# Pre-processing of *Strongyloides ratti* bulk RNA-seq via an alignment-free analysis pipeline

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#### 1 Introduction

The goal of this file is to pre-process the *Strongyloides ratti* RNA-seq dataset originally published by Hunt *et al* 2016.

# 2 Pre-processing Methods Overview

Raw reads are aligned to the *S. ratti* reference transcriptome (PRJEB125.WBPS14.mRNA\_transcripts, downloaded from WormBase Parasite on 17 August 2020), using Kallisto. Kallisto alignments are imported into the R environment and annotated with information imported via the Wormbase ParaSite BioMaRT. Annotation information includes: *C. elegans* homology/percent homology, UniProtKB number, Interproterms, GO terms, and general Description information. Hunt *et al* 2016 establishes two distinct subclades from the four sequenced *Strongyloides* species: *S. venezuelensis-S. papillosus* and *S. ratti-S. stercoralis*. Thus, we also include annotation information for the appropriate in-group (here, *S. stercoralis*), and a reference member of the out-group (*S. papillosus*). Annotation information is saved as an R object that is passed to a Shiny Web App for downstream browsing and on-demand analysis. Note: Raw count data could be saved as a digitial gene expression list if desired (not currently done).

Raw reads were quantified as counts per million using the EdgeR package, then filtered to remove transcripts with low counts (less than 1 count-per-million in at least 1 sample). A list of discarded genes and their expression values across life stages is saved. Non-discarded gene values are normalized using the trimmed

mean of M-values method (TMM, Robinson and Oshlack) to permit between-samples comparisons. The mean-variance relationship was modeled using a precision weights approach Law et al 2014. This dataset includes technical replicates; after voom modeling, data are condensed by replacing within-experiment technical replicates with their average, using the limma:avearrays function.

A variance-stabilized, condensed DGEList object is saved; this file is passed to a Shiny Web App for downstream browsing and on-demand analysis.

Note: samples included in this database were collected in two separate experiments, with FLM and iL3s in one, and FLF and PF in another. Due to the lack of sample overlap between the experiments, it is not possible to correct for batch effects.

### 3 Results/Analysis

#### 3.1 Kallisto read mapping

This script checks the quality of the fastq files and performs an alignment to the *Strongyloides ratti* transcriptome reference with Kallisto. To run this 'shell script' you will need to open your terminal and navigate to the directory where this script resides on your computer. For addditional instructions, see comments at head of script.

#### 3.2 Import Kallisto reads into R

Import Kallisto transcript counts into R using Tximport. Counts are generated from abundance files using the lengthScaledTPM option. This code chunk is not evaluated each time, instead it was run once and an object containing the transcripts per million data is saved. In subsequent chunks, that file is loaded, and analysis progresses. The point of this is so that folks attempting to rerun this analysis do not need to have abundance files loaded on their local machines (and we do not have to upload abundance files to github).

#### 3.3 Gene Annotation

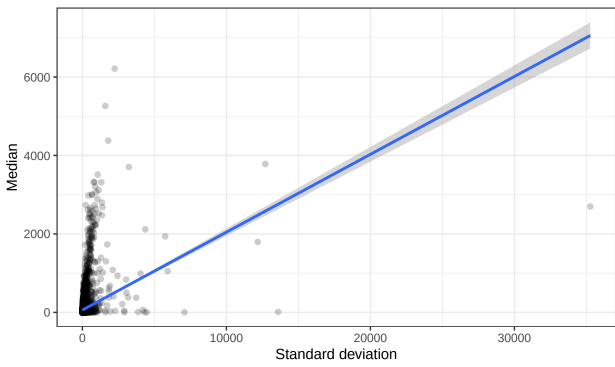
Import gene annotation information for S. ratti genes, including:

- C. elegans homology/percent homology
- S. stercoralis homology/percent homology
- S. papillosus homology/percent homology
- UniProtKB number
- Interpro terms
- GO terms
- general Description information using biomart.

#### 3.4 Generate Digital Gene Expression List

Next we generate a digital gene expression list that can be easily shared/loaded for downstream filtering/normalization. This code chunk generates a scatter plot of unfiltered and non-normalized transcripts per million data.

# Transcripts per million (TPM) unfiltered, non-normalized data



S. ratti RNA-seq Dataset

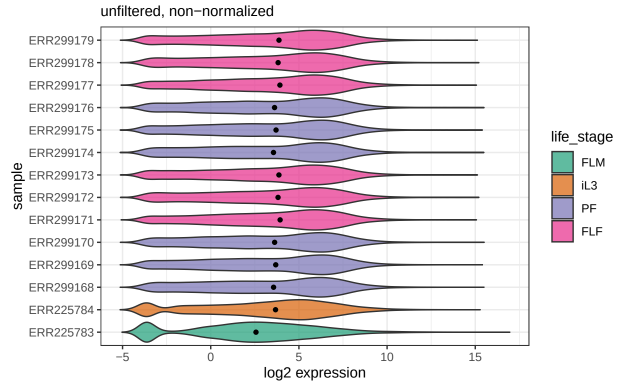
# 3.5 Data Filtering and Normalization

The goal of this chunk is to:

- 1. Filter and normalize data
- 2. Use ggplot2 to visualize the impact of filtering and normalization on the data.

