# Find differentially expressed genes using DESeq2 library functions and reference genome assembly SL4 from September 2019

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### Introduction

This Markdown analyzes RNA-Seq gene expression data from data file results/muday-144-SL4\_counts-salmon.txt documented in Documentation/muday-144\_sample\_sheet.xlsx using differential expression analysis library DESeq2.

The input data file was generated by nf-core/rna-seq pipeline that aligned RNA-Seq data and produced RNA-Seq fragment counts for gene annotations from the September 2019 release of the tomato genome known as "SL4".

In this Markdown, we aim to answer:

• How many genes were differentially expressed in treatment versus control comparisons?

# Input data file summary:

The git hash (version) of results/muday-144-SL4\_counts-salmon.txt was: 62fdd9c2ca32374188634d2d8ba60f9c9413b27d

The data file contains heat-treated and non-heat-treated samples from four tomato genotypes: are (anthocyanin reduced mutant), VF36 (wild-type tomoato cultivar), and a VF36 line designated F3H containing a transgene encoding the are wild-type gene. Samples experienced the heat stress over a time course which included four time points.

In addition to answering the above question, we also aim to create a data file that we will use to compare differential expression results obtained here with differential expression results obtained using a different, but similar, R library called "edgeR." If the results are similar, we can be more confident that we are using these libraries correctly.

# Analysis and Results

Load custom functions for analyzing and visualizing differential expression:

source("Common.R")

Load required counts data:

counts=getCounts(counts\_fname,keep\_description = T)

The table of RNA-Seq counts per gene loaded in the previous code chunk from file results/muday-144-SL4 counts-salmon.txt contained 34,075 rows corresponding to measured genes.

Define false discovery rate threshold for deciding whether a gene is differentially expressed:

```
Q=0.05
```

We will use 0.05, defined in the previous code chunk, to "call" a gene as differentially expressed.

As we test whether the treatment changed gene expression for each genotype and treatment duration combination, we will save results to a single data frame. At the end of the Markdown, we'll write this very large table to a file named results/CvT-DESeq2-SL4.txt.

## Genotype are, anthocyanin-reduced mutant

Comparing group A.28.15 to A.34.15 found 19 genes with changed expression levels with false discovery rate Q less than or equal to 0.05.

Comparing group A.28.30 to A.34.30 found 49 genes with changed expression levels with false discovery rate Q less than or equal to 0.05.

Comparing group A.28.45 to A.34.45 found 80 genes with changed expression levels with false discovery rate Q less than or equal to 0.05.

Comparing group A.28.75 to A.34.75 found 137 genes with changed expression levels with false discovery rate Q less than or equal to 0.05.

The number of DE genes called at Q less than 0.05increased with treatment duration time.

