | T7\_HL58G | T8\_HL58G | T9\_HL58G | T10\_HL58G | T11\_HL58G | T12\_HL58G | T13\_HL58G | T14\_HL58G | T15\_HL58G | T21\_HL58X | T22\_HL58X | T23\_HL58X | T24\_HL58X |
| hotlake\_ucc\_124834 | CD01DRAFT\_3221 | Glycan biosynthesis and metabolism | Peptidoglycan metabolism | UDP-N-acetylmuramate dehydrogenase |  | 11.3358256 | 11.1800462 | 11.5913774 | 12.6287175 | 12.2522154 | 10.6245245 | 11.458865 | 10.8227724 | 11.3105664 | 11.868815 | 10.837703 | 10.8722827 | 10.9260847 | 9.8385927 |
| hotlake\_ucc\_124835 | CD01DRAFT\_3256 | Xenobiotics biodegradation and metabolism | Benzoate degradation | acetyl-CoA acetyltransferase |  | 0.2893289 | 0.0618485 | 0.2695364 | -0.2584539 | 0.0814818 | 1.6733887 | 3.072722 | 1.2447668 | 1.4060076 | 3.627405 | 2.325894 | 2.8275946 | 2.6694170 | 0.6611770 |
| hotlake\_ucc\_80896 | CD01DRAFT\_0003 | Regulatory functions::Signal transduction::Unknown function | Enzymes of unknown specificity::Two-component systems::Taxis::Small molecule interactions | two component signal transduction system histidine kinase |  | -3.9207873 | -3.8945012 | -3.5862830 | -4.0033188 | -4.8527091 | -4.2900475 | -5.839016 | -5.7183508 | -5.5616983 | -6.397703 | -6.397703 | -6.3977031 | -6.3977031 | -4.6529033 |
| hotlake\_ucc\_80898 | CD01DRAFT\_0005 |  |  | outer membrane porin |  | -0.3121074 | -1.1220738 | -0.0941957 | 1.3831640 | -0.9062989 | 0.7489158 | -1.807176 | -0.8968551 | 0.1051990 | -1.515550 | 7.121093 | 7.8805851 | 7.4294346 | 9.2445007 |
| hotlake\_ucc\_80900 | CD01DRAFT\_0006 | Unknown function | Enzymes of unknown specificity | Diacylglycerol O-acyltransferase |  | 0.1854444 | 0.3224793 | 0.0633558 | 0.0108127 | -0.4184039 | 0.5333536 | 1.387705 | -0.2424177 | 0.3700958 | 1.415378 | 2.325894 | -0.3955064 | -0.7815176 | 0.3493803 |
| hotlake\_ucc\_80901 | CD01DRAFT\_0007 |  |  | hypothetical protein |  | 7.5393210 | 7.6051410 | 7.1777729 | 8.2342630 | 7.5543323 | 10.3865789 | 8.438497 | 10.8655177 | 10.2467437 | 8.178904 | 10.794648 | 11.1480828 | 11.0841307 | 10.6393807 |

# HL58.norm.count %>% head  
# # So this simple df includes gene annotations and is normalized...  
# dds.58 %>% assay() %>% head  
# # ...while this is the full object, without normalization. Just like last time  
colData(dds.58)$condition %>% table

## .  
## HL\_48\_58\_G HL\_48\_58\_X HL\_58\_G   
## 5 4 5

more RNA seq data processing for Marinobacter HL-58; generate differential expression output for HL-58 glucose competition treatment HL-58 glucose axenic vs. HL-58/HL-48 glucose co-culture note that HL-58 does not grow on xylose; hence, differential expression cannot be analyzed for HL-58 xylose commensalism

HL58.diff.comp <- deseq\_diff(dds.58, c("condition","HL\_48\_58\_G","HL\_58\_G"), an.58)  
  
write.table(HL58.diff.comp, file="RNA\_seq\_outputs/HL58.diff.comp.csv", quote=FALSE, sep=",", row.names=TRUE, col.names=TRUE)  
kable(head(HL58.diff.comp))

|  |  |  |  |  |  |  |  |  |  |  |  |
| --- | --- | --- | --- | --- | --- | --- | --- | --- | --- | --- | --- |
| GeneID | Acc\_num | Main\_Role | Subrole | Product | Gene | baseMean | log2FoldChange | lfcSE | stat | pvalue | padj |
| hotlake\_ucc\_124834 | CD01DRAFT\_3221 | Glycan biosynthesis and metabolism | Peptidoglycan metabolism | UDP-N-acetylmuramate dehydrogenase |  | 2542.987859 | -0.3681853 | 0.2064011 | -1.7838339 | 0.0744507 | 0.1183062 |
| hotlake\_ucc\_124835 | CD01DRAFT\_3256 | Xenobiotics biodegradation and metabolism | Benzoate degradation | acetyl-CoA acetyltransferase |  | 100.671202 | 1.1086581 | 0.2105873 | 5.2646020 | 0.0000001 | 0.0000008 |
| hotlake\_ucc\_80896 | CD01DRAFT\_0003 | Regulatory functions::Signal transduction::Unknown function | Enzymes of unknown specificity::Two-component systems::Taxis::Small molecule interactions | two component signal transduction system histidine kinase |  | 3.303432 | -2.8295761 | 0.6983407 | -4.0518561 | 0.0000508 | 0.0001784 |
| hotlake\_ucc\_80897 | CD01DRAFT\_0004 |  |  | COG3287 family protein of unknown function |  | 0.000000 | NA | NA | NA | NA | NA |
| hotlake\_ucc\_80898 | CD01DRAFT\_0005 |  |  | outer membrane porin |  | 247.862849 | -0.2903843 | 0.3103231 | -0.9357484 | 0.3494028 | 0.4339249 |
| hotlake\_ucc\_80900 | CD01DRAFT\_0006 | Unknown function | Enzymes of unknown specificity | Diacylglycerol O-acyltransferase |  | 67.393844 | 0.3850037 | 0.1726139 | 2.2304326 | 0.0257187 | 0.0469052 |

generate differential expression output for HL-58 competition over commensalism HL-58/HL-48 xylose co-culture (cmns) vs. HL-58/HL-48 glucose co-culture (comp)

# new graph comparing two treatments  
HL58.diff.coculture <- deseq\_diff(dds.58, c("condition","HL\_48\_58\_G","HL\_48\_58\_X"), an.58)  
  
write.table(HL58.diff.coculture, file="RNA\_seq\_outputs/HL58.diff.coculture.csv", quote=FALSE, sep=",", row.names=TRUE, col.names=TRUE)  
kable(head(HL58.diff.coculture))

|  |  |  |  |  |  |  |  |  |  |  |  |
| --- | --- | --- | --- | --- | --- | --- | --- | --- | --- | --- | --- |
| GeneID | Acc\_num | Main\_Role | Subrole | Product | Gene | baseMean | log2FoldChange | lfcSE | stat | pvalue | padj |
| hotlake\_ucc\_124834 | CD01DRAFT\_3221 | Glycan biosynthesis and metabolism | Peptidoglycan metabolism | UDP-N-acetylmuramate dehydrogenase |  | 2542.987859 | 0.3579844 | 0.2244110 | 1.5952182 | 0.1106635 | 0.1751276 |
| hotlake\_ucc\_124835 | CD01DRAFT\_3256 | Xenobiotics biodegradation and metabolism | Benzoate degradation | acetyl-CoA acetyltransferase |  | 100.671202 | 0.1746891 | 0.2913066 | 0.5996743 | 0.5487233 | 0.6405866 |
| hotlake\_ucc\_80896 | CD01DRAFT\_0003 | Regulatory functions::Signal transduction::Unknown function | Enzymes of unknown specificity::Two-component systems::Taxis::Small molecule interactions | two component signal transduction system histidine kinase |  | 3.303432 | -1.3303797 | 1.2096676 | -1.0997895 | 0.2714238 | 0.3704850 |
| hotlake\_ucc\_80897 | CD01DRAFT\_0004 |  |  | COG3287 family protein of unknown function |  | 0.000000 | NA | NA | NA | NA | NA |
| hotlake\_ucc\_80898 | CD01DRAFT\_0005 |  |  | outer membrane porin |  | 247.862849 | -4.0592778 | 0.3342160 | -12.1456716 | 0.0000000 | 0.0000000 |
| hotlake\_ucc\_80900 | CD01DRAFT\_0006 | Unknown function | Enzymes of unknown specificity | Diacylglycerol O-acyltransferase |  | 67.393844 | 0.0106930 | 0.2899656 | 0.0368768 | 0.9705832 | 0.9780839 |

generate differential expression output for HL-58 xylose commensalism over glucose axenic HL-58 glucose axenic vs. HL-58/HL-48 xylose co-culture

Because HL-58 does not grow on xylose, but can survive in the presence of HL-48, this is used as a proxy for the impossible HL\_58\_X. Note that two treatments are applied (different substrate and introduction of HL-48).

