Hot and Cold Thermal Challenges Astrangia poculata

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Here, we leverage the temperate stony coral, Astrangia poculata, which naturally exhibits a facultative symbiosis with Symbiodiniaceae, to explicitly examine how thermal challenges influence coral hosts in isolation from their symbionts. Aposymbiotic A. poculata were collected from Woods Hole, MA, the northern range limit for this species. Corals were thermally challenged in two independent common garden experiments (Heat challenge: 31C, 10 days; Cold challenge: 6C, 16 days) to determine the effects of divergent thermal stressors. Behavioural responses to food stimuli were monitored throughout the thermal challenges and genome-wide gene expression profiling (TagSeq) was used to characterize molecular underpinnings of the coral's response to stress in its aposymbiotic state.

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Libraries

```
library(DESeq2)
library(ggplot2)
library(affycoretools)
library(arrayQualityMetrics)
library(genefilter)
library(DESeq)
library(cowplot)
library(readr)
library(RColorBrewer)
library(gplots)
library(knitr)
library(plotly)
library(vegan)
library(kableExtra)
library(reshape2)
library(prettydoc)
library(VennDiagram)
library(ggrepel)
library(stringr)
library(brms)
library(tidyverse)
library(ggpubr)
```

#Behavioural Analysis

Coral polyp behaviours were observed every 3-4 days throughout the experiment. The total surface area of the coral that had extended vs retracted polyps was estimated and then scored between 1-5 based on polyp activity

Score	Percent.of.colony.with.extended.polyps
1	0
2	25
3	50
4	75
5	100

##Cold

Read in data, transform to long form and organize.

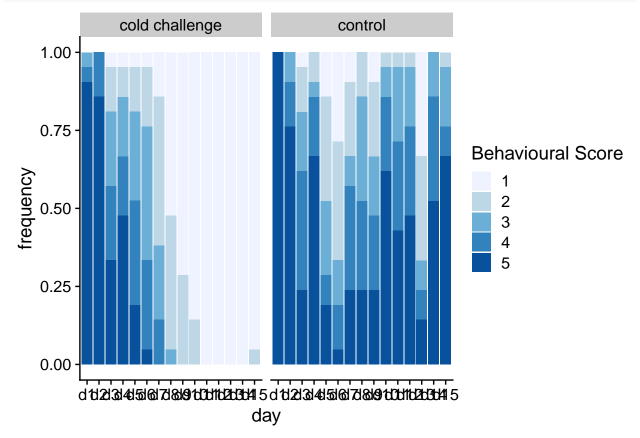
```
cold_behaviour = read.csv("cold_behaviour.csv") %>%
  melt() %>%
  dplyr::rename(day = variable) %>%
  dplyr::rename(polyp_behaviour = value) %>%
  dplyr::rename(treatment = group)
```

This is how we plotted the behavioural figure in the manuscript.

```
cold_stacked = cold_behaviour %>%
  group_by(polyp_behaviour, day, treatment) %>%
  dplyr::summarise(frequency = n())
```

```
## `summarise()` regrouping output by 'polyp_behaviour', 'day' (override with `.groups` argument)
```

```
#Plot it
ggplot(cold_stacked, aes(y = frequency, x = day, fill = as.factor(polyp_behaviour))) +
  geom_bar(stat = "identity", position = "fill") +
  facet_grid(. ~ treatment) +
  scale_fill_brewer(palette = "Blues", direction=1) +
  labs(fill = "Behavioural Score") +
  theme_cowplot()
```



Now to determine if there is any statistical differences between treatment groups. We ran a Bayesian mixed effects ordinal regression model treating coral genotype and the specific aquarium system as crossed random effects using the brms package in R (Bürkner 2017). All population-level fixed effects (e.g. treatment) had weakly informative flat priors.

```
# Do this only once
options(mc.cores=parallel::detectCores())

# model
cold_treatment_model <- brm(polyp_behaviour ~ treatment + day + (1 | genotype) + (1 | system), data = c

## Running /Library/Frameworks/R.framework/Resources/bin/R CMD SHLIB foo.c

## clang -I"/Library/Frameworks/R.framework/Resources/include" -DNDEBUG -I"/Library/Frameworks/R.frameworks/R.framework/Resources/include" -DNDEBUG -I"/Library/Frameworks/R.frameworks/R.framework/Versions/4.0/Resources/library/StanHeaders/inc

## In file included from /Library/Frameworks/R.framework/Versions/4.0/Resources/library/RcppEigen/inclu

## In file included from /Library/Frameworks/R.framework/Versions/4.0/Resources/library/RcppEigen/inclu

## Library/Frameworks/R.framework/Versions/4.0/Resources/library/RcppEigen/inclu

## /Library/Frameworks/R.framework/Versions/4.0/Resources/library/RcppEigen/include/Eigen/src/Core/util

## namespace Eigen {</pre>
```

^

```
## /Library/Frameworks/R.framework/Versions/4.0/Resources/library/RcppEigen/include/Eigen/src/Core/util
## namespace Eigen {
##
##
## In file included from <built-in>:1:
## In file included from /Library/Frameworks/R.framework/Versions/4.0/Resources/library/StanHeaders/inc
## In file included from /Library/Frameworks/R.framework/Versions/4.0/Resources/library/RcppEigen/inclu
## /Library/Frameworks/R.framework/Versions/4.0/Resources/library/RcppEigen/include/Eigen/Core:96:10: f
## #include <complex>
##
## 3 errors generated.
## make: *** [foo.o] Error 1
cold_summary = summary(cold_treatment_model)
cold_summary
    Family: cumulative
     Links: mu = logit; disc = identity
## Formula: polyp_behaviour ~ treatment + day + (1 | genotype) + (1 | system)
      Data: cold_behaviour (Number of observations: 630)
## Samples: 4 chains, each with iter = 2000; warmup = 1000; thin = 1;
##
            total post-warmup samples = 4000
##
## Group-Level Effects:
   ~genotype (Number of levels: 9)
                 Estimate Est.Error 1-95% CI u-95% CI Rhat Bulk_ESS Tail_ESS
## sd(Intercept)
                      0.57
                                0.21
                                         0.28
                                                   1.08 1.00
                                                                  1510
                                                                           2458
##
  ~system (Number of levels: 6)
                 Estimate Est. Error 1-95% CI u-95% CI Rhat Bulk ESS Tail ESS
##
                                0.22
                                          0.01
                                                   0.86 1.00
                                                                  1183
                                                                           1545
##
  sd(Intercept)
                      0.26
## Population-Level Effects:
                    Estimate Est.Error 1-95% CI u-95% CI Rhat Bulk_ESS Tail_ESS
## Intercept[1]
                        -5.78
                                   0.91
                                           -7.83
                                                     -4.25 1.00
                                                                      676
                                                                               670
## Intercept[2]
                        -4.71
                                   0.91
                                           -6.78
                                                     -3.18 1.00
                                                                      675
                                                                               653
                                           -5.84
                                                     -2.29 1.00
## Intercept[3]
                        -3.82
                                   0.91
                                                                      673
                                                                               651
## Intercept[4]
                        -2.70
                                   0.90
                                           -4.71
                                                     -1.19 1.00
                                                                      680
                                                                               575
## treatmentcontrol
                         2.54
                                   0.32
                                            1.92
                                                      3.23 1.00
                                                                     2414
                                                                              1506
## dayd2
                        -1.90
                                   0.96
                                           -4.10
                                                     -0.241.00
                                                                      686
                                                                               761
## dayd3
                        -4.73
                                   0.91
                                           -6.81
                                                     -3.17 1.00
                                                                      685
                                                                               562
## dayd4
                        -3.48
                                   0.91
                                           -5.56
                                                     -1.96 1.00
                                                                      729
                                                                               595
## dayd5
                        -5.26
                                   0.92
                                           -7.38
                                                     -3.68 1.00
                                                                      670
                                                                               587
## dayd6
                        -5.87
                                   0.92
                                           -7.88
                                                     -4.34 1.00
                                                                      668
                                                                               552
## dayd7
                        -5.60
                                   0.90
                                            -7.66
                                                     -4.11 1.00
                                                                      686
                                                                               542
## dayd8
                        -6.14
                                   0.90
                                           -8.18
                                                     -4.61 1.00
                                                                      688
                                                                               650
## dayd9
                        -6.68
                                   0.91
                                           -8.77
                                                     -5.12 1.00
                                                                      707
                                                                               616
## dayd10
                        -5.95
                                   0.91
                                                     -4.42 1.00
                                                                      762
                                                                               650
                                           -8.07
## dayd11
                        -6.45
                                   0.93
                                           -8.54
                                                     -4.90 1.00
                                                                      688
                                                                               641
## dayd12
                                                                      687
                                                                               637
                        -6.39
                                   0.92
                                           -8.49
                                                     -4.83 1.00
## dayd13
                                   0.95
                                                                      733
                        -8.09
                                           -10.19
                                                     -6.441.00
                                                                               757
                                                     -4.70 1.00
## dayd14
                        -6.22
                                   0.92
                                           -8.31
                                                                      677
                                                                               660
                                                                               584
## dayd15
                        -6.11
                                   0.92
                                           -8.23
                                                     -4.57 1.00
                                                                      662
##
## Family Specific Parameters:
```

```
## Estimate Est.Error 1-95% CI u-95% CI Rhat Bulk_ESS Tail_ESS
## disc 1.00 0.00 1.00 1.00 1.00 4000 4000
##
## Samples were drawn using sampling(NUTS). For each parameter, Bulk_ESS
## and Tail_ESS are effective sample size measures, and Rhat is the potential
## scale reduction factor on split chains (at convergence, Rhat = 1).
```

Now, we want to make a statment as to how confident we are about the effects of our treatment on influencing polyp behaviour. We extract the posterior samples for the treatment condition and divide the number of times an iteration was above zero by the total number of iterations.

```
cold_post = posterior_samples(cold_treatment_model)
sum(cold_post$b_treatmentcontrol > 0) / 4000

## [1] 1
write.csv(cold_summary$fixed, "cold_bayes_fixed.csv")
```

Hot

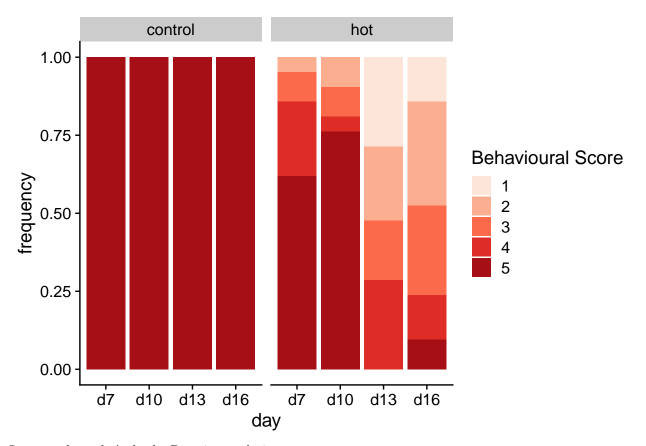
Read in data and organize

```
hot_behaviour = read.csv("hot_behaviour.csv") %>%
  melt() %>%
  rename(day = variable) %>%
  rename(polyp_behaviour = value) %>%
  mutate(genotype = sapply(strsplit(as.character(individual), split = ""), "[[", 2))
```