## 3.5.1 Plot of unfiltered, non-normalized log2CPM data by life stage

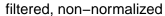
# S. ratti: Log2 Counts per Million (CPM)

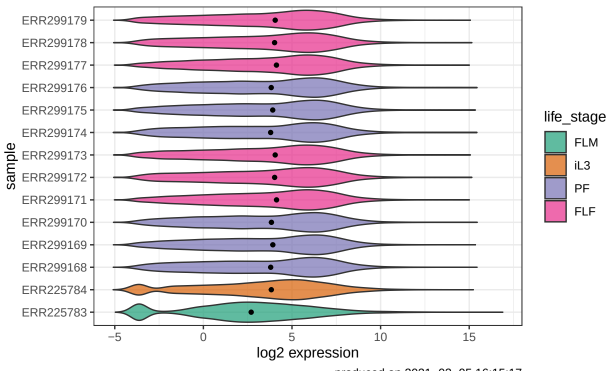


produced on 2021-02-05 16:15:15

#### 3.5.2 Plot of filtered, non-normalized log2CPM data by life stage

# S. ratti: Counts per Million (CPM)



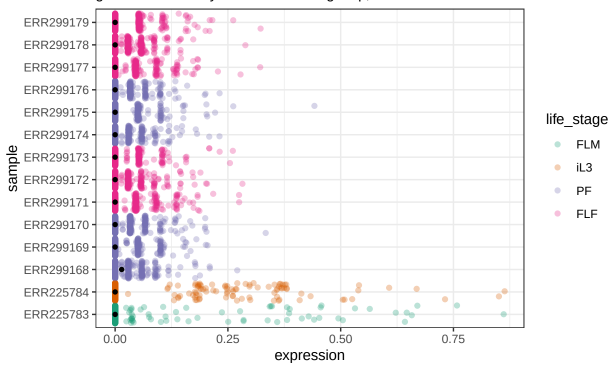


#### produced on 2021-02-05 16:15:17

#### 3.5.3 Plot of genes discarded by low-copy filtering step

The low copy number filtering step excluded a total of dim(myDGEList.discarded)[[1]] genes.

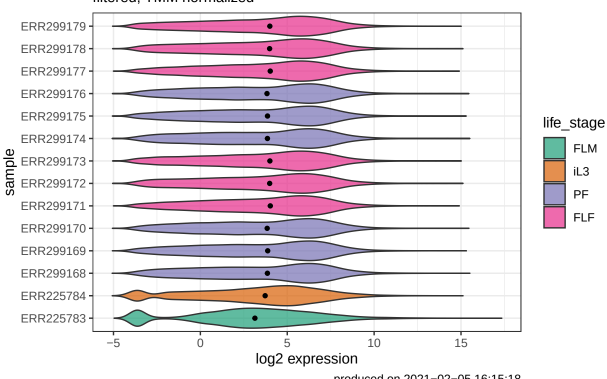
S. ratti: Counts per Million (CPM) genes excluded by low count filtering step, non-normalized



produced on 2021-02-05 16:15:17

#### 3.5.4 Plot of filtered, normalized log2CPM data by life stage

# S. ratti: Log2 Counts per Million (CPM) filtered, TMM normalized

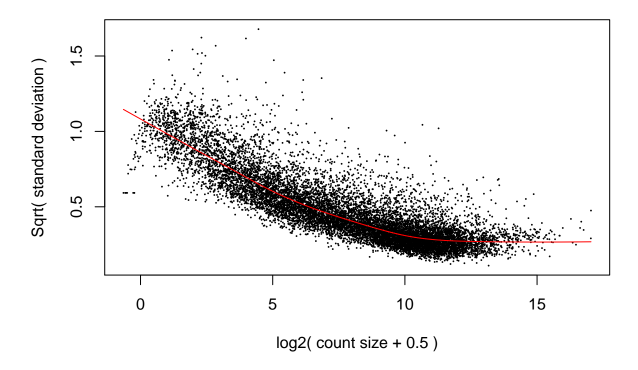


produced on 2021-02-05 16:15:18

#### 3.6 Compute Variance-Stabilized DGEList Object

This chunk uses a DGEList of filtered and normalized abundance data. It will fit data to a linear model for responsively detecting differentially expressed genes (DEGs).

#### voom: Mean-variance trend



#### 3.7 Save Data and Annotations

Finally, we save data and annotations generated in code chunks above. We can separate these saving actions into two groups:

- 1. Data saved for downstream offline analyses, including the SrRNAseq\_data\_preprocessed file which saves filtered, normalized (but not voom adjusted) log2CPM values, gene annotation information, and sample information.
- 2. Files that are required inputs to the Strongyloides RNA-seq Browser App, including:
  - i) a gene annotation R object (Sr\_geneAnnotations)
  - ii) the variance-stabilized vDGEList, saved as an R object (Sr\_vDGEList)
  - iii) a matrix of discarded genes and their raw counts (SrRNAseq\_discardedGene\_counts.csv) this data is downloadable from within the Browser App
  - iv) a matrix of variance-stabilized gene expression data, extracted from the vDGEList (SrRNAseq\_log2cpm\_filtered\_norm\_voom.csv) this data is downloadable from within the Browser App

By default, this code chunk will not be evaluated. To save the files described above, change the chunk option from eval = FALSE to eval = TRUE. These files are saved in an Outputs folder; in order to make them accessible to a local version of the Shiny browser they need to be moved to appropriate subfolders within the App folder - the www sub folder (for .csv files) or the Data subfolder (for R objects). Stable copies are already located within those folders and do not need to be replaced unless the pre-processing steps change.