# new graph control to two treatments  
HL58.diff.proxy <- deseq\_diff(dds.58, c("condition","HL\_48\_58\_X","HL\_58\_G"), an.58)  
  
write.table(HL58.diff.proxy, file="RNA\_seq\_outputs/HL58.diff.proxy.csv", quote=FALSE, sep=",", row.names=TRUE, col.names=TRUE)  
kable(head(HL58.diff.proxy))

|  |  |  |  |  |  |  |  |  |  |  |  |
| --- | --- | --- | --- | --- | --- | --- | --- | --- | --- | --- | --- |
| GeneID | Acc\_num | Main\_Role | Subrole | Product | Gene | baseMean | log2FoldChange | lfcSE | stat | pvalue | padj |
| hotlake\_ucc\_124834 | CD01DRAFT\_3221 | Glycan biosynthesis and metabolism | Peptidoglycan metabolism | UDP-N-acetylmuramate dehydrogenase |  | 2542.987859 | -0.7261697 | 0.2241244 | -3.240030 | 0.0011952 | 0.0034253 |
| hotlake\_ucc\_124835 | CD01DRAFT\_3256 | Xenobiotics biodegradation and metabolism | Benzoate degradation | acetyl-CoA acetyltransferase |  | 100.671202 | 0.9339691 | 0.2889583 | 3.232193 | 0.0012284 | 0.0035027 |
| hotlake\_ucc\_80896 | CD01DRAFT\_0003 | Regulatory functions::Signal transduction::Unknown function | Enzymes of unknown specificity::Two-component systems::Taxis::Small molecule interactions | two component signal transduction system histidine kinase |  | 3.303432 | -1.4991964 | 1.0699322 | -1.401207 | 0.1611522 | 0.2438791 |
| hotlake\_ucc\_80897 | CD01DRAFT\_0004 |  |  | COG3287 family protein of unknown function |  | 0.000000 | NA | NA | NA | NA | NA |
| hotlake\_ucc\_80898 | CD01DRAFT\_0005 |  |  | outer membrane porin |  | 247.862849 | 3.7688935 | 0.3242340 | 11.623992 | 0.0000000 | 0.0000000 |
| hotlake\_ucc\_80900 | CD01DRAFT\_0006 | Unknown function | Enzymes of unknown specificity | Diacylglycerol O-acyltransferase |  | 67.393844 | 0.3743107 | 0.2844672 | 1.315831 | 0.1882308 | 0.2780080 |

## Plot differentially expressed genes

setup volcano plot for HL48 glucose competition this analysis barrows ideas/code from a previously described example source <https://twbattaglia.github.io/2016/12/17/volcano-plot/>

add\_color\_cutoff <- function(df, pvalue\_cutoff = -log10(0.05), up\_name, down\_name, default\_name = "None"){  
 # This function is hard-coded to our data. It's not meant to be a general function.  
 df$color <- default\_name  
 df$color[df$lfc > 0 & df$pvalue > pvalue\_cutoff] <- up\_name  
 df$color[df$lfc < 0 & df$pvalue > pvalue\_cutoff] <- down\_name  
  
 return(df)  
}  
  
# https://stackoverflow.com/questions/7367138/text-wrap-for-plot-titles  
wrap\_strings <- function(x, goal\_width = 40){  
 as.character(sapply(x,FUN=function(x){  
 if(is.na(x)){return("")}  
 if(nchar(x) == 0){return("")}  
 print((nchar(x) / (ceiling(nchar(x)/goal\_width))))  
 paste(strwrap(x, width = (nchar(x) / (ceiling(nchar(x)/goal\_width)))), collapse=" \n")  
 }))  
}  
  
# Test:  
c("test", "", NA) %>% wrap\_strings()

## [1] 4

## [1] "test" "" ""

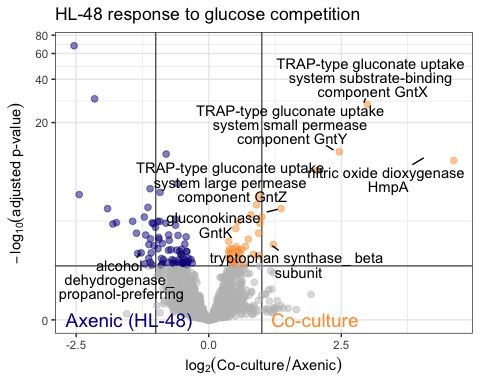
# Also use it on full gp.labels  
gp.labels$Product <- wrap\_strings(gp.labels$Product)

## [1] 29  
## [1] 33  
## [1] 31.5  
## [1] 31.5  
## [1] 18  
## [1] 33  
## [1] 21  
## [1] 24.5  
## [1] 24.5  
## [1] 26  
## [1] 31.5  
## [1] 40  
## [1] 33  
## [1] 25  
## [1] 25.5  
## [1] 26.5  
## [1] 40  
## [1] 21  
## [1] 23  
## [1] 22.5  
## [1] 20.5  
## [1] 38  
## [1] 23  
## [1] 37  
## [1] 35  
## [1] 35

plot\_v <- function(df, df.labels){  
 # This function is hard-coded to our data. It's not meant to be a general function.  
 plot <- ggplot(df, aes(x = lfc, y = pvalue))  
 return(plot +  
 #geom\_vline(xintercept = 0, color = "black") + # add line at 0  
 geom\_vline(xintercept = c(-log2(2),log2(2)), color = "grey40") + # Add cutoffs  
 geom\_hline(yintercept = -log10(0.05), color = "grey40") + # we put our pvalue cutoff in here  
 geom\_point(aes(color = factor(color)), size = 2, alpha = 0.5, na.rm = TRUE) +  
 theme(legend.position = "none") + # remove legend  
 # We let's add these manually so they match the different graph  
 #annotate("text", x = -2, y = 0, label = "Axenic", size = 5, color = "black") + # add Untreated text  
 #annotate("text", x = 2, y = 0, label = "Co-culture", size = 5, color = "red") + # add Treated text  
 #xlab(expression(log[2]("Co-culture" / "Axenic"))) + # x-axis label  
 ylab(expression(-log[10]("adjusted p-value"))) + # y-axis label  
 scale\_y\_continuous(trans = "log1p") + #transform the y-axis  
 scale\_color\_manual(values = c("Co-culture" = viridis(10, option = "C")[8],  
 "Axenic" = viridis(10, option = "C")[1],  
 "None" = "grey")) + # We could add new colors here for Commensalism and Competition plots.  
 geom\_text\_repel(data = df.labels, lineheight = 0.8,  
 mapping = aes(label = Product), min.segment.length = unit(0, "lines")  
 ,box.padding = unit(0.2, "lines"), point.padding = unit(0.2, "lines")  
 )  
 )  
}

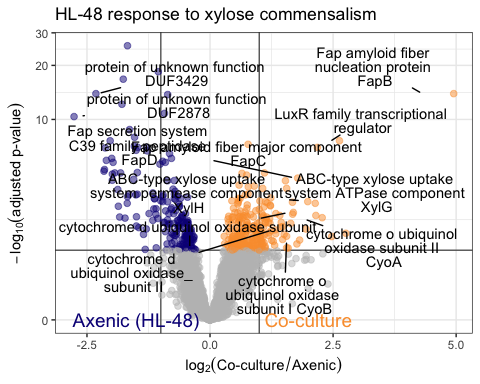
### HL-48 response to glucose competition

Fig2-A, Colored by directionality



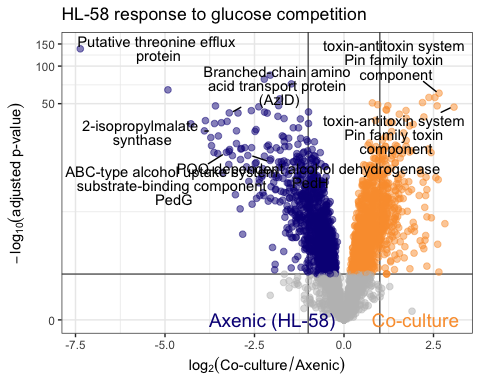
### HL-48 response to xylose commensalism

Colored by directionality



### HL-58 response to glucose competition

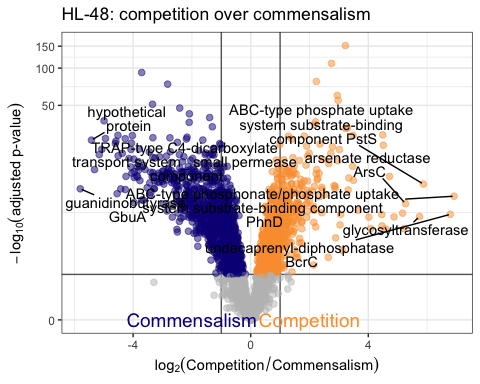
Colored by directionality



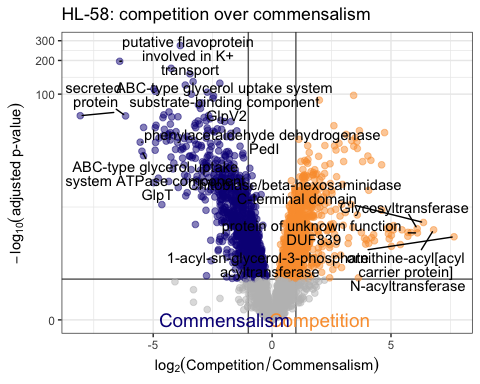
### Three new volcano plots

## [1] 23  
## [1] 32.5  
## [1] 31  
## [1] 19  
## [1] 38.5  
## [1] 20  
## [1] 34  
## [1] 35  
## [1] 22  
## [1] 34.5