Displaying the data as a proportion of overall score with given phenotypes.

```
hot_stacked = hot_behaviour %>%
  group_by(polyp_behaviour, day, treatment) %>%
  summarise(frequency = n())
```

```
## `summarise()` has grouped output by 'polyp_behaviour', 'day'. You can override using the `.groups` a
ggplot(hot_stacked, aes(y = frequency, x = day, fill = as.factor(polyp_behaviour))) +
    geom_bar(stat = "identity", position = "fill") +
    facet_grid(. ~ treatment) +
    scale_fill_brewer(palette = "Reds", direction=1) +
    labs(fill = "Behavioural Score") +
    theme_cowplot()
```



Same as above, let's do the Bayesian analysis.

```
hot_treatment_model <- brm(polyp_behaviour ~ treatment + day + (1 | genotype) + (1 | system), data = ho
## Running /Library/Frameworks/R.framework/Resources/bin/R CMD SHLIB foo.c
## clang -I"/Library/Frameworks/R.framework/Resources/include" -DNDEBUG
                                                                          -I"/Library/Frameworks/R.fram
## In file included from <built-in>:1:
## In file included from /Library/Frameworks/R.framework/Versions/4.0/Resources/library/StanHeaders/inc
## In file included from /Library/Frameworks/R.framework/Versions/4.0/Resources/library/RcppEigen/inclu
## In file included from /Library/Frameworks/R.framework/Versions/4.0/Resources/library/RcppEigen/inclu
## /Library/Frameworks/R.framework/Versions/4.0/Resources/library/RcppEigen/include/Eigen/src/Core/util
## namespace Eigen {
## ^
## /Library/Frameworks/R.framework/Versions/4.0/Resources/library/RcppEigen/include/Eigen/src/Core/util
## namespace Eigen {
##
##
## In file included from <built-in>:1:
## In file included from /Library/Frameworks/R.framework/Versions/4.0/Resources/library/StanHeaders/inc
## In file included from /Library/Frameworks/R.framework/Versions/4.0/Resources/library/RcppEigen/inclu
## /Library/Frameworks/R.framework/Versions/4.0/Resources/library/RcppEigen/include/Eigen/Core:96:10: f
## #include <complex>
##
            ^~~~~~~~
## 3 errors generated.
## make: *** [foo.o] Error 1
hot_summary = summary(hot_treatment_model)
hot_summary
```

```
Family: cumulative
##
    Links: mu = logit; disc = identity
## Formula: polyp_behaviour ~ treatment + day + (1 | genotype) + (1 | system)
      Data: hot_behaviour (Number of observations: 168)
##
## Samples: 4 chains, each with iter = 2000; warmup = 1000; thin = 1;
            total post-warmup samples = 4000
##
##
## Group-Level Effects:
  ~genotype (Number of levels: 9)
##
                 Estimate Est.Error 1-95% CI u-95% CI Rhat Bulk_ESS Tail_ESS
## sd(Intercept)
                                0.39
                                         0.01
                                                   1.44 1.00
                                                                 1118
                                                                            708
##
##
  ~system (Number of levels: 6)
                 Estimate Est.Error 1-95% CI u-95% CI Rhat Bulk_ESS Tail_ESS
##
                                1.27
                                         0.04
## sd(Intercept)
                      1.13
                                                   5.02 1.01
                                                                  401
                                                                            442
##
## Population-Level Effects:
                Estimate Est.Error 1-95% CI u-95% CI Rhat Bulk ESS Tail ESS
                                      -15.42
## Intercept[1]
                  -11.17
                               2.01
                                                -7.34 1.01
                                                                 709
## Intercept[2]
                   -9.64
                               1.96
                                      -13.90
                                                 -5.89 1.01
                                                                 701
                                                                           488
## Intercept[3]
                   -8.56
                               1.92
                                      -12.71
                                                -4.80 1.01
                                                                 709
                                                                          504
                   -7.30
## Intercept[4]
                               1.88
                                      -11.35
                                                -3.58 1.01
                                                                 719
                                                                           491
                   -7.34
## treatmenthot
                               1.92
                                      -11.92
                                                -4.24 1.01
                                                                 681
                                                                          215
## davd2
                    0.48
                               0.71
                                       -0.88
                                                 1.85 1.00
                                                                1580
                                                                          833
## dayd3
                   -3.10
                               0.67
                                       -4.43
                                                -1.79 1.00
                                                                1717
                                                                          2566
## dayd4
                   -2.79
                               0.66
                                       -4.17
                                                -1.55 1.00
                                                                1521
                                                                         1117
##
## Family Specific Parameters:
        Estimate Est.Error 1-95% CI u-95% CI Rhat Bulk_ESS Tail_ESS
##
## disc
            1.00
                      0.00
                                1.00
                                         1.00 1.00
                                                        4000
                                                                 4000
##
## Samples were drawn using sampling(NUTS). For each parameter, Bulk_ESS
## and Tail_ESS are effective sample size measures, and Rhat is the potential
## scale reduction factor on split chains (at convergence, Rhat = 1).
What's our confidence statement.
hot_post = posterior_samples(hot_treatment_model)
sum(hot_post$b_treatmenthot < 0) / 4000</pre>
## [1] 0.9985
write.csv(hot_summary$fixed, "hot_bayes_fixed.csv")
```

Data Cleanup

Here we want to filter out contaminant reads. These could be from various taxa, and so we first get NCBI taxonomic info

```
wget https://ftp.ncbi.nlm.nih.gov/pub/taxonomy/new_taxdump/new_taxdump.tar.gz
tar -zxvf new_taxdump.tar.gz
```

Set up rank lineage file

```
col_types=("c-c-c-c-c-c-c-c"))
```

From this I can make a list of potential contaminants to remove called filter_list

```
filter_list = tax %>%
  filter(k == "Archaea" | k == "Bacteria" | k == "Plantae" | k == "Protozoa" | k == "Chromista" | k == "F
dplyr::select(id) %>%
  dplyr::rename(speciesID = id) %>%
  as.data.frame()
```

Compare this with the iso2gene. First I have to break apart the delimiters to get the taxonomic id's into their own column. *136 genes filtered out with this method

Now we can filter out the contamination

```
cold_counts = read.csv("cold_raw_assembled_transcriptome.csv") %>%
  dplyr::rename(Iso = X) %>%
  filter(!Iso %in% dirty$Iso) %>%
  column_to_rownames(var = "Iso")
hot_counts = read.csv("hot_raw_counts_assembled_transcriptome.csv") %>%
  dplyr::rename(Iso = X) %>%
  filter(!Iso %in% dirty$Iso) %>%
  column_to_rownames(var = "Iso")
```

Outlier Detection

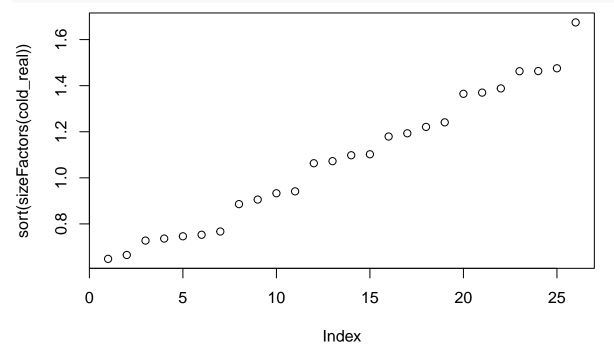
I utilized old school DESeq to look for any samples that didn't sequence properly.

Cold

First I set up experimental designs

Plotting the library size factors

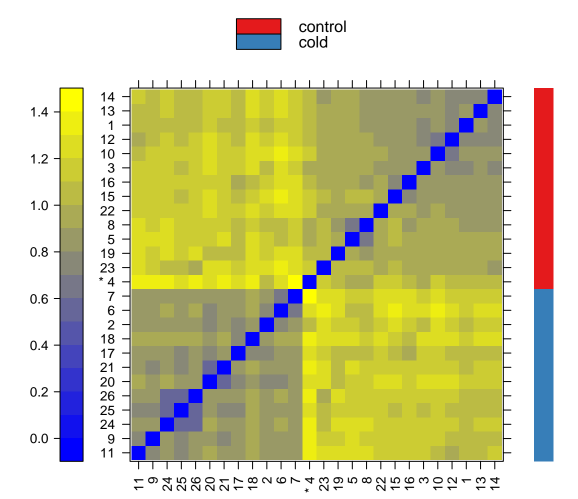
```
cold_real=newCountDataSet(cold_counts,cold_expDesign)
cold_real=estimateSizeFactors(cold_real)
plot(sort(sizeFactors(cold_real)))
```



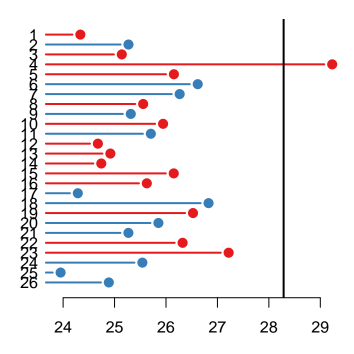
Outliers - here you have to manually inspect the html output.

```
cold_cds=estimateDispersions(cold_real,method="blind")
cold_vsdBlind=DESeq::varianceStabilizingTransformation(cold_cds)
arrayQualityMetrics(cold_vsdBlind,intgroup=c("cold_treatment"), force=TRUE, outdir = "cold_arrayQuality")
```

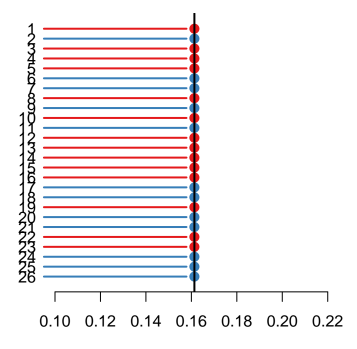
We can see the cold and control group nicely based on distances between arrays

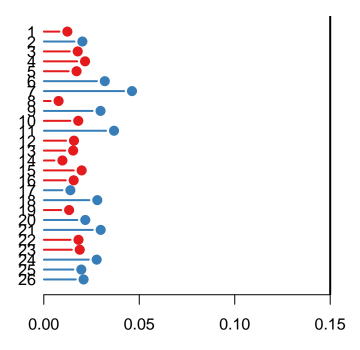


That said, there was one outlier detected based on the previous figures. The bars are shown in the original order of the arrays. Based on the distribution of the values across all arrays, a threshold of 28.3 was determined, which is indicated by the vertical line. One array exceeded the threshold and was considered an outlier. We can see the cold and control group nicely based on distances between arrays



Despite this showing an outlier, all other outlier tests tid not suggest that this sample was an outlier.