### 4 Appendix I: All code for this report

```
knitr::opts_chunk$set(echo = FALSE, message = FALSE, warning = FALSE)
# This script checks the quality of the fastq files and performs an alignment
# to the Strongyloides ratti cDNA transcriptome reference with Kallisto.
# To run this 'shell script' you will need to open your terminal and navigate
# to the directory where this script resides on your computer.
# This should be the same directory where you fastq files and
# reference fasta file are found.
# Change permissions on your computer so that you can run a shell script by
# typing: 'chmod u+x readMapping_sratti.sh' (without the quotes)
# at the terminal prompt
# Then type './readMapping_sratti.sh' (without the quotes) at the prompt.
# This will begin the process of running each line of code in the shell script.
# first use fastqc to check the quality of the fastq files:
fastqc *.gz -t 14
# build index from the reference fasta file
kallisto index -i strongyloides_ratti.PRJEB125.WBPS14.mRNA_transcripts.index
strongyloides_ratti.PRJEB125.WBPS14.mRNA_transcripts.fa
# map reads to the indexed reference host transcriptome
# Parasitic Females: Biological Replicates A-C, Technical replicate set 1
kallisto quant -i strongyloides_ratti.PRJEB125.WBPS14.mRNA_transcripts.index -o
ERR299168 -t 14 ERR299168 1.fastq.gz ERR299168 2.fastq.gz&> ERR299168.log
kallisto quant -i strongyloides_ratti.PRJEB125.WBPS14.mRNA_transcripts.index -o
ERR299169 -t 14 ERR299169 1.fastq.gz ERR299169 2.fastq.gz&> ERR299169.log
kallisto quant -i strongyloides_ratti.PRJEB125.WBPS14.mRNA_transcripts.index -o
ERR299170 -t 14 ERR299170_1.fastq.gz ERR299170_2.fastq.gz&> ERR299170.log
# Free-living Females: Biological Replicates A-C, Technical replicate set 1
kallisto quant -i strongyloides_ratti.PRJEB125.WBPS14.mRNA_transcripts.index -o
ERR299171 -t 14 ERR299171_1.fastq.gz ERR299171_2.fastq.gz&> ERR299171.log
kallisto quant -i strongyloides_ratti.PRJEB125.WBPS14.mRNA_transcripts.index -o
ERR299172 -t 14 ERR299172_1.fastq.gz ERR299172_2.fastq.gz&> ERR299172.log
kallisto quant -i strongyloides_ratti.PRJEB125.WBPS14.mRNA_transcripts.index -o
ERR299173 -t 14 ERR299173_1.fastq.gz ERR299173_2.fastq.gz&> ERR299173.log
# Parasitic Females: Biological Replicates A-C, Technical replicate set 2
kallisto quant -i strongyloides_ratti.PRJEB125.WBPS14.mRNA_transcripts.index -o
ERR299174 -t 14 ERR299174_1.fastq.gz ERR299174_2.fastq.gz&> ERR299174.log
kallisto quant -i strongyloides ratti.PRJEB125.WBPS14.mRNA transcripts.index -o
ERR299175 -t 14 ERR299175_1.fastq.gz ERR299175_2.fastq.gz&> ERR299175.log
kallisto quant -i strongyloides ratti.PRJEB125.WBPS14.mRNA transcripts.index -o
ERR299176 -t 14 ERR299176_1.fastq.gz ERR299176_2.fastq.gz&> ERR299176.log
# Free-living Females: Biological Replicates A-C, Technical replicate set 2
kallisto quant -i strongyloides_ratti.PRJEB125.WBPS14.mRNA_transcripts.index -o
ERR299177 -t 14 ERR299177_1.fastq.gz ERR299177_2.fastq.gz&> ERR299177.log
kallisto quant -i strongyloides_ratti.PRJEB125.WBPS14.mRNA_transcripts.index -o
```

```
kallisto quant -i strongyloides_ratti.PRJEB125.WBPS14.mRNA_transcripts.index -o
ERR299179 -t 14 ERR299179 1.fastq.gz ERR299179 2.fastq.gz&> ERR299179.log
# Free-living Males
kallisto quant -i strongyloides_ratti.PRJEB125.WBPS14.mRNA_transcripts.index -o
# iL3s
kallisto quant -i strongyloides_ratti.PRJEB125.WBPS14.mRNA_transcripts.index -o
ERR225784 -t 14 ERR225784_1.fastq.gz ERR225784_2.fastq.gz&> ERR225784.log
# summarize fastqc and kallisto mapping results using MultiQC
multiqc -d .
echo "Finished"
# load packages ----
suppressPackageStartupMessages({
 library(tidyverse)
 library(tximport)
 library(ensembldb)
 library(biomaRt)
 library(magrittr)
})
# read in the study design ----
targets <- read tsv(".../Data/S ratti/Study Design/SrattiRNAseq study design.txt",
                  na = c("", "NA", "na"))
# create file paths to the abundance files generated by Kallisto
# using the 'file.path' function
path <- file.path("../Data/S_ratti/Reads",targets$sample, "abundance.tsv")</pre>
# get annotations using organism-specific package ----
Tx.Sr <- getBM(attributes=c('wbps_transcript_id',</pre>
                          'wbps_gene_id'),
              # grab the ensembl annotations for Wormbase Parasite genes
              mart = useMart(biomart="parasite_mart",
                            dataset = "wbps_gene",
                            host="https://parasite.wormbase.org",
                            port = 443),
              filters = c('species_id_1010'),
              value = list('strattprjeb125')) %>%
 as_tibble() %>%
 #we need to rename the columns retreived from biomart