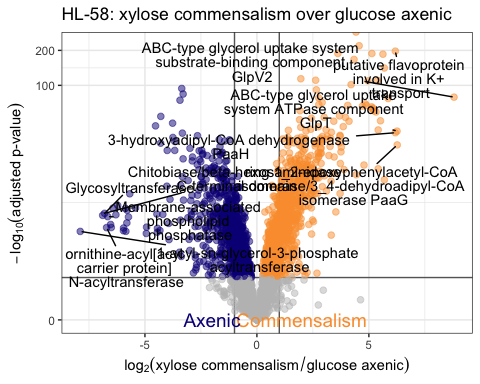
## Warning: Removed 2 rows containing missing values (geom\_text\_repel).



## [1] 24  
## [1] 34  
## [1] 19  
## [1] 23  
## [1] 27  
## [1] 16  
## [1] 26.5  
## [1] 32.5  
## [1] 23  
## [1] 37



## [1] 38  
## [1] 38.5  
## [1] 26.5  
## [1] 32.5  
## [1] 23  
## [1] 24  
## [1] 22  
## [1] 19  
## [1] 23  
## [1] 27



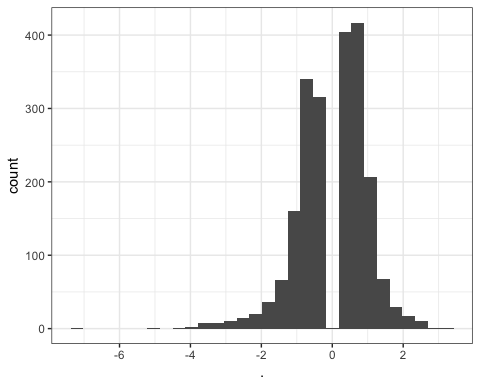
# Functional Enrichment (FE)

Dot plots!

This section applies filters to differentially expressed genes and calculates those gene function categories that are statistically enriched from the genome of each species

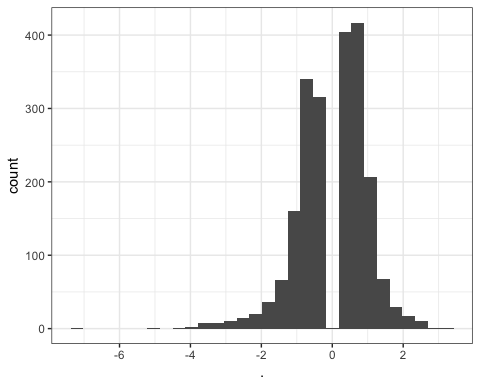
#trim differentially expressed genes by a pvalue cutoff p-adjusted <= 0.05  
HL48.diff.comp.filt <- HL48.diff.comp %>% subset(padj <= 0.05)  
HL48.diff.cmns.filt <- HL48.diff.cmns %>% subset(padj <= 0.05)  
HL58.diff.comp.filt <- HL58.diff.comp %>% subset(padj <= 0.05)  
  
# fold changes that are significant  
HL58.diff.comp.filt$log2FoldChange %>% qplot() + geom\_histogram()

## `stat\_bin()` using `bins = 30`. Pick better value with `binwidth`.  
## `stat\_bin()` using `bins = 30`. Pick better value with `binwidth`.



#trim differentially expressed genes by fold change; FC greater than 2 (< log2 = -1 or > log2 = 1)  
HL48.diff.comp.filtU <- HL48.diff.comp.filt %>% subset(log2FoldChange >= 1)  
HL48.diff.cmns.filtU <- HL48.diff.cmns.filt %>% subset(log2FoldChange >= 1)  
HL58.diff.comp.filtU <- HL58.diff.comp.filt %>% subset(log2FoldChange >= 1)  
# HL48.diff.comp.filt <- HL48.diff.comp.filt %>% subset(abs(log2FoldChange) >= 1)  
# HL48.diff.cmns.filt <- HL48.diff.cmns.filt %>% subset(abs(log2FoldChange) >= 1)  
# HL58.diff.comp.filt <- HL58.diff.comp.filt %>% subset(abs(log2FoldChange) >= 1)  
HL48.diff.comp.filtD <- HL48.diff.comp.filt %>% subset(log2FoldChange <= -1)  
HL48.diff.cmns.filtD <- HL48.diff.cmns.filt %>% subset(log2FoldChange <= -1)  
HL58.diff.comp.filtD <- HL58.diff.comp.filt %>% subset(log2FoldChange <= -1)  
  
# HL58.diff.comp.filtU$log2FoldChange %>% qplot() + geom\_histogram() # all up  
# HL58.diff.comp.filtD$log2FoldChange %>% qplot() + geom\_histogram() # all down  
  
# fold changes that are significant and OVER 2 (but not under). So this capture enrichment only.  
HL58.diff.comp.filt$log2FoldChange %>% qplot() + geom\_histogram()

## `stat\_bin()` using `bins = 30`. Pick better value with `binwidth`.  
## `stat\_bin()` using `bins = 30`. Pick better value with `binwidth`.



#prepare input files for FE  
HL48.comp.FE.U <- HL48.diff.comp.filtU %>% select(GeneID) %>% data.frame(., ModuleID = "HL\_48\_58\_G\_v\_HL\_48\_G")  
HL48.cmns.FE.U <- HL48.diff.cmns.filtU %>% select(GeneID) %>% data.frame(., ModuleID = "HL\_48\_58\_X\_v\_HL\_48\_X")  
HL58.comp.FE.U <- HL58.diff.comp.filtU %>% select(GeneID) %>% data.frame(., ModuleID = "HL\_48\_58\_G\_v\_HL\_58\_G")  
  
HL48.comp.FE.D <- HL48.diff.comp.filtD %>% select(GeneID) %>% data.frame(., ModuleID = "HL\_48\_58\_G\_v\_HL\_48\_G")  
HL48.cmns.FE.D <- HL48.diff.cmns.filtD %>% select(GeneID) %>% data.frame(., ModuleID = "HL\_48\_58\_X\_v\_HL\_48\_X")  
HL58.comp.FE.D <- HL58.diff.comp.filtD %>% select(GeneID) %>% data.frame(., ModuleID = "HL\_48\_58\_G\_v\_HL\_58\_G")

FE calculations of main role categories

#first, set up the subrole FE function  
#input x is a FE input e.g. HL48.comp.FE  
#input y is an annotation file e.g. an.48  
mainroleeModuleEnrichment <- function(x,y)  
{  
 y[grepl('::', y$Main\_Role), 'Main\_Role'] <- 'Ambiguous\_Function'  
 fModuleData <- x  
 fAnnotData <- y[, c("GeneID", "Main\_Role")]  
 colnames(fAnnotData) <- c("GeneID", "Main\_Role")  
 uniqueFunCats <- unique(fAnnotData[c("Main\_Role")])  
 Main\_Role <- unique(fAnnotData$Main\_Role)  
 modules <- unique(fModuleData$ModuleID)  
 numGenesInGenome <- nrow(fAnnotData)  
 outputData <- NULL  
 for (mID in modules)  
 {  
 genesInSet <- fModuleData[fModuleData$ModuleID == mID, "GeneID"]  
 numGenesInSet <- length(genesInSet)  
 for (i in 1:nrow(uniqueFunCats))  
 {  
 Main\_Role <- uniqueFunCats[i, "Main\_Role"]  
 genesInGenomeWithAnnot <- fAnnotData[fAnnotData$Main\_Role == Main\_Role, "GeneID"]  
 numGenesInGenomeWithAnnot <- length(genesInGenomeWithAnnot)  
 numGenesInSetWithAnnot <- length(intersect(genesInSet, genesInGenomeWithAnnot))  
 #=====================================================================================  
 # Run Fisher's exact test  
 counts <- matrix(c(numGenesInSetWithAnnot, numGenesInSet-numGenesInSetWithAnnot,  
 numGenesInGenomeWithAnnot, numGenesInGenome-numGenesInGenomeWithAnnot), nrow=2)  
 res <- fisher.test(counts)  
 if (res$p.value <= 0.05)  
 {  
 pModule <- numGenesInSetWithAnnot/numGenesInSet  
 pGenome <- numGenesInGenomeWithAnnot/numGenesInGenome  
 ratio <- pModule/pGenome  
 if (pModule > pGenome)  
 {  
 outputData <- rbind(outputData, cbind(ModuleID=mID, Main\_Role=Main\_Role, PVal=res$p.value, Ratio=ratio, PercentageInModule=pModule, PercentageInGenome=pGenome))  
 }  
 }  
 }  
 }  
 outputData <- data.frame(outputData)  
 outputData  
}  
  