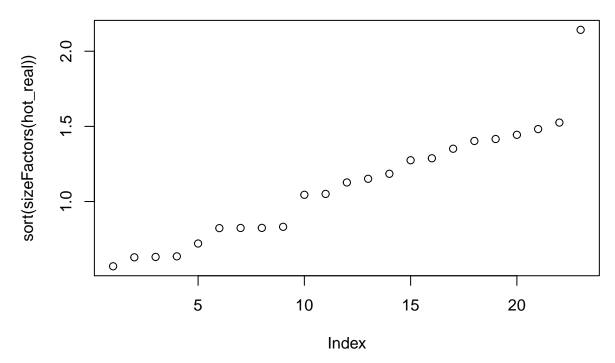
So, this sample was not deemed an outlier and it was kept in for the remainder of the analysis.

Hot

First I set up experimental designs

Plotting the library size factors

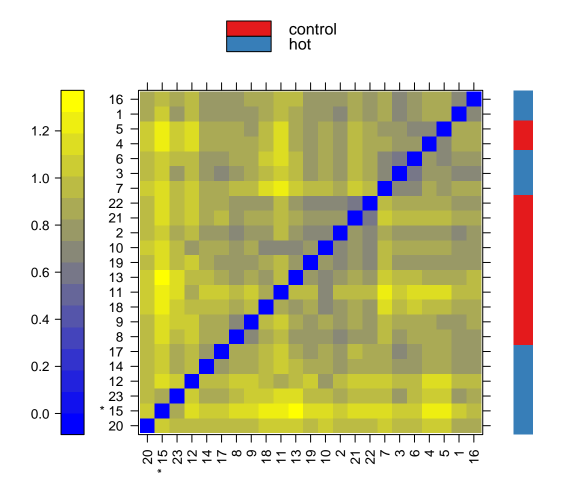
```
hot_real=newCountDataSet(hot_counts,hot_expDesign)
hot_real=estimateSizeFactors(hot_real)
plot(sort(sizeFactors(hot_real)))
```



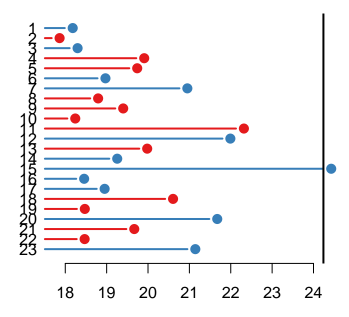
Outliers - here you have to manually inspect the html output.

```
hot_cds=estimateDispersions(hot_real,method="blind")
hot_vsdBlind=DESeq::varianceStabilizingTransformation(hot_cds)
arrayQualityMetrics(hot_vsdBlind,intgroup=c("hot_treatment"), force=TRUE, outdir = "hot_arrayQualityMet.")
```

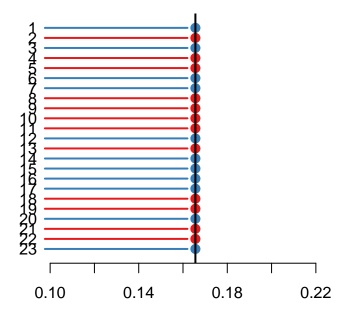
We can see the hot and control group mostly together, however not as strong a discrimination as the cold experiment based on distances between arrays

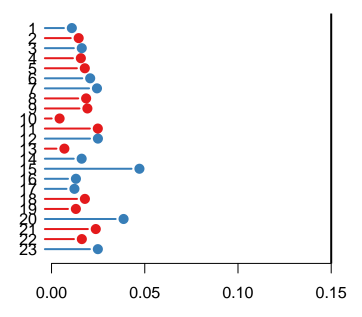


Much like in the cold, there was one sample that failed a single outlier quality metric. The bars are shown in the original order of the arrays. Based on the distribution of the values across all arrays, a threshold of 24.6 was determined, which is indicated by the vertical line. One array exceeded the threshold and was considered an outlier.



Despite this showing an outlier, all other outlier tests tid not suggest that this sample was an outlier.





All samples were kept for differential expression analysis.

Differential Expression

DESeq2 model

```
Cold
cold_dds = DESeqDataSetFromMatrix(countData = cold_counts, colData = cold_expDesign, design = ~ cold_ge
cold_dds = DESeq(cold_dds)
cold_rlogged = DESeq2::rlog(cold_dds, blind = TRUE) #for use later on
write.csv(assay(cold_rlogged), "cold_rlogged.csv")
cold_results = results(cold_dds, alpha = 0.05, contrast = c("cold_treatment", "cold", "control"))
summary(cold results)
##
## out of 13108 with nonzero total read count
## adjusted p-value < 0.05
## LFC > 0 (up)
                       : 2244, 17%
                       : 3074, 23%
## LFC < 0 (down)
## outliers [1]
                       : 1, 0.0076%
## low counts [2]
                       : 0, 0%
## (mean count < 0)
## [1] see 'cooksCutoff' argument of ?results
## [2] see 'independentFiltering' argument of ?results
write.csv(cold_results, "cold_results.csv", row.names = TRUE)
Hot
hot_dds = DESeqDataSetFromMatrix(countData = hot_counts, colData = hot_expDesign, design = ~ hot_genoty
hot_dds = DESeq(hot_dds)
hot_rlogged = rlogTransformation(hot_dds, blind = TRUE)
write.csv(assay(hot_rlogged), "hot_rlogged.csv")
hot_results = results(hot_dds, alpha = 0.05, contrast = c("hot_treatment", "hot", "control"))
```

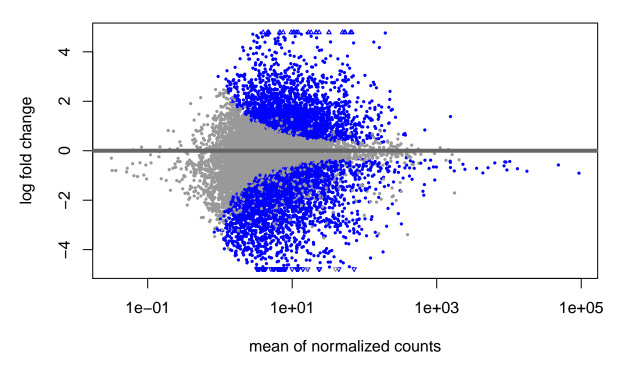
```
##
## out of 13109 with nonzero total read count
## adjusted p-value < 0.05
## LFC > 0 (up) : 410, 3.1%
## LFC < 0 (down) : 644, 4.9%
## outliers [1] : 10, 0.076%
## low counts [2] : 3304, 25%
## (mean count < 2)
## [1] see 'cooksCutoff' argument of ?results
## [2] see 'independentFiltering' argument of ?results
write.csv(hot_results, "hot_results.csv", row.names = TRUE)</pre>
```

Volcano Plots

Cold

DESeq2::plotMA(cold_results, main = "Cold vs Control")

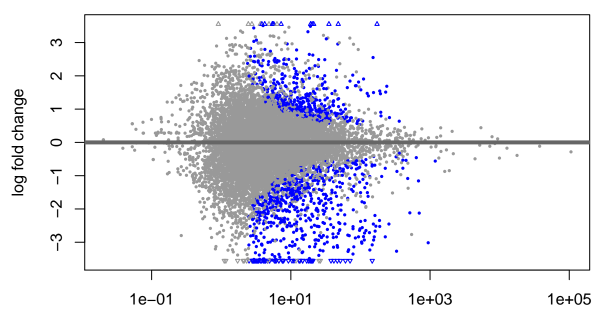
Cold vs Control



Hot

DESeq2::plotMA(hot_results, main = "Hot vs Control")

Hot vs Control



mean of normalized counts

##PCAs ###Cold

```
cold_pcadata = DESeq2::plotPCA(cold_rlogged, intgroup = c( "cold_treatment", "cold_genotype"), returnDa
cold_percentVar = round(100 * attr(cold_pcadata, "percentVar"))
cold_pca = prcomp(t(assay(cold_rlogged)), center = TRUE, scale. = FALSE)

PCA_cold = as.data.frame(cold_pca$x)%>%
    dplyr::select(PC1, PC2) %>%
    rownames_to_column("sample") %>%
    left_join(cold_expDesign)
```

```
## Joining, by = "sample"

cold_cols = c("control" = "grey", "cold" = "#68a2ff")

ggplot(PCA_cold, aes(PC1, PC2)) +

geom_point(aes(colour = cold_treatment), size = 3) +

stat_ellipse(geom = "polygon", alpha = 1/10, aes(fill = cold_treatment)) +

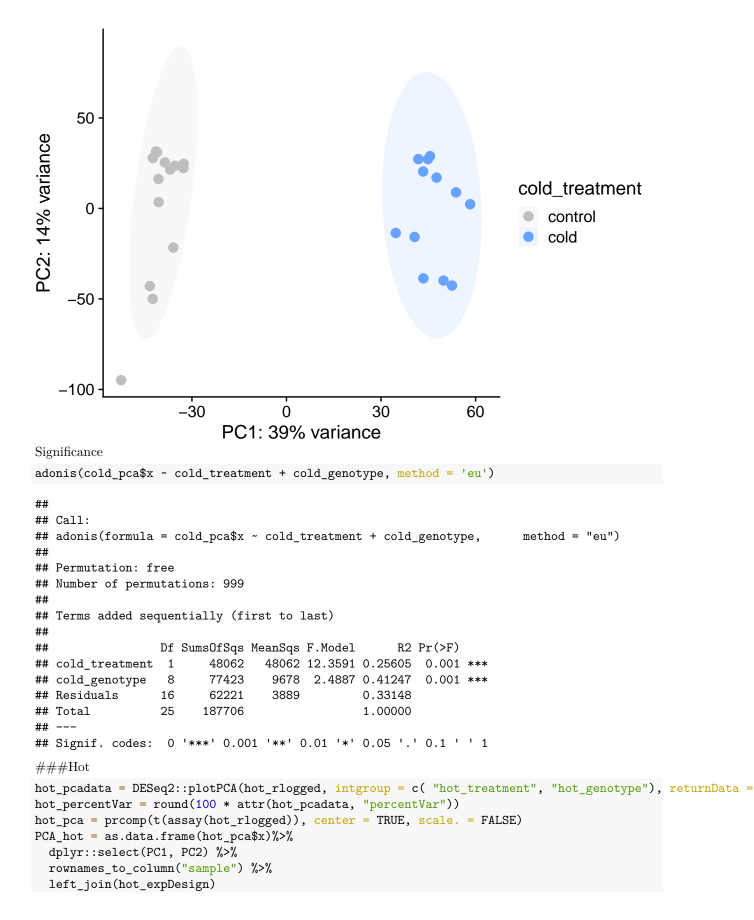
scale_colour_manual(values = cold_cols) +

scale_fill_manual(values = cold_cols) +

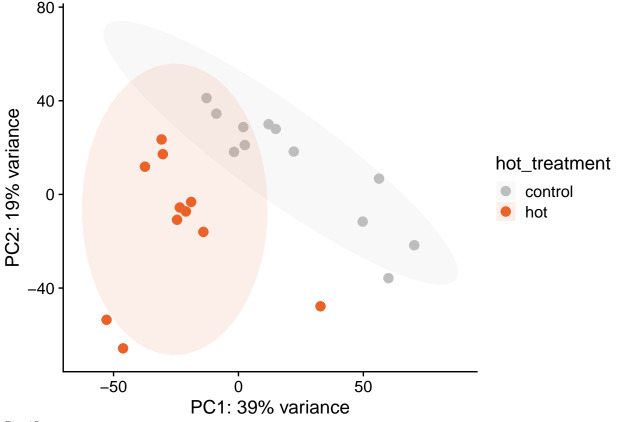
xlab(paste0("PC1: ",cold_percentVar[1],"% variance")) +

ylab(paste0("PC2: ",cold_percentVar[2],"% variance")) +

theme_cowplot()
```



```
## Joining, by = "sample"
hot_cols = c("control" = "grey", "hot" = "#ea6227")
ggplot(PCA_hot, aes(PC1, PC2)) +
    geom_point(aes(colour = hot_treatment), size = 3) +
    stat_ellipse(geom = "polygon", alpha = 1/10, aes(fill = hot_treatment)) +
    scale_colour_manual(values = hot_cols) +
    scale_fill_manual(values = hot_cols) +
    xlab(paste0("PC1: ",hot_percentVar[1],"% variance")) +
    ylab(paste0("PC2: ",hot_percentVar[2],"% variance")) +
    theme_cowplot()
```