Display volcano plots that summarize the above results:

```
common.legend = T, # COMMON LEGEND
                                                                                                        legend = "right", # legend position
                                                                                                        align = "hv", # Align them both, horizontal and vertical
                                                                                                        nrow = 2,
                                                                                                        ncol = 2)
A_combined
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                        A.28.15 vs A.34.15
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                        Horizontal dashed line shows FDR 0.05
                                                                                                                                                                                                                                                            Horizontal dashed line shows FDR 0.05
          50
                                                                                                                                          Solvc06d072430.3
                                                                                                                                                                                                                                                                                                                                                                          Solyc04g011440.4
                                                                                                                                                                                                                                               40
                                                                                                                              Solyc11g066100.2
           40
                                                                                                                                                                                                                                                                                                                                                                      Solyc01g104740.3
                                                                                                                                                                                                                                              30
                                                                                                                                                                                                                                                                                                                                                                       Solyc08g062437.1
          30
                                                                                                                            Solyc03g115230.3
-Log<sub>10</sub> P
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                                                                                                                                                                                                                                                                                                                                                                               Solyc09g092690.3
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Solyc04g072160.3
          20
                                                                                                                                            Solyc08g078700.2
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Solyc18g0783700.2

Solyc18g078330.2

Solyc08g006880.3 Solyc08g082420.3

Solyc08g006880.3 Solyc03g082420.3

Solyc08g07630.1

Solyc08g076520.1
                                                                                                                                        Solyc08g062437.1
                                                                                                                                        Solyc04g011440.4
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           10
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Solyc01g081250.3
Solyc10g084170.1
Solyc06g076020.3
                                                                                                                                                                                                                                                                                                                                              Solyc16g076020.3 + Solyc13g118060.3 Solyc16g076020.3 + Solyc16g20330.1 Solyc16g66.4 - Solyc16g66
                           Solyc03g005555.1
Solyc05g053970.2
              0
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Mid-regulated
Up-regulated
                                                                                              Log<sub>2</sub> fold change
                                                                                                                                                                                                                                                                                                                                  Log<sub>2</sub> fold change
                                                                                                                                                                                                                                                                                                                                                                                                   total = 12673 variables
                                                                                                                                                                total = 17381 variables
C
                        A.28.45 vs A.34.45
                                                                                                                                                                                                                                    D
                                                                                                                                                                                                                                                           A.28.75 vs A.34.75
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                NS
Log<sub>2</sub> FC
                        Horizontal dashed line shows FDR 0.05
                                                                                                                                                                                                                                                           Horizontal dashed line shows FDR 0.05
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          20
                                                                                                                                                                                                                                                                                                                                                                                        Solvc08a062960.4
                                                                                                                                    Solyc02g077670.3
Solyc08g062960.4
           15
                                                                                                                                           Solyc09g008970.1
Solyc03g082420.3
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                                                                                                                                                                                                                                                                                                                                                                                Solyc03g117630.1
Solyc11g020040.2
Solyc09g015010.2
Solyc04g07/2160.3
Solyc06g083190.4
                                                                                                                                      Solyc04g072160.3
Solyc01g095320.4
                                                                                                                                                                                                                                  -Log<sub>10</sub> P
                                                                                                                                    Solyc03g118060.3
Solyc08g062340.3
Solyc03g007890.3
Solyc03g123540.3
          10
                                                                                                   Solyc02g079930.3

+Solyc05g007895.1

Solyc05g052410.3
```

The volcano plots confirm the previous observation that the number of genes found to be differentially expressed in response to the heat treatment increased with treatment duration.

Solyc03g083170.3 Solyc06g007300

Log<sub>2</sub> fold change

-5

10

0

total = 11800 variables

Log<sub>2</sub> fold change

5

0

Solyc10g086060,2

lyc01g0203 Solyc05g0

Solyc12g099980:3 Solyc10g086410.3 Solyc01g107080.3 Solyc01g102960.3 Solyc01g102960.3 Solyc03g115230.3

+ Solyc05g046310.3 120.2 Solyc11g020330.1 670.3 Solycho

10

total = 12875 variables

They also show that most of the changes were in the positive direction. Most of the genes that met the significance threshold (the horizontal line) were in the upper right quadrant of the plots, indicating that their expression levels increased in response to the treatment.

## Genotype VF36, wild-type

Comparing group V.28.15 to V.34.15 found 0 genes with changed expression levels with false discovery rate Q less than or equal to 0.05.

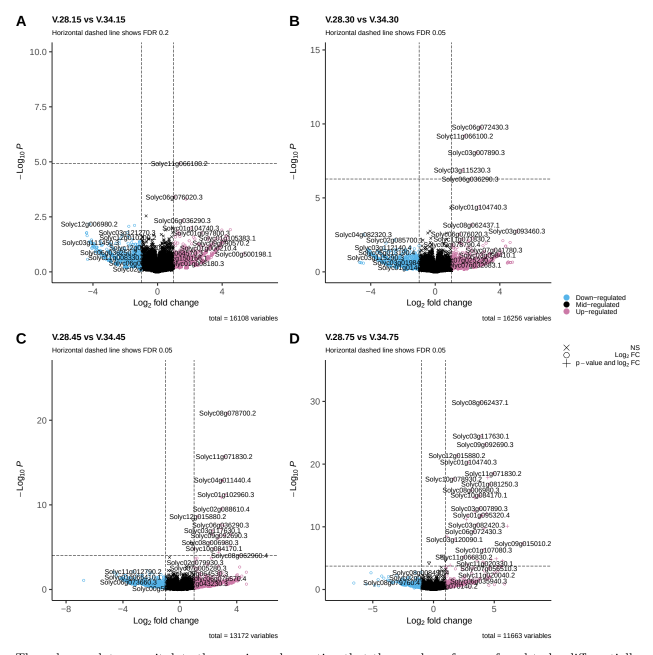
Comparing group V.28.30 to V.34.30 found 5 genes with changed expression levels with false discovery rate Q less than or equal to 0.05.

Comparing group V.28.45 to V.34.45 found 27 genes with changed expression levels with false discovery rate Q less than or equal to 0.05.