 dplyr::rename(target_id = wbps_transcript_id,
               WB_geneID = wbps_gene_id) %>%
 dplyr::mutate(gene_name = str_remove_all(target_id, "\\.[0-9]$")) %>%
 dplyr::mutate(gene_name = str_remove_all(gene_name, "[a-c]$")) %>%
 dplyr::select(!WB_geneID)
# import Kallisto transcript counts into R using Tximport ----
# copy the abundance files to the working directory and rename
# so that each sample has a unique name
```

```
Txi_gene <- tximport(path,</pre>
                     type = "kallisto",
                     tx2gene = Tx.Sr[,1:2],
                     txOut = FALSE,
                     countsFromAbundance = "lengthScaledTPM",
                     ignoreTxVersion = FALSE)
# Save the raw transcript counts ----
save(Txi_gene,
     file = file.path("../Data/S_ratti",
                      "SrRNAseq TPM"))
# Introduction to this chunk -----
# This chunk imports gene annotation information for S. ratti genes, including:
# C. elegans homologs/percent homology,
# S. stercoralis homologs/percent homology,
# S. papillosus homologs/percent homology,
# UniProtKB number, Interpro terms, GO terms,
# and general Description information using biomart.
# It will generate a table that can be saved.
# Load packages -----
suppressPackageStartupMessages({
 library(tidyverse)
 library(tximport)
 library(ensembldb)
 library(biomaRt)
 library(magrittr)
})
# Load data & study design ----
load(file = file.path("../Data/S_ratti",
                      "SrRNAseq_TPM"))
targets <- read_tsv("../Data/S_ratti/Study_Design/SrattiRNAseq_study_design.txt",</pre>
                    na = c("", "NA", "na"))
# Get In-subclade group homologs for S. ratti
# genes from BioMart and filter ----
Annt.temp.1 <- getBM(attributes=c('wbps_transcript_id',</pre>
                                   'ststerprjeb528_gene',
                                   'ststerprjeb528_homolog_perc_id'),
                     # grab the ensembl annotations for Wormbase Parasite genes
                     mart = useMart(biomart="parasite_mart",
                                    dataset = "wbps_gene",
                                    host="https://parasite.wormbase.org",
                                    port = 443),
                     filters = c('species_id_1010'),
                     value = list('strattprjeb125')) %>%
  as_tibble() %>%
  #rename columns
  dplyr::rename(geneID = wbps_transcript_id,
                In.subclade_geneID = ststerprjeb528_gene,
```

```
In.subclade_percent_homology= ststerprjeb528_homolog_perc_id
 ) %>%
 dplyr::group by(geneID)
# Get Out-subclade group homologs for S. ratti
# genes from BioMart and filter ----
Annt.temp.2 <- getBM(attributes=c('wbps_transcript_id',</pre>
                                  'stpapiprjeb525_gene',
                                  'stpapiprjeb525_homolog_perc_id'
                     # grab the ensembl annotations for Wormbase Parasite genes
                     mart = useMart(biomart="parasite_mart",
                                    dataset = "wbps_gene",
                                    host="https://parasite.wormbase.org",
                                    port = 443),
                     filters = c('species_id_1010'),
                     value = list('strattprjeb125')) %>%
 as tibble() %>%
 #rename columns
 dplyr::rename(geneID = wbps_transcript_id,
                Out.subclade geneID = stpapiprjeb525 gene,
                Out.subclade_percent_homology= stpapiprjeb525_homolog_perc_id
 ) %>%
 dplyr::group_by(geneID)
# Get C. elegans homologs and gene information for S. ratti
# genes from BioMart and filter ----
Annt.temp.3 <- getBM(attributes=c('wbps_transcript_id',</pre>
                                   'wbps_gene_id',
                                   'caelegprjna13758_gene_name',
                                   'caelegprjna13758_homolog_perc_id',
                                   'description',
                                   'interpro_short_description',
                                   'go_name_1006',
                                   'uniprot_sptrembl'),
                     # grab the ensembl annotations for Wormbase Parasite genes
                     mart = useMart(biomart="parasite_mart",
                                    dataset = "wbps_gene",
                                    host="https://parasite.wormbase.org",
                                    port = 443),
                     filters = c('species_id_1010',
                                  'wbps_transcript_id'),
                     value = list('strattprjeb125',
                                  rownames(Txi_gene$counts))) %>%
 as_tibble() %>%