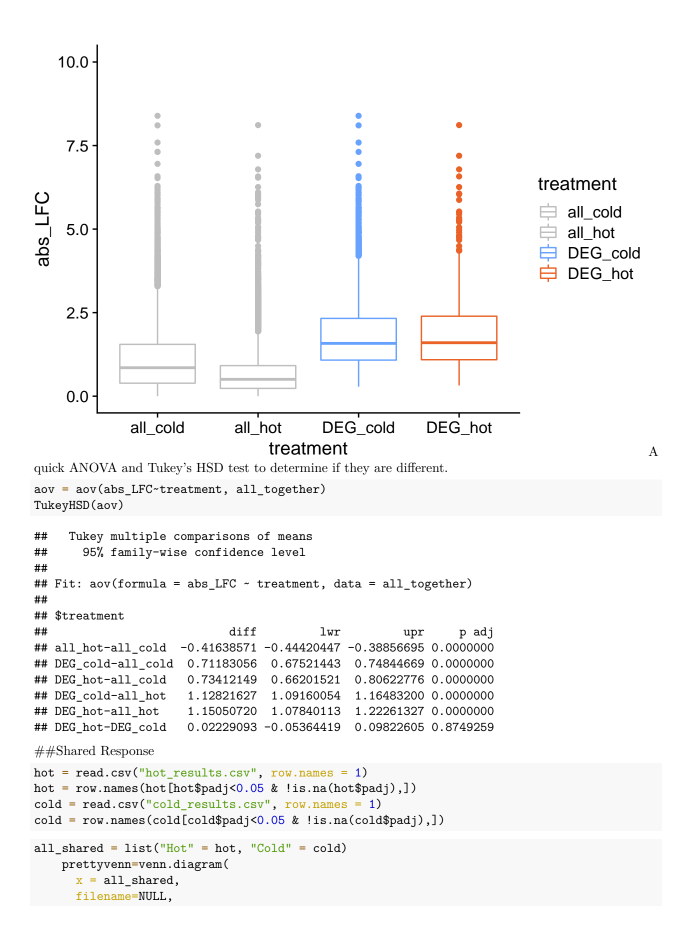
Significance

```
adonis(hot_pca$x ~ hot_treatment + hot_genotype, method = 'eu')
```

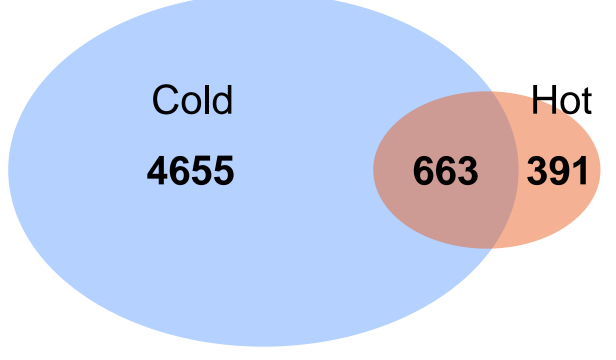
```
##
## Call:
## adonis(formula = hot_pca$x ~ hot_treatment + hot_genotype, method = "eu")
##
## Permutation: free
## Number of permutations: 999
##
## Terms added sequentially (first to last)
##
## Df SumsOfSqs MeanSqs F.Model R2 Pr(>F)
## hot_treatment 1 19385 19385.3 4.5248 0.13748 0.001 ***
## hot_genotype 8 65922 8240.3 1.9234 0.46753 0.001 ***
```

```
## Residuals
               13
                       55695 4284.2
                                              0.39499
## Total
                 22
                       141002
                                              1.00000
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
Here we are making a supplemental figure of overall LFC for each experiment
cold_results = read.csv("cold_results.csv")
hot_results = read.csv("hot_results.csv")
all LFC results = cold results %>%
  select(X, log2FoldChange) %>%
  dplyr::rename(all_cold = log2FoldChange) %>%
  left_join(hot_results) %>%
  select(X, all_cold, log2FoldChange) %>%
  dplyr::rename(all_hot = log2FoldChange) %>%
  gather(treatment, LFC, all_cold:all_hot )
## Joining, by = "X"
DEG_cold = cold_results %>%
  filter(padj < 0.05) %>%
  select(X, log2FoldChange) %>%
  dplyr::rename(DEG_cold =log2FoldChange)
DEG_hot = hot_results %>%
  filter(padj < 0.05) %>%
  select(X, log2FoldChange) %>%
  dplyr::rename(DEG_hot =log2FoldChange)
DEG_LFC_results = DEG_cold %>%
  full_join(DEG_hot) %>%
  gather(treatment, LFC, DEG_cold:DEG_hot)
## Joining, by = "X"
all_together = all_LFC_results %>%
  full_join(DEG_LFC_results) %>%
  mutate(abs_LFC = abs(LFC))
## Joining, by = c("X", "treatment", "LFC")
cols2 = c("DEG_hot" = "#ea6227", "all_hot" = "grey", "DEG_cold" = "#68a2ff", "all_cold" = "grey")
ggplot(all_together, aes(x = treatment, y = abs_LFC)) +
  geom_boxplot(aes(colour = treatment)) +
  scale_colour_manual(values = cols2) +
  ylim(0, 10) +
  theme_cowplot()
```

Warning: Removed 5065 rows containing non-finite values (stat_boxplot).



```
col = "transparent",
    fill = c("#ea6227", "#68a2ff"),
    alpha = 0.5,
    # label.col = c("darkred", "white", "darkgreen", "white", "white", "white", "blue4"),
    cex = 2.5,
    fontfamily = "sans",
    fontface = "bold",
    cat.default.pos = "text",
    cat.col = "black",
    cat.cex = 2.5,
    cat.fontfamily = "sans",
    cat.dist = c(0.08, 0.08),
    cat.pos = 1
    );
grid.draw(prettyvenn)
```



Hypergeometric test

```
a = read.csv("hot_results.csv")
h = read.csv("hot_results.csv") %>%
  filter(padj < 0.05)

c = read.csv("cold_results.csv") %>%
  filter(padj < 0.05)

overlap = inner_join(h, c, by = "X")

phyper((nrow(overlap)-1), nrow(h), (nrow(a)-nrow(h)), nrow(c), lower.tail = F, log.p = FALSE)

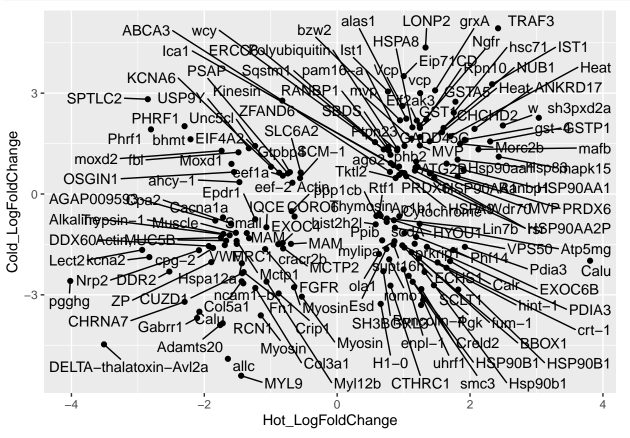
## [1] 1.183221e-52</pre>
```

```
gene = read.delim("astrangia_iso2gene.tab", sep = "\t") %>%
  mutate(gene_symbol = gsub(".* GN=", "", Gene)) %>%  # Remove everything before OX=
  mutate(gene_symbol = gsub(".*", "", gene_symbol)) # Remove everything after species ID

plot = overlap %>%
  rename(Iso = X) %>%
  inner_join(gene) %>%
  dplyr::select(gene_symbol, log2FoldChange.x, log2FoldChange.y) %>%
  rename(Hot_LogFoldChange = log2FoldChange.x) %>%
  rename(Cold_LogFoldChange = log2FoldChange.y)
```

Plotting delta ranks, this figure does not appear like this in the main manuscript. Only select genes are highlighted, for interactive gene plot visit my website: www.wuitchik.weebly.com/bioinformatics.html

```
delta_ranks = ggplot(plot, aes(Hot_LogFoldChange, Cold_LogFoldChange, label = gene_symbol)) +
   geom_point() +
   geom_text_repel()
delta_ranks
```