#mainrolefunctial enrichment function using HL48.comp.FE and an.48 as x and y inputs  
  
HL48.comp.MR.FE.U <- mainroleeModuleEnrichment(HL48.comp.FE.U, an.48)  
HL48.comp.MR.FE.U$Treatment <- "HL-48 Glucose\nCompetition"  
HL48.comp.MR.FE.U$dir <- "Increase"  
  
#mainrolefunctial enrichment function using HL48.cmns.FE and an.48 as x and y inputs  
HL48.cmns.MR.FE.U <- mainroleeModuleEnrichment(HL48.cmns.FE.U, an.48)  
HL48.cmns.MR.FE.U$Treatment <- "HL-48 Xylose\nCommensalism"  
HL48.cmns.MR.FE.U$dir <- "Increase"  
  
#mainrolefunctial enrichment function using HL58.comp.FE and an.58 as x and y inputs  
HL58.comp.MR.FE.U <- mainroleeModuleEnrichment(HL58.comp.FE.U, an.58)  
HL58.comp.MR.FE.U$Treatment <- 'HL-58 Glucose\nCompetition'  
HL58.comp.MR.FE.U$dir <- "Increase"  
  
  
  
# matching set for reduced (D Down) enrichment  
HL48.comp.MR.FE.D <- mainroleeModuleEnrichment(HL48.comp.FE.D, an.48)  
HL48.comp.MR.FE.D$Treatment <- 'HL-48 Glucose\nCompetition'  
HL48.comp.MR.FE.D$dir <- "Decrease"  
# Empty?  
  
HL48.cmns.MR.FE.D <- mainroleeModuleEnrichment(HL48.cmns.FE.D, an.48)  
HL48.cmns.MR.FE.D$Treatment <- 'HL-48 Xylose\nCommensalism'  
HL48.cmns.MR.FE.D$dir <- "Decrease"  
  
HL58.comp.MR.FE.D <- mainroleeModuleEnrichment(HL58.comp.FE.D, an.58)  
HL58.comp.MR.FE.D$Treatment <- 'HL-58 Glucose\nCompetition'  
HL58.comp.MR.FE.D$dir <- "Decrease"  
  
  
#pull it together  
MR.FE <- rbind(HL48.comp.MR.FE.U, HL48.cmns.MR.FE.U, HL58.comp.MR.FE.U,  
 HL48.comp.MR.FE.D, # This is the empty data frame  
 HL48.cmns.MR.FE.D, HL58.comp.MR.FE.D)  
kable(MR.FE, caption = "Functional enrichment of main role gene categories")

Functional enrichment of main role gene categories

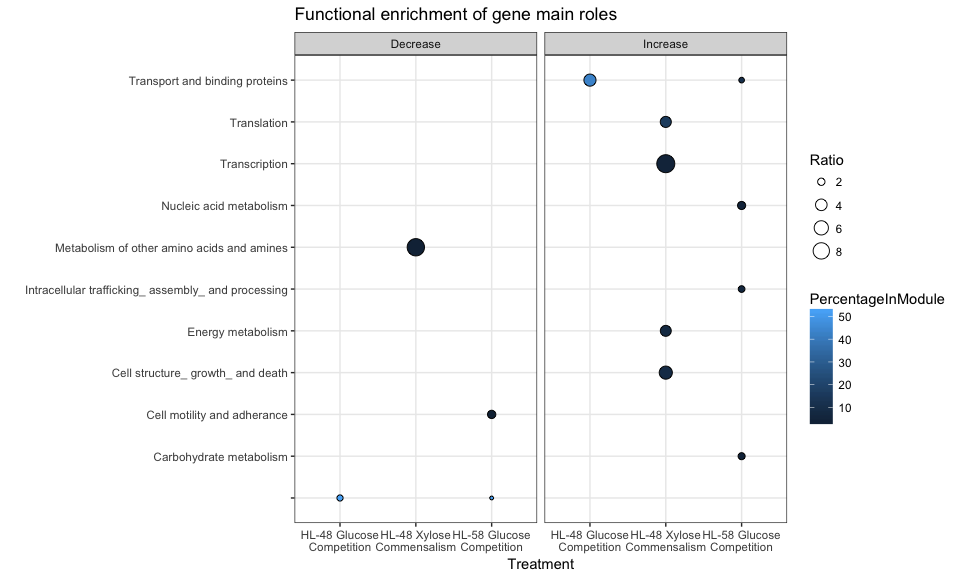
|  |  |  |  |  |  |  |  |
| --- | --- | --- | --- | --- | --- | --- | --- |
| ModuleID | Main\_Role | PVal | Ratio | PercentageInModule | PercentageInGenome | Treatment | dir |
| HL\_48\_58\_G\_v\_HL\_48\_G | Transport and binding proteins | 0.0247849323290425 | 4.3644578313253 | 0.428571428571429 | 0.0981958000591541 | HL-48 Glucose |  |
| Competition Increase |  |  |  |  |  |  |  |
| HL\_48\_58\_X\_v\_HL\_48\_X | Cell structure\_ growth\_ and death | 0.000621398637458999 | 5.10504745470233 | 0.0921052631578947 | 0.018041999408459 | HL-48 Xylose |  |
| Commensalism Increase |  |  |  |  |  |  |  |
| HL\_48\_58\_X\_v\_HL\_48\_X | Energy metabolism | 0.00765840961416274 | 3.60704125177809 | 0.0789473684210526 | 0.0218870156758355 | HL-48 Xylose |  |
| Commensalism Increase |  |  |  |  |  |  |  |
| HL\_48\_58\_X\_v\_HL\_48\_X | Translation | 0.000125291419464542 | 3.68166969147005 | 0.157894736842105 | 0.0428867199053534 | HL-48 Xylose |  |
| Commensalism Increase |  |  |  |  |  |  |  |
| HL\_48\_58\_X\_v\_HL\_48\_X | Transcription | 0.00116854340692983 | 9.8859649122807 | 0.0526315789473684 | 0.00532386867790594 | HL-48 Xylose |  |
| Commensalism Increase |  |  |  |  |  |  |  |
| HL\_48\_58\_G\_v\_HL\_58\_G | Intracellular trafficking\_ assembly\_ and processing | 0.0264067225592166 | 1.82752593907784 | 0.0631970260223048 | 0.0345806451612903 | HL-58 Glucose |  |
| Competition Increase |  |  |  |  |  |  |  |
| HL\_48\_58\_G\_v\_HL\_58\_G | Transport and binding proteins | 0.0137794066704247 | 1.618562298985 | 0.111524163568773 | 0.0689032258064516 | HL-58 Glucose |  |
| Competition Increase |  |  |  |  |  |  |  |
| HL\_48\_58\_G\_v\_HL\_58\_G | Carbohydrate metabolism | 0.0391890775576213 | 1.92069392812887 | 0.0446096654275093 | 0.0232258064516129 | HL-58 Glucose |  |
| Competition Increase |  |  |  |  |  |  |  |
| HL\_48\_58\_G\_v\_HL\_58\_G | Nucleic acid metabolism | 0.00706118571573953 | 2.26598721857901 | 0.0520446096654275 | 0.0229677419354839 | HL-58 Glucose |  |
| Competition Increase |  |  |  |  |  |  |  |
| HL\_48\_58\_G\_v\_HL\_48\_G |  | 0.0270972329550317 | 1.71859237536657 | 0.52 | 0.302573203194321 | HL-48 Glucose |  |
| Competition Decrease |  |  |  |  |  |  |  |
| HL\_48\_58\_X\_v\_HL\_48\_X | Metabolism of other amino acids and amines | 0.0244267716725998 | 9.11320754716981 | 0.0377358490566038 | 0.0041407867494824 | HL-48 Xylose |  |
| Commensalism Decrease |  |  |  |  |  |  |  |
| HL\_48\_58\_G\_v\_HL\_58\_G |  | 6.67985051333306e-08 | 1.48494842015754 | 0.515037593984962 | 0.346838709677419 | HL-58 Glucose |  |
| Competition Decrease |  |  |  |  |  |  |  |
| HL\_48\_58\_G\_v\_HL\_58\_G | Cell motility and adherance | 0.0303716770237368 | 2.33082706766917 | 0.0300751879699248 | 0.0129032258064516 | HL-58 Glucose |  |
| Competition Decrease |  |  |  |  |  |  |  |

## Plot main role functional enrichment results

MR <- data.frame(dir = MR.FE$dir, Treatment = MR.FE$Treatment, Main\_Role = MR.FE$Main\_Role,  
 Ratio = as.numeric(MR.FE$Ratio), PercentageInModule = 100\*(as.numeric(MR.FE$PercentageInModule)))  
  