  #rename columns
 dplyr::rename(geneID = wbps_transcript_id,
                WBgeneID = wbps_gene_id,
                Ce_geneID = caelegprjna13758_gene_name,
                Ce_percent_homology = caelegprjna13758_homolog_perc_id,
                Description = description,
                GO_term = go_name_1006,
                UniProtKB = uniprot_sptrembl
```

```
) %>%
  dplyr::group_by(geneID)
Annt.import <- full_join(Annt.temp.1, Annt.temp.2, by = "geneID") %>%
  full_join(Annt.temp.3, by = "geneID")
Annt.import$geneID <- str remove all(Annt.import$geneID, "\\.[0-9]$")
Annt.import$geneID <- str_remove_all(Annt.import$geneID, "[a-z]$")
# Replace empty string values (mostly in Ce_geneID column) with NAs
Annt.import[Annt.import == ""]<-NA
# Remove any duplications in the possible homolog matches.
# Select based on highest % homology.
# #Give fake value here to make sure genes
#without a homologs aren't filtered out
Annt.import$Ce_percent_homology[
  is.na(Annt.import$Ce_percent_homology)] <- 1000</pre>
Annt.import$In.subclade_percent_homology[
  is.na(Annt.import$In.subclade_percent_homology)] <- 1000</pre>
Annt.import$Out.subclade_percent_homology[
  is.na(Annt.import$Out.subclade_percent_homology)] <- 1000</pre>
Annt.logs <-Annt.import %>%
  dplyr::select(!c(interpro_short_description:GO_term))%>%
  group_by(geneID) %>%
  slice_max(n = 1, order_by = Ce_percent_homology,
            with_ties = FALSE) %>%
  slice_max(n = 1, order_by = In.subclade_percent_homology,
            with_ties = FALSE) %>%
  slice_max(n = 1, order_by = Out.subclade_percent_homology,
            with_ties = FALSE) %>%
  group_by(geneID, Ce_geneID)
# Remove source code to shorten the description
Annt.logs$Description<- Annt.logs$Description %>%
  str_replace_all(string = .,
                  pattern = " \\[Source:.*\\]",
                  replacement = "") %>%
  cbind()
Annt.logs Ce percent homology [
  Annt.logs$Ce percent homology == 1000] <- NA
Annt.logs$In.subclade_percent_homology[
  Annt.logs$In.subclade_percent_homology == 1000] <- NA
Annt.logs$Out.subclade_percent_homology[
  Annt.logs$Out.subclade_percent_homology == 1000] <- NA
# Clean up interprotKB terms, removing duplications and collapsing to one line
Annt.interpro<-Annt.import %>%
  dplyr::select(geneID, Ce_geneID, interpro_short_description) %>%
  group_by(geneID, Ce_geneID) %>%
```

```
dplyr::distinct(interpro_short_description, .keep_all = TRUE) %>%
  dplyr::summarise(InterPro = paste(interpro_short_description,
                                    collapse = ', '))
# Clean up GO terms, removing duplications and collapsing to one line
Annt.goterms<-Annt.import %>%
  dplyr::select(geneID, Ce_geneID, GO_term) %>%
  group_by(geneID, Ce_geneID) %>%
  dplyr::distinct(GO term, .keep all = TRUE) %>%
  dplyr::summarise(GO_term = paste(GO_term, collapse = ', '))
annotations <-dplyr::left_join(Annt.logs, Annt.interpro) %>%
  dplyr::left_join(.,Annt.goterms) %>%
  ungroup() %>%
  dplyr::relocate(WBgeneID,In.subclade_geneID,
                  In.subclade_percent_homology,
                  Out.subclade_geneID,
                  Out.subclade_percent_homology,
                  .after = geneID) %>%
  column_to_rownames(var = "geneID")
# List of S. ratti genes is longer than the number of genes
# in the RNA-seq dataset. So subset the annotations by the geneIDs in Txi_gene
annotations<-annotations[rownames(annotations) %in% rownames(Txi_gene$counts),]
# Goals of this chunk:
# Generate and save a digitial gene expression list
# that could be easily shared/loaded for downstream filtering/normalization
# Load packages -----
suppressPackageStartupMessages({
  library(tidyverse)
 library(edgeR)
 library(matrixStats)
 library(cowplot)
  library(ggthemes)
 library(RColorBrewer)
 library(gprofiler2)
})
# Generate and plot summary stats for the data ----
myTPM.stats <- transform(Txi_gene$abundance,
                         SD=rowSds(Txi gene$abundance),
                         AVG=rowMeans(Txi_gene$abundance),
                         MED=rowMedians(Txi_gene$abundance))
# produce a scatter plot of the transformed data
p1<-ggplot(myTPM.stats) +</pre>
  aes(x = SD, y = MED) +
  geom_point(shape=16, size=2, alpha = 0.2) +
  geom_smooth(method=lm) +
  #geom_hex(show.legend = FALSE) +
  labs(y="Median", x = "Standard deviation",
```

```
title="Transcripts per million (TPM)",
       subtitle="unfiltered, non-normalized data",
       caption="S. ratti RNA-seq Dataset") +
  theme bw()
р1
# make a Digital Gene Expression list using the raw counts and plot ----
myDGEList <- DGEList(Txi gene$counts,</pre>
                     samples = data.frame(samples = targets$source,