#GO Analysis

We are going to do a Mann-Whitney U test which requires that we first -logged the pvalue, and multiply it by -1 if it's less than zero or by 1 if it's greater than zero. Note that this needs to be opened and saved in excel (no other changes) as I believe the way R is saving the file the unicode does not work with Misha's script.

```
cold_go_input = read.csv("cold_results.csv") %>%
  mutate(mutated_p = -log(pvalue)) %>%
  mutate(mutated_p_updown = ifelse(log2FoldChange < 0, mutated_p*-1, mutated_p*1)) %>%
```

```
dplyr::select(X, mutated_p_updown) %>%
  na.omit()

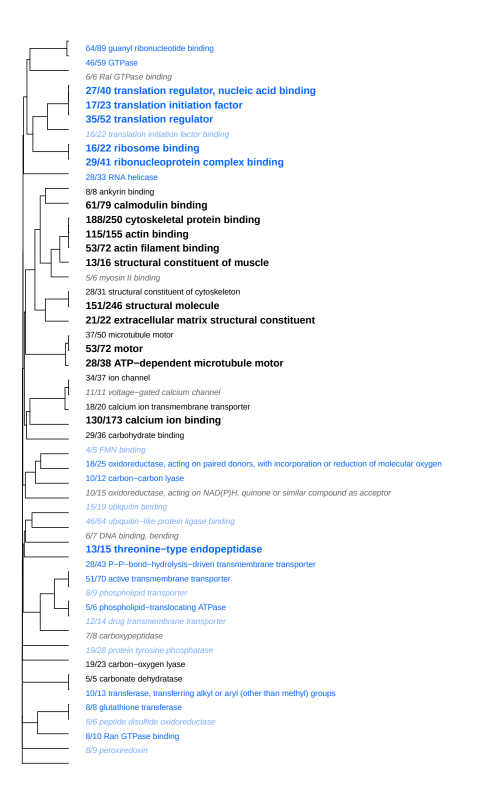
colnames(cold_go_input) = NULL

write.csv(cold_go_input, "cold_go_input.csv", row.names = FALSE)
```

Cold

Molecular functions

These code are all adapted from Dr. Matz and can be found GO_MWU https://github.com/z0on/GO_MWU This will be broken down into three sections

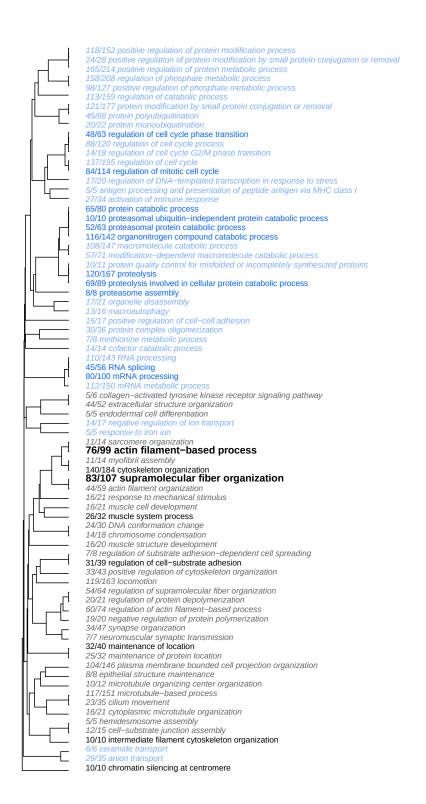


p < 0.01

p < 0.05

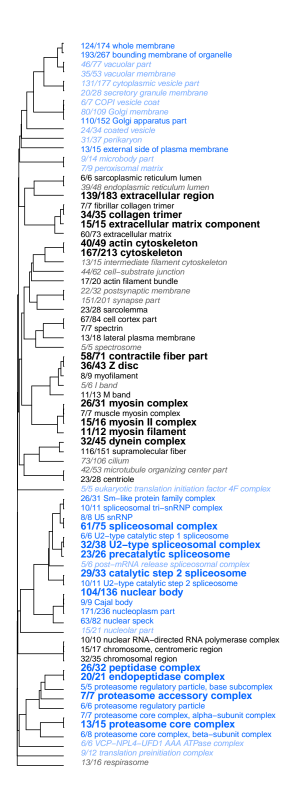
p < 0.1

Biological Process



p < 0.001 p < 0.01 p < 0.05

Cellular Component



p < 0.001 p < 0.01 p < 0.05

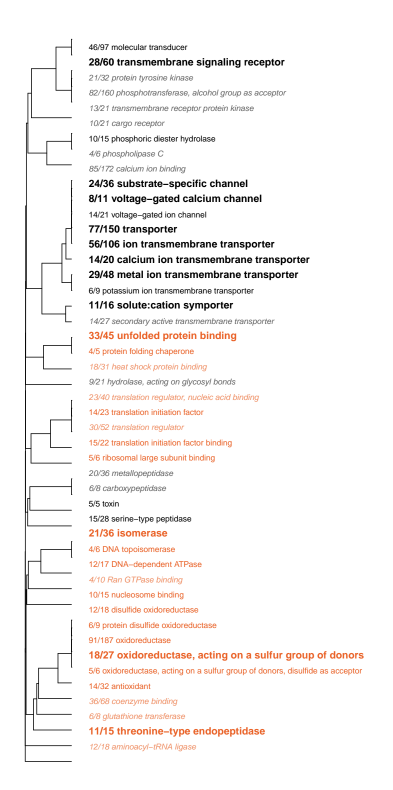
Hot

```
hot_go_input = read.csv("hot_results.csv") %>%
  mutate(mutated_p = -log(pvalue)) %>%
  mutate(mutated_p_updown = ifelse(log2FoldChange < 0, mutated_p*-1, mutated_p*1)) %>%
  dplyr::select(X, mutated_p_updown) %>%
  na.omit()

colnames(hot_go_input) = NULL

write.csv(hot_go_input, "hot_go_input.csv", row.names = FALSE)
```

Molecular functions

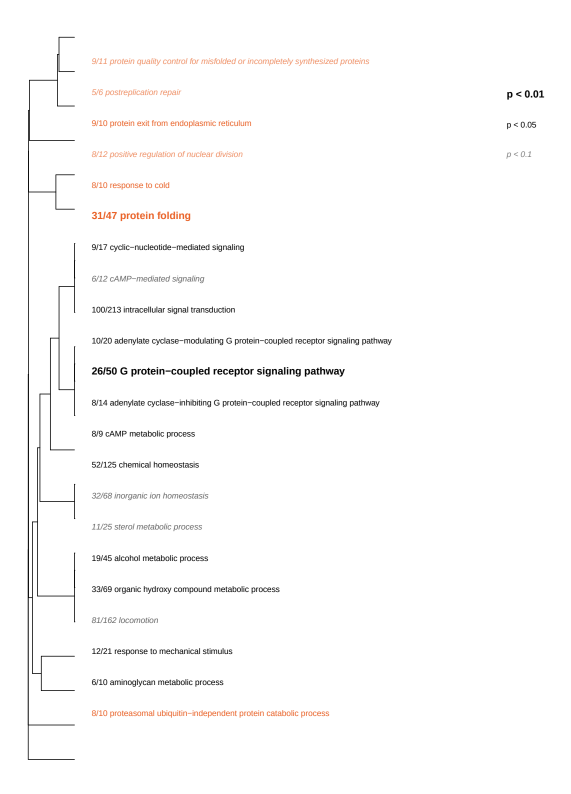


p < 0.01

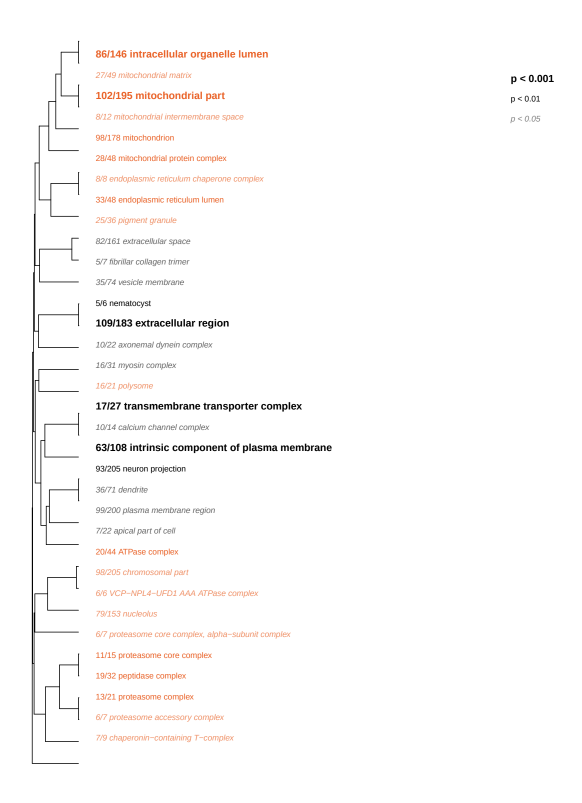
p < 0.05

p < 0.1

Biological Process



Cellular Component

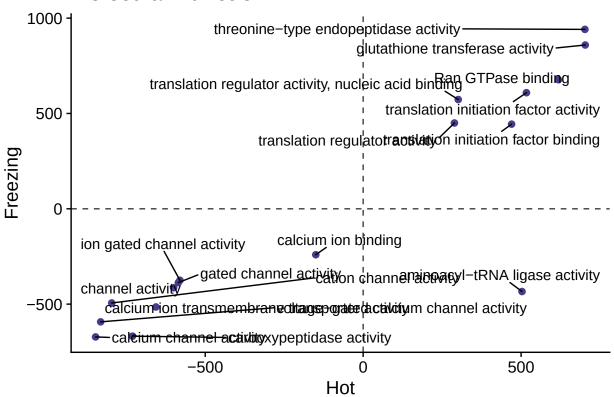


Comparing Hot and Cold

Molecular Functions

```
mf_hotMWU =read.table("Supps/MWU_MF_hot_go_input_excel_saved.csv",header=T)
mf_coldMWU =read.table("Supps/MWU_MF_cold_go_input_excel_saved.csv", header=T)
# Terms in both sets
mf_goods=intersect(mf_hotMWU$term,mf_coldMWU$term)
data1=mf_hotMWU[mf_hotMWU$term %in% mf_goods,]
data2=mf_coldMWU[mf_coldMWU$term %in% mf_goods,]
# Combine them
ress=merge(data1,data2,by="term")
plot = ress %>%
  mutate(colour =
           case_when( p.adj.x < 0.1 & p.adj.y < 0.1 & delta.rank.x > 0 & delta.rank.y > 0 ~ 'red',
                      p.adj.x < 0.1 \& p.adj.y < 0.1 \& delta.rank.x < 0 \& delta.rank.y < 0 ~ 'blue',
                      p.adj.x < 0.1 & p.adj.y < 0.1 & delta.rank.x > 0 & delta.rank.y < 0 ~ 'purple',
                      p.adj.x < 0.1 \& p.adj.y < 0.1 \& delta.rank.x < 0 \& delta.rank.y > 0 ~ 'green')) %
  replace_na(list(colour = "black"))
# This is to manually look for interesting go terms, and you can play with it in excel
mf_interest = plot %>%
  filter(p.adj.x <0.1) %>%
  filter(p.adj.y < 0.1)
write.csv(mf_interest, "mf_interesting.csv")
# Read back in your manipulated csv for those that you want to use as labels
mf_interest = read.csv("mf_interesting.csv")
# Here is the actual plot, lots of it is redundant
mf_plot = ggplot(mf_interest, aes(delta.rank.x, delta.rank.y, label = name.y)) +
  geom_point(aes(color = colour), size = 2, show.legend = FALSE) +
  scale_color_manual(values = c(red = "darkslateblue",
                                blue = "darkslateblue",
                                green = "darkslateblue",
                                red = "darkslateblue",
                                purple = "darkslateblue",
                                black = alpha("black", 0.15))) +
  scale_fill_manual(values = c(red = "orangered",
                               blue = "dodgerblue2",
                               green = "seagreen3",
                               red = "orangered2",
                               purple = "plum2",
                              black = "black")) +
    geom_text_repel(data = mf_interest, aes(),
                   segment.alpha = 1,
                   box.padding = .5,
                   direction = "both") +
  scale_size("size") +
```