MR$Ratio %>% summary

## Min. 1st Qu. Median Mean 3rd Qu. Max.   
## 1.485 1.828 2.331 3.763 4.364 9.886

g.MR.FE <- ggplot(MR, aes(x = Treatment, y = Main\_Role, size = Ratio, fill = PercentageInModule))  
g.MR.FE <- g.MR.FE +  
 #geom\_point(shape = 21, colour = "#000000", fill = "#40b8d0") +  
 geom\_point(shape = 21) +  
 facet\_grid(~dir) +  
 ggtitle("Functional enrichment of gene main roles") +  
 labs(x = "Treatment", y = "")  
g.MR.FE



FE calculations on main role categories

#first, set up the subrole FE function  
#input x is a FE input e.g. HL48.comp.FE  
#input y is an annotation file e.g. an.48  
subroleModuleEnrichment <- function(x,y)  
{  
 y[grepl('::', y$Subrole), 'Subrole'] <- 'Ambiguous\_Function'  
 fModuleData <- x  
 fAnnotData <- y[, c("GeneID", "Subrole")]  
 colnames(fAnnotData) <- c("GeneID", "Subrole")  
 uniqueFunCats <- unique(fAnnotData[c("Subrole")])  
 subRole <- unique(fAnnotData$Subrole)  
 modules <- unique(fModuleData$ModuleID)  
 numGenesInGenome <- nrow(fAnnotData)  
 outputData <- NULL  
 for (mID in modules)  
 {  
 genesInSet <- fModuleData[fModuleData$ModuleID == mID, "GeneID"]  
 numGenesInSet <- length(genesInSet)  
 for (i in 1:nrow(uniqueFunCats))  
 {  
 subRole <- uniqueFunCats[i, "Subrole"]  
 genesInGenomeWithAnnot <- fAnnotData[fAnnotData$Subrole == subRole, "GeneID"]  
 numGenesInGenomeWithAnnot <- length(genesInGenomeWithAnnot)  
 numGenesInSetWithAnnot <- length(intersect(genesInSet, genesInGenomeWithAnnot))  
 #=====================================================================================  
 # Run Fisher's exact test  
 counts <- matrix(c(numGenesInSetWithAnnot, numGenesInSet-numGenesInSetWithAnnot,  
 numGenesInGenomeWithAnnot, numGenesInGenome-numGenesInGenomeWithAnnot), nrow=2)  
 res <- fisher.test(counts)  
 if (res$p.value <= 0.05)  
 {  
 pModule <- numGenesInSetWithAnnot/numGenesInSet  
 pGenome <- numGenesInGenomeWithAnnot/numGenesInGenome  
 ratio <- pModule/pGenome  
 if (pModule > pGenome)  
 {  
 outputData <- rbind(outputData, cbind(ModuleID=mID, Subrole=subRole, PVal=res$p.value, Ratio=ratio, PercentageInModule=pModule, PercentageInGenome=pGenome))  
 }  
 }  
 }  
 }  
 outputData <- data.frame(outputData)  
 outputData  
}  
  
#subrolefunctial enrichment function using HL48.comp.FE and an.48 as x and y inputs  
HL48.comp.SR.FE.U <- subroleModuleEnrichment(HL48.comp.FE.U, an.48)  
HL48.comp.SR.FE.U$Treatment <- 'HL-48 Glucose\nCompetition'  
HL48.comp.SR.FE.U$dir <- "Increase"  
  
#subrolefunctial enrichment function using HL48.cmns.FE and an.48 as x and y inputs  
HL48.cmns.SR.FE.U <- subroleModuleEnrichment(HL48.cmns.FE.U, an.48)  
HL48.cmns.SR.FE.U$Treatment <- 'HL-48 Xylose\nCommensalism'  
HL48.cmns.SR.FE.U$dir <- "Increase"  
  
#subrolefunctial enrichment function using HL58.comp.FE and an.58 as x and y inputs  
HL58.comp.SR.FE.U <- subroleModuleEnrichment(HL58.comp.FE.U, an.58)  
HL58.comp.SR.FE.U$Treatment <- 'HL-58 Glucose\nCompetition'  
HL58.comp.SR.FE.U$dir <- "Increase"  
  
  
  
# matching set for reduced (D Down) enrichment  
HL48.comp.SR.FE.D <- subroleModuleEnrichment(HL48.comp.FE.D, an.48)  
HL48.comp.SR.FE.D$Treatment <- 'HL-48 Glucose\nCompetition'  
HL48.comp.SR.FE.D$dir <- "Decrease"  
  
HL48.cmns.SR.FE.D <- subroleModuleEnrichment(HL48.cmns.FE.D, an.48)  
HL48.cmns.SR.FE.D$Treatment <- 'HL-48 Xylose\nCommensalism'  
HL48.cmns.SR.FE.D$dir <- "Decrease"  
  
HL58.comp.SR.FE.D <- subroleModuleEnrichment(HL58.comp.FE.D, an.58)  
HL58.comp.SR.FE.D$Treatment <- 'HL-58 Glucose\nCompetition'  
HL58.comp.SR.FE.D$dir <- "Decrease"  
  
  
  
#pull it together  
SR.FE <- rbind(HL48.comp.SR.FE.U, HL48.cmns.SR.FE.U, HL58.comp.SR.FE.U,  
 HL48.comp.SR.FE.D, HL48.cmns.SR.FE.D, HL58.comp.SR.FE.D)  
kable(SR.FE, caption = "Functional enrichment of subrole gene categories")