Comparing group V.28.75 to V.34.75 found 43 genes with changed expression levels with false discovery rate Q less than or equal to 0.05.

The number of DE genes called at Q less than or equal to 0.05 increased with treatment duration time.

Display volcano plots that summarize the above results:



The volcano plots recapitulate the previous observation that the number of genes found to be differentially expressed in response to the heat treatment increased with treatment duration.

Second, nearly all changes that did meet achieve the target false discovery rate of 0.05 were changes in the positive direction with the average estimated treatment expression higher than control. This can be seen by observing how most of the points above the horizontal line, when it was drawn, were also the in the right half of the plot.

Note that plot A draws at line at a different FDR, a value larger than 0.05. If no gene achieved FDR of 0.05 or higher, then our code picks a new Q indicating the p value for the gene with the smallest adjusted p value observed.

## Genotype F3H-OX3, F3H overexpression genotype

Comparing group F.28.15 to F.34.15 found 1 genes with changed expression levels with false discovery rate Q less than or equal to 0.05.

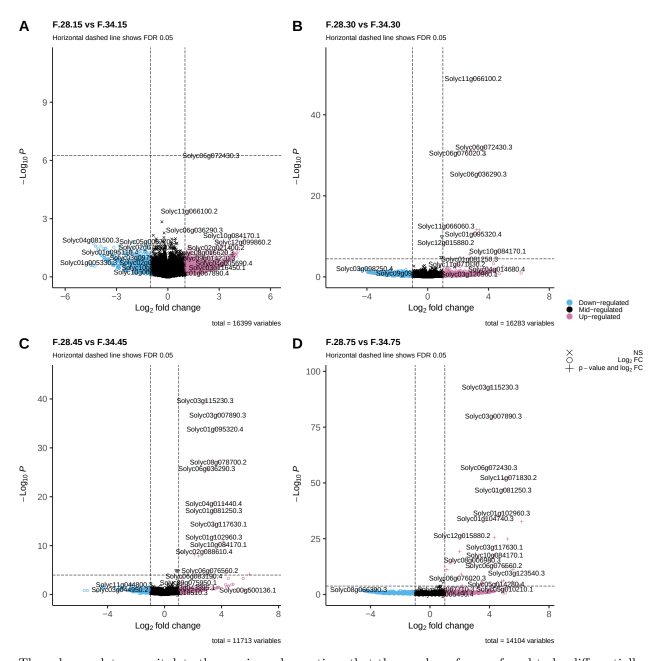
Comparing group F.28.30 to F.34.30 found 18 genes with changed expression levels with false discovery rate Q less than or equal to 0.05.

Comparing group F.28.45 to F.34.45 found 25 genes with changed expression levels with false discovery rate Q less than or equal to 0.05.

Comparing group F.28.75 to F.34.75 found 48 genes with changed expression levels with false discovery rate Q less than or equal to 0.05.

The number of DE genes called at Q less than or equal to 0.05 increased with treatment duration time.

Display volcano plots that summarize the above results:



The volcano plots recapitulate the previous observations that the number of genes found to be differentially expressed in response to the heat treatment increased with treatment duration.

Second, nearly all changes that did meet achieve the target false discovery rate of 0.05 were changes in the positive direction with the average estimated treatment expression higher than control. This can be seen by observing how most of the points above the horizontal line were also the in the right half of the plot.

# Write results to a file for analysts' convenience

All DE results were saved to a large data frame, saved to data frame all, with 170427 rows and 10 columns.

Organize results and round numeric results to 3 significant digits:

```
all = all[,c("gene","group1","group2","baseMean","padj","pvalue","log2FoldChange","lfcSE","stat","descr
for (i in 4:9) {
```

```
all[,i]=signif(all[,i],3)
}
```

Write the data file:

```
write.table(all,file=out_fname,quote=F,row.names = F,sep="\t")
```

A file was created named results/CvT-DESeq2-SL4.txt that contains all the results.

All numeric values are rounded to three significant digits.

Explanation of columns:

- gene SL4 gene measured
- group 1 control group
- group 2 treatment group
- $\bullet\,$  baseMean mean across samples
- padj false discovery rate; adjusted p-value computed using method of Benjamini and Hochberg
- log2FoldChange log2(group 2 average/group 1 average)
- lfcsE log2FoldChange standard error
- stat test statistic used to assess significance
- description gene description

### Discussion

Within each genotype, the number of genes exhibiting expression changes increased with treatment duration.