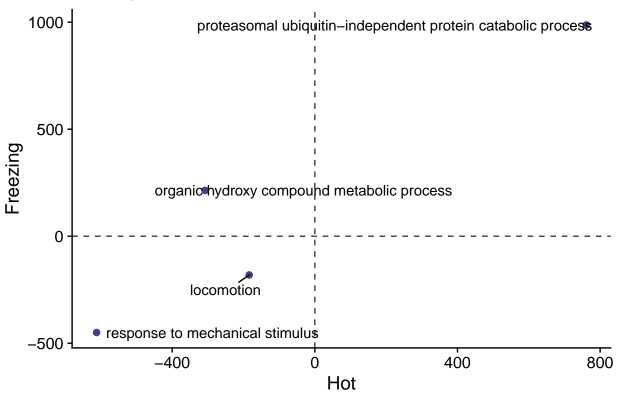
Molecular Function



Biological Process

```
p.adj.x < 0.1 \& p.adj.y < 0.1 \& delta.rank.x < 0 & delta.rank.y > 0 ~ 'green')) %
  replace_na(list(colour = "black"))
# This is to manually look for interesting go terms, and you can play with it in excel
bp_interest = plot %>%
 filter(p.adj.x <0.1) %>%
 filter(p.adj.y < 0.1)
write.csv(bp_interest, "bp_interesting.csv")
# Read back in your manipulated csv for those that you want to use as labels
bp_interest = read.csv("bp_interesting.csv")
# Here is the actual plot, lots of it is redundant
bp_plot = ggplot(bp_interest, aes(delta.rank.x, delta.rank.y, label = name.y)) +
 geom_point(aes(color = colour), size = 2, show.legend = FALSE) +
  scale_color_manual(values = c(red = "darkslateblue",
                                blue = "darkslateblue",
                                green = "darkslateblue",
                                red = "darkslateblue",
                                purple = "darkslateblue",
                                black = alpha("black", 0.15))) +
  scale_fill_manual(values = c(red = "orangered",
                               blue = "dodgerblue2",
                               green = "seagreen3",
                               red = "orangered2",
                               purple = "plum2",
                              black = "black")) +
   geom_text_repel(data = bp_interest, aes(),
                   segment.alpha = 1,
                   box.padding = .5,
                   direction = "both") +
  scale_size("size") +
  labs(x = "Hot",
        y = "Freezing") +
  labs(title = "Biological Process") +
  geom_vline(xintercept = 0, linetype = 2, alpha = 0.75) +
  geom_hline(yintercept = 0, linetype = 2, alpha = 0.75) +
  theme_cowplot()
bp_plot
```

Biological Process

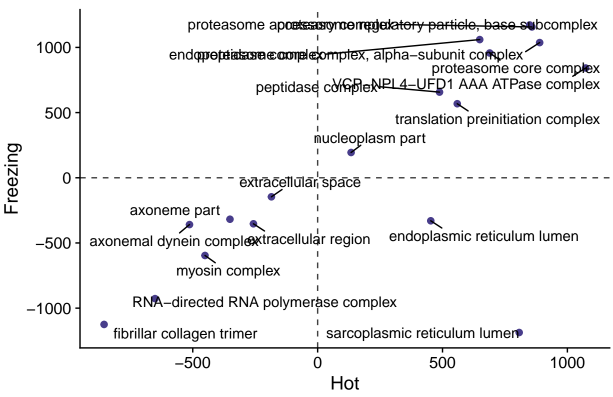


Cellular Components

```
cc_hotMWU =read.table("MWU_cc_hot_go_input_excel_saved.csv",header=T)
cc_coldMWU =read.table("MWU_cc_cold_go_input_excel_saved.csv", header=T)
# Terms in both sets
cc_goods=intersect(cc_hotMWU$term,cc_coldMWU$term)
data1=cc hotMWU[cc hotMWU$term %in% cc goods,]
data2=cc_coldMWU[cc_coldMWU$term %in% cc_goods,]
# Combine them
ress=merge(data1,data2,by="term")
plot = ress %>%
  mutate(colour =
           case_when( p.adj.x < 0.1 & p.adj.y < 0.1 & delta.rank.x > 0 & delta.rank.y > 0 ~ 'red',
                      p.adj.x < 0.1 \& p.adj.y < 0.1 \& delta.rank.x < 0 \& delta.rank.y < 0 ~ 'blue',
                      p.adj.x < 0.1 & p.adj.y < 0.1 & delta.rank.x > 0 & delta.rank.y < 0 ~ 'purple',
                      p.adj.x < 0.1 \& p.adj.y < 0.1 \& delta.rank.x < 0 \& delta.rank.y > 0 ~ 'green')) %
 replace_na(list(colour = "black"))
# This is to manually look for interesting go terms, and you can play with it in excel
cc_interest = plot %>%
  filter(p.adj.x <0.1) %>%
  filter(p.adj.y < 0.1)
write.csv(cc_interest, "cc_interesting.csv")
```

```
# Read back in your manipulated csv for those that you want to use as labels
cc_interest = read.csv("cc_interesting.csv")
# Here is the actual plot, lots of it is redundant
cc_plot = ggplot(cc_interest, aes(delta.rank.x, delta.rank.y, label = name.y)) +
 geom_point(aes(color = colour), size = 2, show.legend = FALSE) +
 scale color manual(values = c(red = "darkslateblue",
                                blue = "darkslateblue",
                                green = "darkslateblue",
                                red = "darkslateblue",
                                purple = "darkslateblue",
                                black = alpha("black", 0.15))) +
  scale_fill_manual(values = c(red = "orangered",
                               blue = "dodgerblue2",
                               green = "seagreen3",
                               red = "orangered2",
                               purple = "plum2",
                              black = "black")) +
   geom_text_repel(data = cc_interest, aes(),
                   segment.alpha = 1,
                   box.padding = .5,
                   direction = "both") +
  scale_size("size") +
 labs( x = "Hot",
       y = "Freezing") +
  labs(title = "Cellular Components") +
  geom_vline(xintercept = 0, linetype = 2, alpha = 0.75) +
  geom_hline(yintercept = 0, linetype = 2, alpha = 0.75) +
  theme_cowplot()
cc_plot
```

Cellular Components



Heatmaps

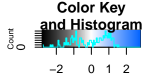
The purpose of these heatmaps is not to provide comprehensive heatmaps used in the manuscript. Rather, it is to highlight the code used to form a basis on how it was applied for each individual heatmap.

```
iso2go = read_tsv("astrangia_iso2go.tab") %>%
  dplyr::rename(Iso = Gene id)
cold_results_df = read.csv("cold_results.csv") %>%
  dplyr::rename("Iso" = "X")
hot_results_df = read.csv("hot_results.csv") %>%
  dplvr::rename("Iso" = "X")
cold_rlog = read.csv("cold_rlogged.csv") %>%
  dplyr::rename("Iso" = "X") %>%
  left_join(cold_results_df) %>%
  filter(padj < 0.1) %>%
  dplyr::select(-baseMean, -log2FoldChange, -lfcSE, -stat, -pvalue, -padj)
hot_rlog = read.csv("hot_rlogged.csv") %>%
  dplyr::rename("Iso" = "X") %>%
  left_join(hot_results_df) %>%
  filter(padj < 0.1) %>%
  dplyr::select(-baseMean, -log2FoldChange, -lfcSE, -stat, -pvalue, -padj)
hot_colour = colorRampPalette(rev(c("#ea6227","#f09167","white", "grey40","black")))(100)
```

```
cold_colour = colorRampPalette(rev(c("#0666ff","#7caeff","white", "grey40","black")))(100)
```

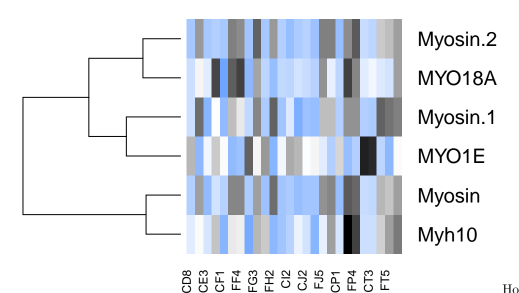
GO:0016459 myosin complex Cold

```
GO_0016459_cold = iso2go %>%
  filter(str_detect(GO_id, "GO:0016459")) %>%
  left_join(gene) %>%
  left_join(cold_rlog) %>%
 mutate(gene_symbol = make.names(gene_symbol, unique = TRUE)) %>%
  column_to_rownames(var = "gene_symbol") %>%
  dplyr::select(-GO_id, -Gene, -Iso) %>%
  drop_na()
  #dplyr::select(sort(current_vars()))
GO_0016459_cold_means=apply(GO_0016459_cold,1,mean) # means of rows
explc=GO_0016459_cold-GO_0016459_cold_means # subtracting them
heatmap.2(as.matrix(GO_0016459_cold), col = cold_colour, Rowv = TRUE, Colv = FALSE, scale = "row",
          dendrogram = "both",
          trace = "none",
         main = "GO:0016459 myosin complex",
         margin = c(5,15))
```