                                           source = targets$sample),
                     group = targets$group,
                     genes = annotations)
# Goals of this chunk:
# 1 - Filter and normalize data
#2 - use ggplot2 to visualize the impact of filtering and
# normalization on the data.
# Notes:
# recall that abundance data are TPM, while the counts are
# read counts mapping to each gene or transcript
# Load packages -----
suppressPackageStartupMessages({
  library(tidyverse)
 library(edgeR)
 library(matrixStats)
 library(cowplot)
 library(ggthemes)
  library(RColorBrewer)
  library(gprofiler2)
})
# calculate and plot log2 counts per million ----
# Generate life stage IDs
ids <- rep(cbind(targets$group),</pre>
           times = nrow(myDGEList$counts)) %>%
  as_factor()
# use the 'cpm' function from EdgeR to get log2 counts per million
# then coerce into a tibble
log2.cpm.df.pivot <-cpm(myDGEList, log=TRUE) %>%
  as_tibble(rownames = "geneID") %>%
  setNames(nm = c("geneID", targets$sample)) %>%
  pivot_longer(cols = -geneID,
               names_to = "samples",
               values_to = "expression") %>%
  add_column(life_stage = ids)
# plot the pivoted data
p2 <- ggplot(log2.cpm.df.pivot) +</pre>
  aes(x=samples, y=expression, fill=life_stage) +
```

```
geom_violin(trim = FALSE, show.legend = T, alpha= 0.7) +
  stat_summary(fun = "median",
               geom = "point",
               shape = 20,
               size = 2,
               color = "black",
               show.legend = FALSE) +
  labs(y="log2 expression", x = "sample",
       title="S. ratti: Log2 Counts per Million (CPM)",
       subtitle="unfiltered, non-normalized",
       caption=paste0("produced on ", Sys.time())) +
  theme bw() +
  scale_fill_brewer(palette = "Dark2") +
  coord flip()
p2
# Filter the data ----
# filter genes/transcripts with low counts
# how many genes had more than 1 CPM (TRUE) in at least n samples
# Note: The cutoff "n" is adjusted for the number of
# samples in the smallest group of comparison.
keepers <- cpm(myDGEList) %>%
  rowSums(.>1)>=1
myDGEList.filtered <- myDGEList[keepers,]</pre>
ids.filtered <- rep(cbind(targets$group),</pre>
                    times = nrow(myDGEList.filtered)) %>%
  as_factor()
log2.cpm.filtered.df.pivot <- cpm(myDGEList.filtered, log=TRUE) %>%
  as_tibble(rownames = "geneID") %>%
  setNames(nm = c("geneID", targets$sample)) %>%
  pivot_longer(cols = -geneID,
               names_to = "samples",
               values_to = "expression") %>%
  add_column(life_stage = ids.filtered)
p3 <- ggplot(log2.cpm.filtered.df.pivot) +
  aes(x=samples, y=expression, fill=life_stage) +
  geom_violin(trim = FALSE, show.legend = T, alpha= 0.7) +
  stat_summary(fun = "median",
               geom = "point",
               shape = 20,
               size = 2,
               color = "black",
               show.legend = FALSE) +
  labs(y="log2 expression", x = "sample",
       title="S. ratti: Counts per Million (CPM)",
       subtitle="filtered, non-normalized",
       caption=paste0("produced on ", Sys.time())) +
  theme_bw() +
  scale_fill_brewer(palette = "Dark2") +
```

```
coord_flip()
рЗ
# Look at the genes excluded by the filtering step ----
# just to check that there aren't any with high expression that are in few samples
# Discarded genes
myDGEList.discarded <- myDGEList[!keepers,]</pre>
ids.discarded <- rep(cbind(targets$group),</pre>
                    times = nrow(myDGEList.discarded)) %>%
    as factor()
log2.cpm.discarded.df.pivot <- cpm(myDGEList.discarded, log=F) %>%
    as_tibble(rownames = "geneID") %>%
    setNames(nm = c("geneID", targets$sample)) %>%
    pivot_longer(cols = -geneID,
                 names_to = "samples",
                 values_to = "expression") %>%
    add_column(life_stage = ids.discarded)
p.discarded <- ggplot(log2.cpm.discarded.df.pivot) +</pre>
    aes(x=samples, y=expression, color=life_stage) +
    #geom_violin(trim = FALSE, show.legend = T, alpha= 0.7) +
    geom_jitter(alpha = 0.3, show.legend = T)+
    stat summary(fun = "median",
                 geom = "point",
                 shape = 20,
                 size = 2,
                 color = "black",
                 show.legend = FALSE) +
    labs(y="expression", x = "sample",
         title="S. ratti: Counts per Million (CPM)",
         subtitle="genes excluded by low count filtering step, non-normalized",
         caption=paste0("produced on ", Sys.time())) +
    theme_bw() +
    scale_color_brewer(palette = "Dark2") +
    coord flip()
p.discarded