The different genotypes exhibited different numbers of differentially expressed genes, with the are genotype exhibiting the greatest number. This is consistent with the known are lower fertility mutant phenotype.

### Conclusion

- The number of genes detected as changed increased with treatment duration.
- Most fold-changes for genes that changed in respond to the treatment were positive, indicating that
  expression levels increased.
- We observed more temperature-dependent gene expression differences in the are genotype than for the other two genotypes tested.

### Session info

```
sessionInfo()
```

```
## R version 4.3.1 (2023-06-16)
## Platform: x86_64-apple-darwin20 (64-bit)
## Running under: macOS Big Sur 11.7.9
##
## Matrix products: default
## BLAS: /Library/Frameworks/R.framework/Versions/4.3-x86_64/Resources/lib/libRblas.0.dylib
## LAPACK: /Library/Frameworks/R.framework/Versions/4.3-x86_64/Resources/lib/libRlapack.dylib; LAPACK
```

```
##
## locale:
## [1] en_US.UTF-8/en_US.UTF-8/en_US.UTF-8/C/en_US.UTF-8/en_US.UTF-8
## time zone: America/New York
## tzcode source: internal
## attached base packages:
## [1] stats4
                 stats
                           graphics grDevices utils
                                                          datasets methods
## [8] base
##
## other attached packages:
  [1] EnhancedVolcano_1.18.0
                                    ggrepel_0.9.3
## [3] ggplot2_3.4.3
                                    DESeq2_1.40.2
## [5] SummarizedExperiment_1.30.2 Biobase_2.60.0
##
   [7] MatrixGenerics_1.12.3
                                    matrixStats_1.0.0
## [9] GenomicRanges_1.52.0
                                    GenomeInfoDb_1.36.3
## [11] IRanges 2.34.1
                                    S4Vectors 0.38.1
## [13] BiocGenerics_0.46.0
                                    edgeR_3.42.4
## [15] limma 3.56.2
                                    readxl 1.4.3
## [17] readr_2.1.4
                                    stringr_1.5.0
## [19] git2r_0.32.0
##
## loaded via a namespace (and not attached):
## [1] gtable 0.3.4
                                xfun 0.40
                                                         rstatix_0.7.2
## [4] lattice_0.21-8
                                tzdb 0.4.0
                                                         vctrs_0.6.3
## [7] tools_4.3.1
                                bitops_1.0-7
                                                         generics_0.1.3
## [10] parallel_4.3.1
                                tibble_3.2.1
                                                         fansi_1.0.4
## [13] pkgconfig_2.0.3
                                Matrix_1.6-1
                                                         lifecycle_1.0.3
## [16] GenomeInfoDbData_1.2.10 farver_2.1.1
                                                         compiler_4.3.1
## [19] munsell_0.5.0
                                codetools_0.2-19
                                                         carData_3.0-5
## [22] htmltools_0.5.6
                                RCurl_1.98-1.12
                                                         yaml_2.3.7
## [25] car_3.1-2
                                tidyr_1.3.0
                                                         ggpubr_0.6.0
## [28] pillar_1.9.0
                                crayon_1.5.2
                                                         BiocParallel_1.34.2
## [31] DelayedArray_0.26.7
                                abind 1.4-5
                                                         tidyselect 1.2.0
## [34] locfit_1.5-9.8
                                digest_0.6.33
                                                         stringi_1.7.12
## [37] purrr 1.0.2
                                dplyr_1.1.3
                                                         labeling 0.4.3
## [40] cowplot_1.1.1
                                fastmap_1.1.1
                                                         grid_4.3.1
## [43] colorspace_2.1-0
                                cli_3.6.1
                                                         magrittr_2.0.3
## [46] S4Arrays_1.0.6
                                utf8_1.2.3
                                                         broom_1.0.5
## [49] withr 2.5.0
                                backports_1.4.1
                                                         scales 1.2.1
## [52] rmarkdown 2.24
                                XVector_0.40.0
                                                         gridExtra_2.3
## [55] ggsignif_0.6.4
                                cellranger_1.1.0
                                                         hms_1.1.3
## [58] evaluate_0.21
                                knitr_1.44
                                                         rlang_1.1.1
## [61] Rcpp_1.0.11
                                glue_1.6.2
                                                         rstudioapi_0.15.0
## [64] R6_2.5.1
                                zlibbioc_1.46.0
```