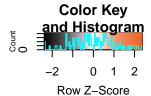
Row Z-Score

016459 myosin complex

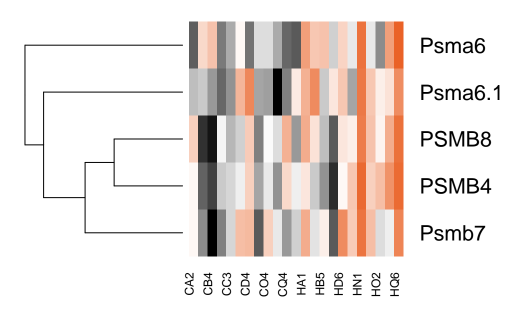


 $GO_0010499_hot = iso2go \%>\%$ filter(str_detect(GO_id, "GO:0010499")) %>% left_join(gene) %>% left_join(hot_rlog) %>% mutate(gene_symbol = make.names(gene_symbol, unique = TRUE)) %>%

Hot



piquitin-independent protein catabolic p



Comparison with Dixon et al. (2020) meta analaysis

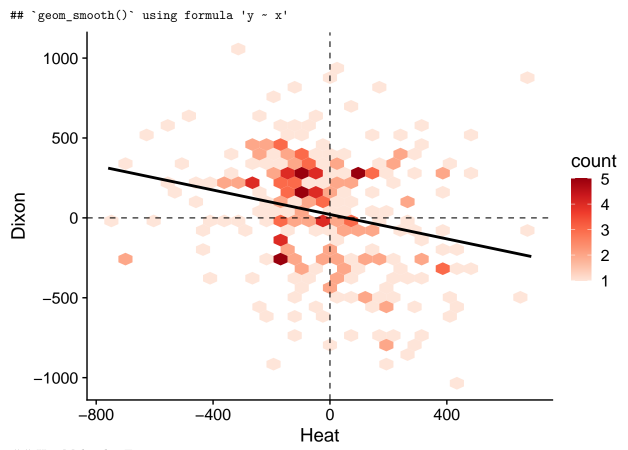
```
Import data
```

```
Dixon_MF = read.table("Dixon_MF.csv", header = T)
Dixon_CC = read.table("Dixon_CC.csv", header = T)
Dixon_BP = read.table("Dixon_BP.csv", header = T)
hot_MF = read.table("MWU_MF_hot_go_input_excel_saved.csv", header = T)
hot_CC = read.table("MWU_CC_hot_go_input_excel_saved.csv", header = T)
hot_BP = read.table("MWU_BP_hot_go_input_excel_saved.csv", header = T)
```

```
cold_MF = read.table("MWU_MF_cold_go_input_excel_saved.csv", header = T)
cold_CC = read.table("MWU_CC_cold_go_input_excel_saved.csv", header = T)
cold_BP = read.table("MWU_BP_cold_go_input_excel_saved.csv", header = T)
```

Hot Cellular component

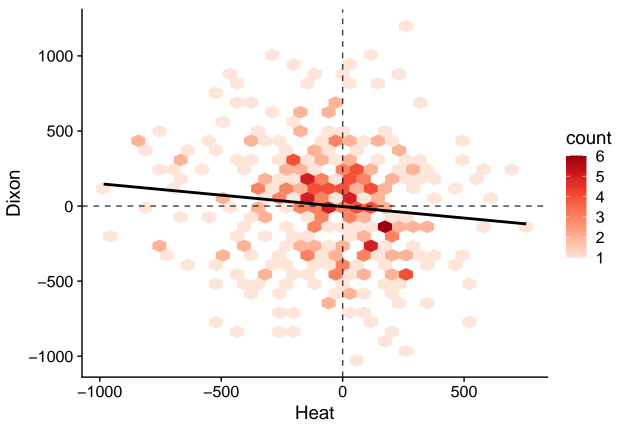
```
# Terms in both sets
hot_cc_goods=intersect(hot_CC$term,Dixon_CC$term)
data1=hot_CC[hot_CC$term %in% hot_cc_goods,]
data2=Dixon_CC[Dixon_CC$term %in% hot_cc_goods,]
# Combine them
hot_cc_plot=merge(data1,data2,by="term")
cc_hot_plot = ggplot(hot_cc_plot, aes(delta.rank.x, delta.rank.y, label = name.y)) +
  scale_fill_distiller(palette = "Reds", direction = 1) +
  geom_hex() +
  labs(x = "Heat",
       y = "Dixon") +
  geom_vline(xintercept = 0, linetype = 2, alpha = 0.75) +
  geom_hline(yintercept = 0, linetype = 2, alpha = 0.75) +
  geom_smooth(method=lm, color="black", se =FALSE) +
  theme_cowplot()
cc_hot_plot
```



Hot Molecular Function

```
# Terms in both sets
hot_MF_goods=intersect(hot_MF$term,Dixon_MF$term)
hot MF data1=hot MF[hot MF$term %in% hot MF goods,]
hot_MF_data2=Dixon_MF[Dixon_MF$term %in% hot_MF_goods,]
# Combine them
hot_MF_plot=merge(hot_MF_data1,hot_MF_data2,by="term")
MF_hot_plot = ggplot(hot_MF_plot, aes(delta.rank.x, delta.rank.y, label = name.y)) +
  scale_fill_distiller(palette = "Reds", direction = 1) +
  geom_hex()+
  scale_size("size") +
  labs(x = "Heat",
       y = "Dixon") +
  geom_vline(xintercept = 0, linetype = 2, alpha = 0.75) +
  geom_hline(yintercept = 0, linetype = 2, alpha = 0.75) +
  geom_smooth(method=lm, color="black", se =FALSE) +
  theme_cowplot()
MF_hot_plot
```

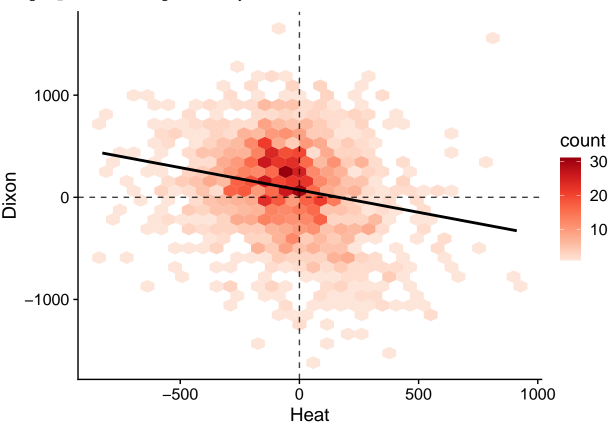
`geom_smooth()` using formula 'y ~ x'



Hot Biological Process

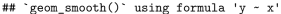
```
# Terms in both sets
hot_BP_goods=intersect(hot_BP$term,Dixon_BP$term)
hot_BP_data1=hot_BP[hot_BP$term %in% hot_BP_goods,]
hot_BP_data2=Dixon_BP[Dixon_BP$term %in% hot_BP_goods,]
```

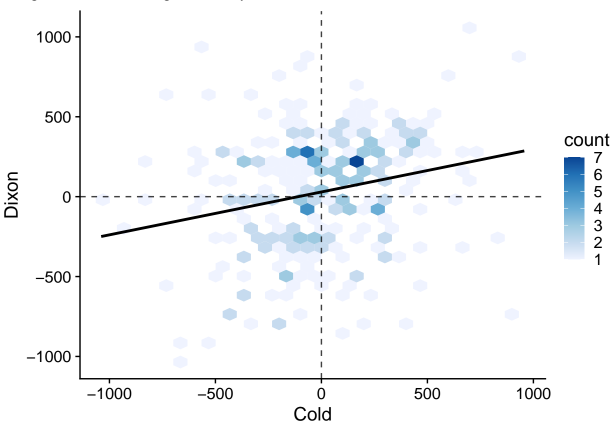
`geom_smooth()` using formula 'y ~ x'



Cold Cellular Component

```
# Terms in both sets
cold_cc_goods=intersect(cold_CC$term,Dixon_CC$term)
data1=cold_CC[cold_CC$term %in% cold_cc_goods,]
data2=Dixon_CC[Dixon_CC$term %in% cold_cc_goods,]
# Combine them
```



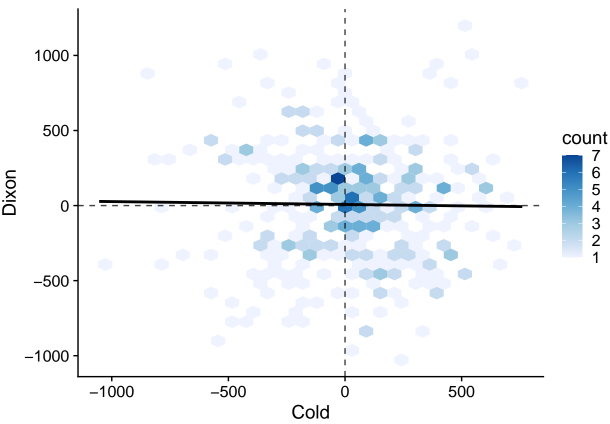


Cold Molecular Function

```
# Terms in both sets
cold_MF_goods=intersect(cold_MF$term,Dixon_MF$term)
cold_MF_data1=cold_MF[cold_MF$term %in% cold_MF_goods,]
cold_MF_data2=Dixon_MF[Dixon_MF$term %in% cold_MF_goods,]

# Combine them
cold_MF_plot=merge(cold_MF_data1,cold_MF_data2,by="term")
```

$geom_smooth()$ using formula 'y ~ x'



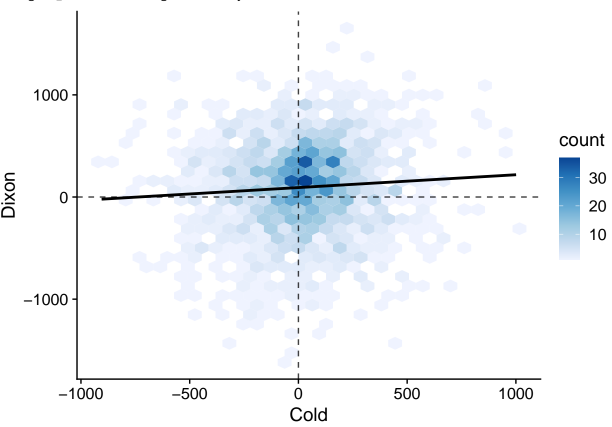
Cold Biological Process

```
# Terms in both sets
cold_BP_goods=intersect(cold_BP$term,Dixon_BP$term)
cold_BP_data1=cold_BP[cold_BP$term %in% cold_BP_goods,]
cold_BP_data2=Dixon_BP[Dixon_BP$term %in% cold_BP_goods,]

# Combine them
cold_BP_plot=merge(cold_BP_data1,cold_BP_data2,by="term")

BP_cold_plot = ggplot(cold_BP_plot, aes(delta.rank.x, delta.rank.y, label = name.y)) +
    scale_fill_distiller(palette = "Blues", direction = 1) +
    geom_hex()+
```