# # Carry out GO enrichment of discarded gene set using gProfiler2 ----
# discarded.geneID <- unique(log2.cpm.discarded.df.pivot$geneID)</pre>
# gost.res <- gost(list(Discarded_genes = discarded.geneID),</pre>
                   organism = "strattprjeb125", correction_method = "fdr")
# gostplot(gost.res, interactive = T, capped = T)
# Generate a matrix of discarded genes and their raw counts ----
discarded.gene.df <- log2.cpm.discarded.df.pivot %>%
  pivot_wider(names_from = c(life_stage, samples),
              names_sep = "-",
              values_from = expression,
              id_cols = geneID)
```

```
# Normalize the data using a between samples normalization ----
# Source for TMM sample normalization here:
# https://genomebiology.biomedcentral.com/articles/10.1186/gb-2010-11-3-r25
myDGEList.filtered.norm <- calcNormFactors(myDGEList.filtered, method = "TMM")
log2.cpm.filtered.norm <- cpm(myDGEList.filtered.norm, log=TRUE)</pre>
log2.cpm.filtered.norm.df<- cpm(myDGEList.filtered.norm, log=TRUE) %>%
  as tibble(rownames = "geneID") %>%
  setNames(nm = c("geneID", targets$sample))
log2.cpm.filtered.norm.df.pivot<-log2.cpm.filtered.norm.df %>%
  pivot_longer(cols = -geneID,
               names_to = "samples",
               values_to = "expression") %>%
  add_column(life_stage = ids.filtered)
p4 <- ggplot(log2.cpm.filtered.norm.df.pivot) +
  aes(x=samples, y=expression, fill=life_stage) +
  geom_violin(trim = FALSE, show.legend = T, alpha = 0.7) +
  stat_summary(fun = "median",
               geom = "point",
               shape = 20,
               size = 2,
               color = "black",
               show.legend = FALSE) +
  labs(y="log2 expression", x = "sample",
       title="S. ratti: Log2 Counts per Million (CPM)",
       subtitle="filtered, TMM normalized",
       caption=paste0("produced on ", Sys.time())) +
  theme_bw() +
  scale_fill_brewer(palette = "Dark2") +
  coord_flip()
p4
# Introduction to this chunk ----
# This chunk uses a DGEList of filtered and normalized abundance data
# It will fit data to a linear model for responsively detecting
# differentially expressed genes (DEGs)
# Load packages ----
suppressPackageStartupMessages({
 library(tidyverse)
 library(limma) # differential gene expression using linear modeling
 library(edgeR)
})
# Set up the design matrix ----
group <- factor(targets$group)</pre>
biolrep <- factor(targets$source)</pre>
#block <- factor(targets$block)</pre>
# ideally, I would use a blocking design to handle possibility of batch effects
# between "PRJEB1376" and "PRJEB3187", comparisons across group.
```

```
# However, because there aren't any overlaping samples, between the two groups,
# I don't think I can use the (~block + group) setup for the model matrix.
# At this point, the batch is likely a confounding variable.
design <- model.matrix(~0 + group)</pre>
colnames(design) <- levels(group)</pre>
# Model mean-variance trend and fit linear model to data ----
colnames(myDGEList.filtered.norm$counts) <- targets$group</pre>
v.DEGList.filtered.norm.withtechreplicates <- voom(counts = myDGEList.filtered.norm,
                                 design = design,
                                 plot = T)
colnames(v.DEGList.filtered.norm.withtechreplicates$E) <- targets$group</pre>
# Condense data by replacing within-experiment technical
# replicates with their average
v.DEGList.filtered.norm <- avearrays(v.DEGList.filtered.norm.withtechreplicates,
                                      biolrep)
colnames(v.DEGList.filtered.norm$E) <- paste(v.DEGList.filtered.norm$targets$group,</pre>
                                              levels(biolrep),sep = '-')
# Check for presence of output folder, generate if it doesn't exist
output.path <- "../Outputs"</pre>
if (!dir.exists(output.path)){
 dir.create(output.path)
# Save full gene annotations ----
# This object is required for the Shiny Browser
save(annotations,
     file = file.path(output.path,
                      "Sr geneAnnotations"))
# Save a matrix of discarded genes and their raw counts ----
discarded.gene.df %>%
write.csv(file = file.path(output.path,
                      "SrRNAseq discardedGene counts.csv"))
# Save matrix of genes and their filtered, normalized, voom-transformed counts ----
# This is the count data that underlies the differential expression analyses in the Shiny app.
# Saving it here so that users of the app can access the input information.
write.csv(v.DEGList.filtered.norm$E,
          file = file.path(output.path,
                            "SrRNAseq log2cpm filtered norm voom.csv"))
# Save v.DEGList ----
# This file will can be imported into Shiny App
save(v.DEGList.filtered.norm,
     file = file.path(output.path,
                      "Sr_vDGEList"))
```