Functional enrichment of subrole gene categories

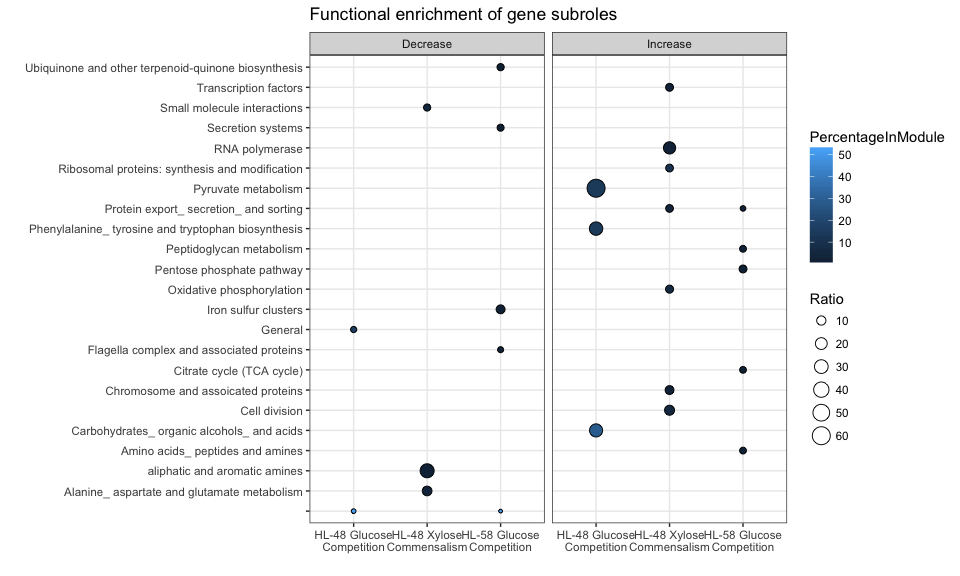
|  |  |  |  |  |  |  |  |
| --- | --- | --- | --- | --- | --- | --- | --- |
| ModuleID | Subrole | PVal | Ratio | PercentageInModule | PercentageInGenome | Treatment | dir |
| HL\_48\_58\_G\_v\_HL\_48\_G | Carbohydrates\_ organic alcohols\_ and acids | 0.00248325134199355 | 26.8333333333333 | 0.285714285714286 | 0.0106477373558119 | HL-48 Glucose |  |
| Competition Increase |  |  |  |  |  |  |  |
| HL\_48\_58\_G\_v\_HL\_48\_G | Phenylalanine\_ tyrosine and tryptophan biosynthesis | 0.0366344810198446 | 28.4117647058824 | 0.142857142857143 | 0.00502809819580006 | HL-48 Glucose |  |
| Competition Increase |  |  |  |  |  |  |  |
| HL\_48\_58\_G\_v\_HL\_48\_G | Pyruvate metabolism | 0.0184637316007577 | 60.375 | 0.142857142857143 | 0.00236616385684709 | HL-48 Glucose |  |
| Competition Increase |  |  |  |  |  |  |  |
| HL\_48\_58\_X\_v\_HL\_48\_X | Protein export\_ secretion\_ and sorting | 0.016590769967938 | 6.06638755980861 | 0.0394736842105263 | 0.00650695060632949 | HL-48 Xylose |  |
| Commensalism Increase |  |  |  |  |  |  |  |
| HL\_48\_58\_X\_v\_HL\_48\_X | Cell division | 0.000522576883149083 | 12.7105263157895 | 0.0526315789473684 | 0.0041407867494824 | HL-48 Xylose |  |
| Commensalism Increase |  |  |  |  |  |  |  |
| HL\_48\_58\_X\_v\_HL\_48\_X | Transcription factors | 0.0469103929027543 | 6.35526315789474 | 0.0263157894736842 | 0.0041407867494824 | HL-48 Xylose |  |
| Commensalism Increase |  |  |  |  |  |  |  |
| HL\_48\_58\_X\_v\_HL\_48\_X | Oxidative phosphorylation | 0.00128086052095729 | 6.74043062200957 | 0.0657894736842105 | 0.00976042590949423 | HL-48 Xylose |  |
| Commensalism Increase |  |  |  |  |  |  |  |
| HL\_48\_58\_X\_v\_HL\_48\_X | Chromosome and assoicated proteins | 0.0272987844560061 | 8.89736842105263 | 0.0263157894736842 | 0.00295770482105886 | HL-48 Xylose |  |
| Commensalism Increase |  |  |  |  |  |  |  |
| HL\_48\_58\_X\_v\_HL\_48\_X | Ribosomal proteins: synthesis and modification | 8.62947452254726e-05 | 6.0321141837645 | 0.105263157894737 | 0.0174504584442473 | HL-48 Xylose |  |
| Commensalism Increase |  |  |  |  |  |  |  |
| HL\_48\_58\_X\_v\_HL\_48\_X | RNA polymerase | 0.00675724604672468 | 22.2434210526316 | 0.0263157894736842 | 0.00118308192842354 | HL-48 Xylose |  |
| Commensalism Increase |  |  |  |  |  |  |  |
| HL\_48\_58\_G\_v\_HL\_58\_G | Peptidoglycan metabolism | 0.00174061263396245 | 4.1157727031333 | 0.0297397769516729 | 0.0072258064516129 | HL-58 Glucose |  |
| Competition Increase |  |  |  |  |  |  |  |
| HL\_48\_58\_G\_v\_HL\_58\_G | Protein export\_ secretion\_ and sorting | 0.0369465840600951 | 2.40086741016109 | 0.0260223048327138 | 0.0108387096774194 | HL-58 Glucose |  |
| Competition Increase |  |  |  |  |  |  |  |
| HL\_48\_58\_G\_v\_HL\_58\_G | Pentose phosphate pathway | 0.0077778747775878 | 6.40231309376291 | 0.0148698884758364 | 0.00232258064516129 | HL-58 Glucose |  |
| Competition Increase |  |  |  |  |  |  |  |
| HL\_48\_58\_G\_v\_HL\_58\_G | Amino acids\_ peptides and amines | 0.0257608076700826 | 4.1157727031333 | 0.0148698884758364 | 0.00361290322580645 | HL-58 Glucose |  |
| Competition Increase |  |  |  |  |  |  |  |
| HL\_48\_58\_G\_v\_HL\_58\_G | Citrate cycle (TCA cycle) | 0.0141938924524084 | 4.00144568360182 | 0.0185873605947955 | 0.00464516129032258 | HL-58 Glucose |  |
| Competition Increase |  |  |  |  |  |  |  |
| HL\_48\_58\_G\_v\_HL\_48\_G |  | 0.0270972329550317 | 1.71859237536657 | 0.52 | 0.302573203194321 | HL-48 Glucose |  |
| Competition Decrease |  |  |  |  |  |  |  |
| HL\_48\_58\_G\_v\_HL\_48\_G | General | 0.0372163936778414 | 3.14511627906977 | 0.16 | 0.0508725229222124 | HL-48 Glucose |  |
| Competition Decrease |  |  |  |  |  |  |  |
| HL\_48\_58\_X\_v\_HL\_48\_X | Small molecule interactions | 0.0277507186204185 | 4.78443396226415 | 0.0566037735849057 | 0.0118308192842354 | HL-48 Xylose |  |
| Commensalism Decrease |  |  |  |  |  |  |  |
| HL\_48\_58\_X\_v\_HL\_48\_X | Alanine\_ aspartate and glutamate metabolism | 0.016352582304608 | 11.598627787307 | 0.0377358490566038 | 0.00325347530316474 | HL-48 Xylose |  |
| Commensalism Decrease |  |  |  |  |  |  |  |
| HL\_48\_58\_X\_v\_HL\_48\_X | aliphatic and aromatic amines | 0.0456038265878301 | 31.8962264150943 | 0.0188679245283019 | 0.000591540964211772 | HL-48 Xylose |  |
| Commensalism Decrease |  |  |  |  |  |  |  |
| HL\_48\_58\_G\_v\_HL\_58\_G |  | 6.67985051333306e-08 | 1.48494842015754 | 0.515037593984962 | 0.346838709677419 | HL-58 Glucose |  |
| Competition Decrease |  |  |  |  |  |  |  |
| HL\_48\_58\_G\_v\_HL\_58\_G | Ubiquinone and other terpenoid-quinone biosynthesis | 0.0373922695734922 | 4.85588972431078 | 0.0112781954887218 | 0.00232258064516129 | HL-58 Glucose |  |
| Competition Decrease |  |  |  |  |  |  |  |
| HL\_48\_58\_G\_v\_HL\_58\_G | Secretion systems | 0.0203596330863447 | 4.48235974551764 | 0.0150375939849624 | 0.00335483870967742 | HL-58 Glucose |  |
| Competition Decrease |  |  |  |  |  |  |  |
| HL\_48\_58\_G\_v\_HL\_58\_G | Flagella complex and associated proteins | 0.0103865233371454 | 2.91353383458647 | 0.0300751879699248 | 0.0103225806451613 | HL-58 Glucose |  |
| Competition Decrease |  |  |  |  |  |  |  |
| HL\_48\_58\_G\_v\_HL\_58\_G | Iron sulfur clusters | 0.000883817191067721 | 9.10479323308271 | 0.018796992481203 | 0.00206451612903226 | HL-58 Glucose |  |
| Competition Decrease |  |  |  |  |  |  |  |

## Plot subrole functional enrichment results

SR <- data.frame(dir = SR.FE$dir, Treatment = SR.FE$Treatment, Subrole = SR.FE$Subrole,  
 Ratio = as.numeric(SR.FE$Ratio), PercentageInModule = 100\*(as.numeric(SR.FE$PercentageInModule)))  
  
SR$Ratio %>% summary

## Min. 1st Qu. Median Mean 3rd Qu. Max.   
## 1.485 4.116 6.066 11.267 11.599 60.375

g.SR.FE <- ggplot(SR, aes(x = Treatment, y = Subrole, size = Ratio, fill = PercentageInModule))  
g.SR.FE <- g.SR.FE +  
 facet\_grid(~dir) +  
 #geom\_point(shape = 21, colour = "#000000", fill = "#40b8d0") +  
 geom\_point(shape = 21) +  
 ggtitle("Functional enrichment of gene subroles") +  
 labs(x = "", y = "")  
g.SR.FE



## Combined plot of main-role and subrole functional enrichment results

names(MR)[3] <- "Main Role"  
names(SR)[3] <- "Subrole"  
  
head(MR)

## dir Treatment  
## 1 Increase HL-48 Glucose\nCompetition  
## 2 Increase HL-48 Xylose\nCommensalism  
## 3 Increase HL-48 Xylose\nCommensalism  
## 4 Increase HL-48 Xylose\nCommensalism  
## 5 Increase HL-48 Xylose\nCommensalism  
## 6 Increase HL-58 Glucose\nCompetition  
## Main Role Ratio  
## 1 Transport and binding proteins 4.364458  
## 2 Cell structure\_ growth\_ and death 5.105047  
## 3 Energy metabolism 3.607041  
## 4 Translation 3.681670  
## 5 Transcription 9.885965  
## 6 Intracellular trafficking\_ assembly\_ and processing 1.827526  
## PercentageInModule  
## 1 42.857143  
## 2 9.210526  
## 3 7.894737  
## 4 15.789474  
## 5 5.263158  
## 6 6.319703

head(SR)

## dir Treatment  
## 1 Increase HL-48 Glucose\nCompetition  
## 2 Increase HL-48 Glucose\nCompetition  
## 3 Increase HL-48 Glucose\nCompetition  
## 4 Increase HL-48 Xylose\nCommensalism  
## 5 Increase HL-48 Xylose\nCommensalism  
## 6 Increase HL-48 Xylose\nCommensalism  
## Subrole Ratio  
## 1 Carbohydrates\_ organic alcohols\_ and acids 26.833333  
## 2 Phenylalanine\_ tyrosine and tryptophan biosynthesis 28.411765  
## 3 Pyruvate metabolism 60.375000  
## 4 Protein export\_ secretion\_ and sorting 6.066388  
## 5 Cell division 12.710526  
## 6 Transcription factors 6.355263  
## PercentageInModule  
## 1 28.571429  
## 2 14.285714  
## 3 14.285714  
## 4 3.947368  
## 5 5.263158  
## 6 2.631579