`geom_smooth()` using formula 'y ~ x'



All together now

```
ggarrange(BP_cold_plot, BP_hot_plot, MF_cold_plot, MF_hot_plot, cc_cold_plot, cc_hot_plot, ncol = 2, nr
## `geom_smooth()` using formula 'y ~ x'
```

```
Dixon B
Dixon A
                                                                                      30
     1000
                                                   1000
                                         30
        0
                                                      0
                                                                                      20
                                         20
   -1000
                                                 -1000
                                                                                      10
                                         10
        -1000-500
                         500 1000
                                                                       500 1000
                     0
                                                           -500
                                                                   0
                   Cold
                                                                 Heat
                                    count
                                                                                  count
                                         7
                                                                                       6
     1000 -
                                                   1000 -
                                         6
                                                                                       5
      500
                                                    500
                                         5
        0
                                                      0
                                                                                       4
                                         4
     -500
                                                  -500
                                                                                       3
                                         3
   -1000 -
                                                 -1000 -
                                                                                       2
                                         2
          -1000-500
                            500
                                                       -1000-500
                                                                         500
                       0
                                                                     0
                                                                                       1
                    Cold
                                                                 Heat
                                    count
                                                                                  count
                                         7
                                                                                       5
     1000 -
                                                   1000 -
Dixon |
                                         6
                                              Dixon
      500
                                                    500
                                                                                       4
                                         5
        0
                                                      0
                                         4
3
2
                                                                                       3
                                                   -500
    -500
   -1000
                                                 -1000
                                                                                       2
                          500 1000
                                                                         400
         -1000-500
                      0
                                                       -800 - 400
                                                                    0
                                                                                       1
                   Cold
                                                                 Heat
#Session Info
sessionInfo()
## R version 4.0.2 (2020-06-22)
## Platform: x86_64-apple-darwin17.0 (64-bit)
## Running under: macOS 10.16
##
## Matrix products: default
           /Library/Frameworks/R.framework/Versions/4.0/Resources/lib/libRblas.dylib
## LAPACK: /Library/Frameworks/R.framework/Versions/4.0/Resources/lib/libRlapack.dylib
## locale:
  [1] en_US.UTF-8/en_US.UTF-8/en_US.UTF-8/C/en_US.UTF-8/en_US.UTF-8
##
## attached base packages:
##
    [1] grid
                  parallel
                            stats4
                                       stats
                                                 graphics grDevices utils
    [8] datasets methods
                            base
##
##
## other attached packages:
   [1] ggpubr_0.4.0
                                    forcats_0.5.1
##
##
   [3] dplyr_1.0.6
                                    purrr_0.3.4
    [5] tidyr_1.1.3
                                    tibble_3.1.2
##
                                    brms_2.15.0
##
   [7] tidyverse_1.3.1
   [9] Rcpp_1.0.6
                                    stringr 1.4.0
## [11] ggrepel_0.9.1
                                    VennDiagram_1.6.20
  [13] futile.logger_1.4.3
                                    prettydoc 0.4.1
## [15] reshape2_1.4.4
                                    kableExtra_1.3.4
```

```
## [17] vegan_2.5-7
                                    permute 0.9-5
## [19] plotly_4.9.3
                                    knitr 1.33
## [21] gplots_3.1.1
                                    RColorBrewer 1.1-2
                                    cowplot_1.1.1
## [23] readr_1.4.0
## [25] DESeq_1.39.0
                                    lattice_0.20-44
## [27] locfit 1.5-9.4
                                    genefilter 1.72.1
## [29] arrayQualityMetrics 3.46.0
                                    affycoretools 1.62.0
## [31] ggplot2_3.3.3
                                    DESeq2_1.30.1
## [33] SummarizedExperiment_1.20.0 Biobase_2.50.0
## [35] MatrixGenerics_1.2.1
                                    matrixStats_0.59.0
## [37] GenomicRanges_1.42.0
                                    GenomeInfoDb_1.26.7
## [39] IRanges_2.24.1
                                    S4Vectors_0.28.1
## [41] BiocGenerics_0.36.1
##
## loaded via a namespace (and not attached):
##
     [1] Hmisc_4.5-0
                                  svglite_2.0.0
                                                            ps_1.6.0
##
     [4] Rsamtools_2.6.0
                                  foreach_1.5.1
                                                            projpred_2.0.2
                                                            MASS_7.3-54
     [7] crayon 1.4.1
                                  V8 3.4.2
  [10] nlme_3.1-152
                                                            reprex_2.0.0
                                  backports_1.2.1
   [13] colourpicker 1.1.0
                                  rlang_0.4.11
                                                            readxl_1.3.1
## [16] XVector_0.30.0
                                  nloptr_1.2.2.2
                                                            callr_3.7.0
                                                            GOstats 2.56.0
## [19] limma_3.46.0
                                  gridSVG_1.7-2
## [22] BiocParallel_1.24.1
                                  bit64_4.0.5
                                                            glue_1.4.2
## [25] loo_2.4.1
                                  rstan_2.21.2
                                                            processx 3.5.2
## [28] AnnotationDbi_1.52.0
                                  vsn 3.58.0
                                                            haven_2.4.1
## [31] tidyselect_1.1.1
                                  rio_0.5.26
                                                            XML_3.99-0.6
## [34] zoo_1.8-9
                                  GenomicAlignments_1.26.0 xtable_1.8-4
## [37] magrittr_2.0.1
                                  evaluate_0.14
                                                            cli_2.5.0
## [40] zlibbioc_1.36.0
                                  hwriter_1.3.2
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                                                            ensembldb_2.14.1
## [43] miniUI_0.1.1.1
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##
   [46] lambda.r_1.2.4
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                                                            shiny_1.6.0
## [49] xfun_0.23
                                  askpass_1.1
                                                            inline_0.3.19
  [52] pkgbuild_1.2.0
                                  cluster_2.1.2
                                                            bridgesampling_1.1-2
## [55] caTools_1.18.2
                                                            ff_4.0.4
                                  Brobdingnag_1.2-6
   [58] base64 2.0
                                                            threejs_0.3.3
                                  biovizBase_1.38.0
## [61] Biostrings_2.58.0
                                  png_0.1-7
                                                            reshape_0.8.8
## [64] withr 2.4.2
                                  bitops 1.0-7
                                                            cellranger 1.1.0
## [67] RBGL_1.66.0
                                  plyr_1.8.6
                                                            GSEABase_1.52.1
## [70] AnnotationFilter_1.14.0
                                  PFAM.db_3.12.0
                                                            coda_0.19-4
## [73] pillar_1.6.1
                                  RcppParallel_5.1.4
                                                            cachem_1.0.5
## [76] GenomicFeatures 1.42.3
                                  fs 1.5.0
                                                            xts_0.12.1
## [79] vctrs_0.3.8
                                  ellipsis_0.3.2
                                                            generics_0.1.0
## [82] dygraphs_1.1.1.6
                                  tools_4.0.2
                                                            gcrma_2.62.0
## [85] foreign_0.8-81
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                                                            munsell_0.5.0
## [88] gamm4_0.2-6
                                  emmeans_1.6.1
                                                            DelayedArray_0.16.3
## [91] fastmap_1.1.0
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                                                            abind_1.4-5
## [94] httpuv_1.6.1
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## [97] gridExtra_2.3
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                                                            BiocFileCache_1.14.0
## [103] jsonlite_1.7.2
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                                                            GGally_2.1.1
## [106] scales_1.1.1
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                                  graph_1.68.0
## [109] estimability_1.3
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## [112] promises_1.2.0.1
                                  car_3.0-10
                                                            latticeExtra_0.6-29
## [115] R.utils_2.10.1
                                  checkmate 2.0.0
                                                            openxlsx_4.2.3
```

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##	[121]	BSgenome_1.58.0	igraph_1.2.6	survival_3.2-11
##	[124]	rsconnect_0.8.18	yaml_2.2.1	systemfonts_1.0.2
##	[127]	Glimma_2.0.0	bayesplot_1.8.0	htmltools_0.5.1.1
##	[130]	rstantools_2.1.1	memoise_2.0.0	VariantAnnotation_1.36.0
##	[133]	viridisLite_0.4.0	digest_0.6.27	assertthat_0.2.1
##	[136]	mime_0.10	rappdirs_0.3.3	futile.options_1.0.1
##	[139]	RSQLite_2.2.7	beadarray_2.40.0	data.table_1.14.0
##	[142]	blob_1.2.1	R.oo_1.24.0	ReportingTools_2.30.2
##	[145]	preprocessCore_1.52.1	labeling_0.4.2	shinythemes_1.2.0
##	[148]	splines_4.0.2	Formula_1.2-4	illuminaio_0.32.0
##	[151]	OrganismDbi_1.32.0	ProtGenerics_1.22.0	RCurl_1.98-1.3
##		broom_0.7.6	hms_1.1.0	modelr_0.1.8
##		colorspace_2.0-1	base64enc_0.1-3	BiocManager_1.30.15
##		nnet_7.3-16	mvtnorm_1.1-1	fansi_0.5.0
##		R6_2.5.0	ggridges_0.5.3	lifecycle_1.0.0
##		formatR_1.10	StanHeaders_2.21.0-7	zip_2.2.0
##		ggsignif_0.6.1	curl_4.3.1	minqa_1.2.4
##		affyio_1.60.0	Matrix_1.3-3	ggbio_1.38.0
		BeadDataPackR_1.42.0	iterators_1.0.13	htmlwidgets_1.5.3
##		biomaRt_2.46.3	markdown_1.1	crosstalk_1.1.1
##		rvest_1.0.0	mgcv_1.8-35	openssl_1.4.4
##		htmlTable_2.2.1	<pre>lubridate_1.7.10</pre>	codetools_0.2-18
##		GO.db_3.12.1	gtools_3.8.2	<pre>prettyunits_1.1.1</pre>
##		dbplyr_2.1.1	R.methodsS3_1.8.1	gtable_0.3.0
##		DBI_1.1.1	highr_0.9	httr_1.4.2
##		KernSmooth_2.23-20	stringi_1.6.2	oligoClasses_1.52.0
##		progress_1.2.2	farver_2.1.0	annotate_1.68.0
##		hexbin_1.28.2	Rgraphviz_2.34.0	DT_0.18
##		xml2_1.3.2	boot_1.3-28	shinyjs_2.0.0
##		lme4_1.1-27	geneplotter_1.68.0	Category_2.56.0
##		bit_4.0.4	jpeg_0.1-8.1	pkgconfig_2.0.3
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