# 5 Appendix II: Session Info

```
## R version 3.6.3 (2020-02-29)
## Platform: x86_64-apple-darwin15.6.0 (64-bit)
## Running under: macOS Catalina 10.15.5
##
## Matrix products: default
         /Library/Frameworks/R.framework/Versions/3.6/Resources/lib/libRblas.0.dylib
## BLAS:
## LAPACK: /Library/Frameworks/R.framework/Versions/3.6/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] stats4
                parallel stats
                                     graphics grDevices utils
                                                                   datasets
## [8] methods
                 base
## other attached packages:
## [1] gprofiler2_0.1.9
                                RColorBrewer_1.1-2
                                                        ggthemes_4.2.0
## [4] cowplot_1.0.0
                                matrixStats_0.56.0
                                                        edgeR_3.28.1
## [7] limma_3.42.2
                                magrittr_1.5
                                                        biomaRt_2.42.1
## [10] ensembldb 2.10.2
                                AnnotationFilter_1.10.0 GenomicFeatures_1.38.2
## [13] AnnotationDbi_1.48.0
                                Biobase_2.46.0
                                                        GenomicRanges_1.38.0
## [16] GenomeInfoDb_1.22.1
                                IRanges_2.20.2
                                                        S4Vectors_0.24.4
## [19] BiocGenerics_0.32.0
                                tximport_1.14.2
                                                        forcats_0.5.0
## [22] stringr_1.4.0
                                                        purrr_0.3.4
                                dplyr_1.0.1
                                                        tibble_3.0.3
## [25] readr 1.3.1
                                tidyr 1.1.1
## [28] ggplot2_3.3.2
                                tidyverse_1.3.0
##
## loaded via a namespace (and not attached):
## [1] colorspace_1.4-1
                                    ellipsis_0.3.1
## [3] XVector_0.26.0
                                    fs_1.4.2
## [5] rstudioapi_0.11
                                    farver_2.0.3
## [7] bit64_0.9-7
                                    fansi_0.4.1
## [9] lubridate_1.7.9
                                    xm12_1.3.2
## [11] splines_3.6.3
                                    knitr_1.29
```

```
## [13] jsonlite_1.7.0
                                    Rsamtools_2.2.3
## [15] broom_0.5.6
                                    dbplyr_1.4.4
## [17] compiler 3.6.3
                                    httr 1.4.2
## [19] backports_1.1.8
                                    assertthat_0.2.1
## [21] Matrix 1.2-18
                                    lazyeval_0.2.2
## [23] cli 2.0.2
                                    htmltools 0.5.0
## [25] prettyunits 1.1.1
                                    tools 3.6.3
## [27] gtable_0.3.0
                                    glue_1.4.1
## [29] GenomeInfoDbData_1.2.2
                                    rappdirs 0.3.1
## [31] Rcpp_1.0.5
                                     cellranger_1.1.0
## [33] vctrs_0.3.2
                                    Biostrings_2.54.0
## [35] nlme_3.1-148
                                    rtracklayer_1.46.0
## [37] xfun_0.15
                                    rvest_0.3.5
## [39] lifecycle_0.2.0
                                    XML_3.99-0.3
## [41] zlibbioc_1.32.0
                                     scales_1.1.1
## [43] hms_0.5.3
                                    ProtGenerics_1.18.0
## [45] SummarizedExperiment_1.16.1 yaml_2.2.1
## [47] curl 4.3
                                    memoise 1.1.0
## [49] stringi_1.4.6
                                    RSQLite_2.2.0
## [51] BiocParallel 1.20.1
                                    rlang 0.4.7
## [53] pkgconfig_2.0.3
                                    bitops_1.0-6
## [55] evaluate 0.14
                                    lattice 0.20-41
## [57] labeling_0.3
                                    GenomicAlignments_1.22.1
## [59] htmlwidgets 1.5.1.9001
                                    bit 1.1-15.2
## [61] tidyselect_1.1.0
                                    R6 2.4.1
## [63] generics 0.0.2
                                    DelayedArray_0.12.3
## [65] DBI_1.1.0
                                    mgcv_1.8-31
## [67] pillar_1.4.6
                                    haven_2.3.1
## [69] withr_2.2.0
                                    RCurl_1.98-1.2
## [71] modelr_0.1.8
                                     crayon_1.3.4
## [73] BiocFileCache_1.10.2
                                    plotly_4.9.2.9000
## [75] rmarkdown_2.3
                                    progress_1.2.2
## [77] locfit_1.5-9.4
                                    grid_3.6.3
## [79] readxl_1.3.1
                                    data.table_1.12.8
## [81] blob 1.2.1
                                    reprex 0.3.0
## [83] digest_0.6.25
                                    openssl_1.4.2
## [85] munsell 0.5.0
                                    viridisLite_0.3.0
## [87] askpass_1.1
```