# melt then combine into Enriched Roles  
mr.melt <- melt(MR, measure.vars = "Main Role")  
sr.melt <- melt(SR, measure.vars = "Subrole")  
head(mr.melt)

## dir Treatment Ratio PercentageInModule  
## 1 Increase HL-48 Glucose\nCompetition 4.364458 42.857143  
## 2 Increase HL-48 Xylose\nCommensalism 5.105047 9.210526  
## 3 Increase HL-48 Xylose\nCommensalism 3.607041 7.894737  
## 4 Increase HL-48 Xylose\nCommensalism 3.681670 15.789474  
## 5 Increase HL-48 Xylose\nCommensalism 9.885965 5.263158  
## 6 Increase HL-58 Glucose\nCompetition 1.827526 6.319703  
## variable value  
## 1 Main Role Transport and binding proteins  
## 2 Main Role Cell structure\_ growth\_ and death  
## 3 Main Role Energy metabolism  
## 4 Main Role Translation  
## 5 Main Role Transcription  
## 6 Main Role Intracellular trafficking\_ assembly\_ and processing

head(sr.melt)

## dir Treatment Ratio PercentageInModule  
## 1 Increase HL-48 Glucose\nCompetition 26.833333 28.571429  
## 2 Increase HL-48 Glucose\nCompetition 28.411765 14.285714  
## 3 Increase HL-48 Glucose\nCompetition 60.375000 14.285714  
## 4 Increase HL-48 Xylose\nCommensalism 6.066388 3.947368  
## 5 Increase HL-48 Xylose\nCommensalism 12.710526 5.263158  
## 6 Increase HL-48 Xylose\nCommensalism 6.355263 2.631579  
## variable value  
## 1 Subrole Carbohydrates\_ organic alcohols\_ and acids  
## 2 Subrole Phenylalanine\_ tyrosine and tryptophan biosynthesis  
## 3 Subrole Pyruvate metabolism  
## 4 Subrole Protein export\_ secretion\_ and sorting  
## 5 Subrole Cell division  
## 6 Subrole Transcription factors

el.melt <- rbind(mr.melt, sr.melt)  
  
# rename treatments  
el.melt$Treatment <- el.melt$Treatment %>% factor(labels =  
 c("HL-48 \nGlucose\nCompetition", "HL-48 \nXylose\nCommensalism", "HL-58 \nGlucose\nCompetition"))  
# rename treatments (without mentioning sugar type)  
#el.melt$Treatment <- el.melt$Treatment %>% factor(labels = c("HL-48 \nCompetition", "HL-48 \nCommensalism", "HL-58 \nCompetition"))  
  
# reorder levels  
el.melt$Treatment <- factor(el.melt$Treatment, levels(el.melt$Treatment)[c(1,3,2)])  
  
  
# Remove underscore  
el.melt$value <- gsub("\_", "", fixed = T, x = el.melt$value)  
# Manually edit y-axis elements in this graph. Easy way to change text text  
# pico(el.melt$value) # Get text  
# el.melt$value <- c("the string"")  
  
# Also remove the unidentified item  
el.melt <- el.melt %>% subset(value != "")  
  
g.el.FE <- ggplot(el.melt, aes(x = Treatment, y = value, size = Ratio, fill = PercentageInModule))  
g.el.FE + geom\_point(shape = 21) + labs(x = "", y = "") + facet\_grid(variable~dir, scales = "free\_y", space = "free") +  
scale\_size\_continuous(range = c(2, 10)) + scale\_fill\_viridis(option = "B", begin = .1, end = .9) +  
theme(  
 #legend.position = c(-0.9, -0.12), legend.direction = "horizontal", legend.box = "vertical", plot.margin = unit(c(10, 5.5, 30, 5.5), "points"),  
 legend.position = c(-0.5, -0.10), legend.direction = "horizontal", legend.box = "vertical", plot.margin = unit(c(10, 5.5, 25, 5.5), "points"),  
 strip.background = element\_blank(), strip.text = element\_text(size = 12), legend.spacing.y = unit(0, "points"),  
 axis.text.x = element\_text(angle = -30, hjust=0, vjust = 1, size = 10)) # angle of 30

ggsave("figures/fig2.pdf", width = 120, height = 120, units = "mm", scale = 1.5)

### Additional FE analysis of data from the three sup. volcano plots

# Input data sets  
  
HL48.diff.coculture.f <- HL48.diff.coculture %>% subset(padj <= 0.05)  
HL58.diff.coculture.f <- HL58.diff.coculture %>% subset(padj <= 0.05)  
HL58.diff.proxy.f <- HL58.diff.proxy %>% subset(padj <= 0.05)  
  
HL48.diff.coculture.U <- HL48.diff.coculture.f %>% subset(log2FoldChange >= 1)  
HL58.diff.coculture.U <- HL58.diff.coculture.f %>% subset(log2FoldChange >= 1)  
HL58.diff.proxy.U <- HL58.diff.proxy.f %>% subset(log2FoldChange >= 1)  
HL48.diff.coculture.D <- HL48.diff.coculture.f %>% subset(log2FoldChange <= -1)  
HL58.diff.coculture.D <- HL58.diff.coculture.f %>% subset(log2FoldChange <= -1)  
HL58.diff.proxy.D <- HL58.diff.proxy.f %>% subset(log2FoldChange <= -1)  
  
#prepare input files for FE  
HL48.diff.coculture.U.FE <- HL48.diff.coculture.U %>% select(GeneID) %>% data.frame(., ModuleID = "test")  
HL58.diff.coculture.U.FE <- HL58.diff.coculture.U %>% select(GeneID) %>% data.frame(., ModuleID = "test2")  
HL58.diff.proxy.U.FE <- HL58.diff.proxy.U %>% select(GeneID) %>% data.frame(., ModuleID = "test3")  
HL48.diff.coculture.D.FE <- HL48.diff.coculture.D %>% select(GeneID) %>% data.frame(., ModuleID = "test4")  
HL58.diff.coculture.D.FE <- HL58.diff.coculture.D %>% select(GeneID) %>% data.frame(., ModuleID = "test5")  
HL58.diff.proxy.D.FE <- HL58.diff.proxy.D %>% select(GeneID) %>% data.frame(., ModuleID = "test6")

# main role  
# Up  
HL48.diff.coculture.U.FE.MR <- mainroleeModuleEnrichment(HL48.diff.coculture.U.FE, an.48)  
HL48.diff.coculture.U.FE.MR$Treatment <- 'HL-48 Coculture'  
HL48.diff.coculture.U.FE.MR$dir <- "Increase"  
HL58.diff.coculture.U.FE.MR <- mainroleeModuleEnrichment(HL58.diff.coculture.U.FE, an.58)  
HL58.diff.coculture.U.FE.MR$Treatment <- 'HL-58 Coculture'  
HL58.diff.coculture.U.FE.MR$dir <- "Increase"  
HL58.diff.proxy.U.FE.MR <- mainroleeModuleEnrichment(HL58.diff.proxy.U.FE, an.58)  
HL58.diff.proxy.U.FE.MR$Treatment <- 'HL-58 Proxy'  
HL58.diff.proxy.U.FE.MR$dir <- "Increase"  
# down  
HL48.diff.coculture.D.FE.MR <- mainroleeModuleEnrichment(HL48.diff.coculture.D.FE, an.48)  
#HL48.diff.coculture.D.FE.MR$Treatment <- 'HL48 Coculture' # empty  
#HL48.diff.coculture.D.FE.MR$dir <- "Decrease" # empty  
HL58.diff.coculture.D.FE.MR <- mainroleeModuleEnrichment(HL58.diff.coculture.D.FE, an.58)  
HL58.diff.coculture.D.FE.MR$Treatment <- 'HL-58 Coculture'  
HL58.diff.coculture.D.FE.MR$dir <- "Decrease"  
HL58.diff.proxy.D.FE.MR <- mainroleeModuleEnrichment(HL58.diff.proxy.D.FE, an.58)  
HL58.diff.proxy.D.FE.MR$Treatment <- 'HL-58 Proxy'  
HL58.diff.proxy.D.FE.MR$dir <- "Decrease"  
  
  
#pull it together  
new.MR.FE <- rbind(HL48.diff.coculture.U.FE.MR, HL58.diff.coculture.U.FE.MR, HL58.diff.proxy.U.FE.MR,  
 HL48.diff.coculture.D.FE.MR, HL58.diff.coculture.D.FE.MR, HL58.diff.proxy.D.FE.MR)  
#kable(new.MR.FE) #caption = "Functional enrichment of main role gene categories")

# Sub role  
# Up  
HL48.diff.coculture.U.FE.SR <- subroleModuleEnrichment(HL48.diff.coculture.U.FE, an.48)  
HL48.diff.coculture.U.FE.SR$Treatment <- 'HL-48 Coculture'  
HL48.diff.coculture.U.FE.SR$dir <- "Increase"  
HL58.diff.coculture.U.FE.SR <- subroleModuleEnrichment(HL58.diff.coculture.U.FE, an.58)  
HL58.diff.coculture.U.FE.SR$Treatment <- 'HL-58 Coculture'  
HL58.diff.coculture.U.FE.SR$dir <- "Increase"  
HL58.diff.proxy.U.FE.SR <- subroleModuleEnrichment(HL58.diff.proxy.U.FE, an.58)  
HL58.diff.proxy.U.FE.SR$Treatment <- 'HL-58 Proxy'  
HL58.diff.proxy.U.FE.SR$dir <- "Increase"  
# down  
HL48.diff.coculture.D.FE.SR <- subroleModuleEnrichment(HL48.diff.coculture.D.FE, an.48)  
HL48.diff.coculture.D.FE.SR$Treatment <- 'HL-48 Coculture'  
HL48.diff.coculture.D.FE.SR$dir <- "Decrease"  
HL58.diff.coculture.D.FE.SR <- subroleModuleEnrichment(HL58.diff.coculture.D.FE, an.58)  
HL58.diff.coculture.D.FE.SR$Treatment <- 'HL-58 Coculture'  
HL58.diff.coculture.D.FE.SR$dir <- "Decrease"  
HL58.diff.proxy.D.FE.SR <- subroleModuleEnrichment(HL58.diff.proxy.D.FE, an.58)  
HL58.diff.proxy.D.FE.SR$Treatment <- 'HL-58 Proxy'  
HL58.diff.proxy.D.FE.SR$dir <- "Decrease"  
  
  
#pull it together  
new.SR.FE <- rbind(HL48.diff.coculture.U.FE.SR, HL58.diff.coculture.U.FE.SR, HL58.diff.proxy.U.FE.SR,  
 HL48.diff.coculture.D.FE.SR, HL58.diff.coculture.D.FE.SR, HL58.diff.proxy.D.FE.SR)  
#kable(new.SR.FE) #caption = "Functional enrichment of sub-role gene categories")

new.MR <- data.frame(dir = new.MR.FE$dir, Treatment = new.MR.FE$Treatment, Main\_role = new.MR.FE$Main\_Role,  
 Ratio = as.numeric(new.MR.FE$Ratio), PercentageInModule = 100\*(as.numeric(new.MR.FE$PercentageInModule)))  
  
new.SR <- data.frame(dir = new.SR.FE$dir, Treatment = new.SR.FE$Treatment, Subrole = new.SR.FE$Subrole,  
 Ratio = as.numeric(new.SR.FE$Ratio), PercentageInModule = 100\*(as.numeric(new.SR.FE$PercentageInModule)))  
  
names(new.MR)[3] <- "Main Role"  
names(new.SR)[3] <- "Subrole"  
  
head(new.MR)

## dir Treatment Main Role  
## 1 Increase HL-48 Coculture Carbohydrate metabolism  
## 2 Increase HL-58 Coculture Carbohydrate metabolism  
## 3 Increase HL-58 Proxy Xenobiotics biodegradation and metabolism  
## 4 Increase HL-58 Proxy Fatty acid and lipid metabolism  
## 5 Decrease HL-58 Coculture Xenobiotics biodegradation and metabolism  
## 6 Decrease HL-58 Coculture   
## Ratio PercentageInModule  
## 1 2.396173 6.024096  
## 2 1.847878 4.291845  
## 3 5.740741 1.481481  
## 4 1.845238 3.333333  
## 5 5.391304 1.391304  
## 6 1.328772 46.086957

head(new.SR)

## dir Treatment Subrole  
## 1 Increase HL-48 Coculture Polysacharide and lipopolysaccharide metabolism  
## 2 Increase HL-48 Coculture Invasion response  
## 3 Increase HL-48 Coculture Glycolysis / Gluconeogenesis  
## 4 Increase HL-48 Coculture Pyruvate metabolism  
## 5 Increase HL-48 Coculture Butanoate metabolism  
## 6 Increase HL-48 Coculture Anions  
## Ratio PercentageInModule  
## 1 2.586345 3.212851  
## 2 5.222428 2.008032  
## 3 4.849398 2.008032  
## 4 5.091867 1.204819  
## 5 10.183735 1.204819  
## 6 5.657631 4.016064

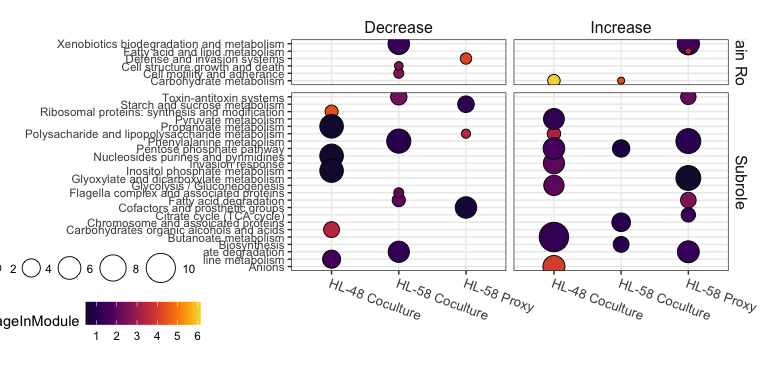
# melt then combine into Enriched Roles  
new.mr.melt <- melt(new.MR, measure.vars = "Main Role")  
new.sr.melt <- melt(new.SR, measure.vars = "Subrole")  
head(new.mr.melt)

## dir Treatment Ratio PercentageInModule variable  
## 1 Increase HL-48 Coculture 2.396173 6.024096 Main Role  
## 2 Increase HL-58 Coculture 1.847878 4.291845 Main Role  
## 3 Increase HL-58 Proxy 5.740741 1.481481 Main Role  
## 4 Increase HL-58 Proxy 1.845238 3.333333 Main Role  
## 5 Decrease HL-58 Coculture 5.391304 1.391304 Main Role  
## 6 Decrease HL-58 Coculture 1.328772 46.086957 Main Role  
## value  
## 1 Carbohydrate metabolism  
## 2 Carbohydrate metabolism  
## 3 Xenobiotics biodegradation and metabolism  
## 4 Fatty acid and lipid metabolism  
## 5 Xenobiotics biodegradation and metabolism  
## 6

head(new.sr.melt)

## dir Treatment Ratio PercentageInModule variable  
## 1 Increase HL-48 Coculture 2.586345 3.212851 Subrole  
## 2 Increase HL-48 Coculture 5.222428 2.008032 Subrole  
## 3 Increase HL-48 Coculture 4.849398 2.008032 Subrole  
## 4 Increase HL-48 Coculture 5.091867 1.204819 Subrole  
## 5 Increase HL-48 Coculture 10.183735 1.204819 Subrole  
## 6 Increase HL-48 Coculture 5.657631 4.016064 Subrole  
## value  
## 1 Polysacharide and lipopolysaccharide metabolism  
## 2 Invasion response  
## 3 Glycolysis / Gluconeogenesis  
## 4 Pyruvate metabolism  
## 5 Butanoate metabolism  
## 6 Anions

new.el.melt <- rbind(new.mr.melt, new.sr.melt)  
  
# rename treatments. (Maybe not needed based on my new names for 'modules')  
# new.el.melt$Treatment <- el.melt$Treatment %>% factor(labels = c("HL48 \nGlucose\nCompetition", "HL48 \nXylose\nCommensalism", "HL58 \nGlucose\nCompetition"))  
  
# Remove underscore  
new.el.melt$value <- gsub("\_", "", fixed = T, x = new.el.melt$value)  
# Manually edit y-axis elements in this graph. Easy way to change text text  
# pico(el.melt$value) # Get text  
# el.melt$value <- c("the string"")  
  
# Also remove the unidentified item  
new.el.melt <- new.el.melt %>% subset(value != "")  
  
g.new.el.FE <- ggplot(new.el.melt, aes(x = Treatment, y = value, size = Ratio, fill = PercentageInModule))  
g.new.el.FE + geom\_point(shape = 21) + labs(x = "", y = "") + facet\_grid(variable~dir, scales = "free\_y", space = "free") +  
scale\_size\_continuous(range = c(2, 10)) + scale\_fill\_viridis(option = "B", begin = .1, end = .9) +  
theme(  
 #legend.position = c(-0.9, -0.12), legend.direction = "horizontal", legend.box = "vertical", plot.margin = unit(c(10, 5.5, 30, 5.5), "points"),  
 legend.position = c(-0.5, -0.10), legend.direction = "horizontal", legend.box = "vertical", plot.margin = unit(c(10, 10, 35, 5.5), "points"),  
 strip.background = element\_blank(), strip.text = element\_text(size = 12), legend.spacing.y = unit(0, "points"),  
 axis.text.x = element\_text(angle = -20, hjust=0, vjust = 1, size = 10))



ggsave("figures/sub-FE-new.pdf", width = 114, height = 140, units = "mm", scale = 1.5)

#Save and exit.  
knitr::knit\_exit(F)

FALSE