### Project 1: Function to Summarize and Plot Gene Expression

2024-08-07

### Import and summarize data

### Clean up metadata

### Transpose Genes dataframe

```
#transposing the genes x samples matrix to samples x genes
rownames(genes) <- genes$X #assigning row names
genes_t <- as.data.frame(t(dplyr::select(genes, -c('X')))) #transpose as a dataframe
genes_t$participant_id <- rownames(genes_t) #assigning participant id column for later joining
#which(genes_t$participant_id == "COVID_06_.y_male_NonICU") #finding column with mismatch to metadata
genes_t$participant_id[6] <- "COVID_06_:y_male_NonICU" #reassinging value to match metadata</pre>
```

### Combining metadata and genes dataframe

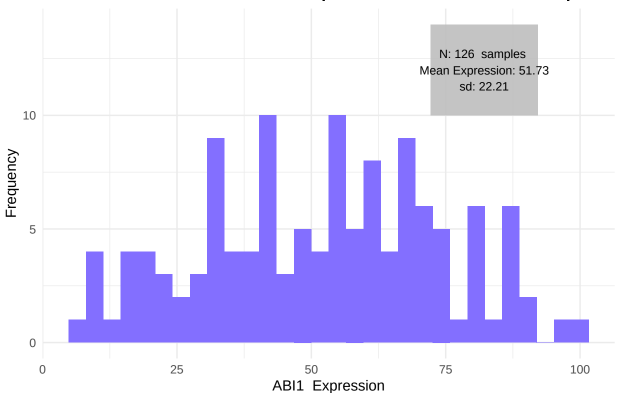
```
#combining the samples x genes matrix to the metadata
all_data <- dplyr::inner_join(genes_t, metadata_clean, by = 'participant_id')</pre>
```

### Write function for plotting

```
plots <- function(df, genes_list, cont_cov, cat_cov) {</pre>
  df_filtered <- df %>% dplyr::select(participant_id, all_of(genes_list), all_of(cont_cov), all_of(cat_
  df_filtered_long <- df_filtered %>% tidyr::pivot_longer(cols = genes_list, names_to = 'Gene', values_
    dplyr::arrange(Gene, .data[[cont_cov]])
  for (gene in genes_list){
    one_gene <- df_filtered_long %>% dplyr::filter(Gene == gene)
    mean_expression <- round(mean(one_gene$Expression), 2)</pre>
    sd_expression <- round(sd(one_gene$Expression), 2)</pre>
    x_min_box <- quantile(one_gene$Expression, 3/4)</pre>
    n <- length(one_gene$Expression)</pre>
    hist_title <- paste('Distribution of ', gene, ' expression across samples')
    hist_x <- paste(gene, ' Expression')</pre>
    hist <- one_gene %>% ggplot(aes(Expression)) +
      geom_histogram(bins = 30, fill = 'slateblue1') +
      theme minimal() +
      theme(plot.title = element_text(hjust = .4, size = 20)) +
      labs(title = hist title,
      x = hist_x,
      y = 'Frequency',
      size = 20) +
      scale_color_manual(values = c('slateblue1')) +
      annotate('rect',
           xmin = x_min_box + 5,
           xmax = x_min_box + 25,
           ymin = 10,
           ymax = 14, fill = 'grey', alpha = .9)+
  annotate(geom = 'text', x = x_min_box + 15, y = 12, label = paste('N:', n,' samples',
                                                          '\n Mean Expression:', mean_expression,
                                                          '\nsd:', sd expression), size = 3)
    print(hist)
    scatter_filtered <- one_gene %% filter(one_gene[[cont_cov]] != ' unknown') #filter out unknowns
    scatter_filtered[cont_cov] <- as.numeric(scatter_filtered[[cont_cov]])</pre>
    if (cont_cov == 'crp.mg.l.') {
      scatter_filtered$crp.level <- cut(scatter_filtered$crp.mg.l., breaks = c(0, 3, 10, 100, 500000),</pre>
                                         labels = c('normal', 'moderate', 'high', 'severe'))
    colorPalette_scatter <- c('azure4', 'skyblue2', 'slateblue2', 'midnightblue')</pre>
    title_scatter <- paste(toupper(cont_cov), ' vs ', gene, ' Expression')</pre>
    x_scatter <- paste(toupper(cont_cov), '(mg/L)')</pre>
    y_scatter <- paste(gene, ' Expression')</pre>
    scatter <- scatter_filtered %>% ggplot(aes(x = .data[[cont_cov]], y = Expression, color = crp.level
      geom_point(na.rm = TRUE) +
      theme(
        axis.text.x = element_text(angle=90),
```

```
plot.title = element_text(hjust = .4, size = 20),
        legend.position = c(.85,.75)) +
      labs(title = title_scatter, x = x_scatter, y = y_scatter) +
      scale_color_manual(labels = c('Normal (<3mg/L)', 'Moderate (3-10mg/L)', 'High (10 -100mg/L)', 'Se</pre>
                           colorPalette_scatter)
    print(scatter)
    title_box <- paste(gene, ' Expression across ', cat_cov[1], ' and ', cat_cov[2])</pre>
    x_lab_box <- cat_cov[1]</pre>
    y_lab_box <- paste(gene, ' Expression')</pre>
    n_covid_pos <- length(dplyr::filter(</pre>
      one_gene, disease_status == 'disease state: COVID-19') $Expression)
    n_covid_neg <- length(dplyr::filter(</pre>
      one_gene, disease_status != 'disease state: COVID-19')$Expression)
    y_annot <- max(one_gene$Expression)</pre>
    box <- one_gene %>% ggplot(aes_string(x = cat_cov[1], y = 'Expression', fill = cat_cov[2])) +
      geom_boxplot() +
      theme_minimal() +
      theme(legend.position = 'bottom') +
      labs(title = title_box,
           x = x_{lab_box}
           y = y_{lab_box} +
      scale_fill_manual(labels = c('no ventilator', 'ventilator'),
                        values = c('ghostwhite', 'mediumvioletred')) +
      scale_x_discrete(labels = c('COVID', 'NON-COVID')) +
      annotate(geom = 'text', x = 1, y = y_annot + 5,
               label = paste('Covid positive: N = ', n_covid_pos),
               color = 'black')+
      annotate(geom = 'text', x = 2, y = y_annot + 5,
               label = paste('Covid negative: N = ', n_covid_neg),
               color = 'black')
    print(box)
  }
  }
plots(all_data, c('ABI1', 'AAMP', 'ABHD17A'), c('crp.mg.l.'), c('disease_status', 'mechanical_ventilati
## Warning: Using an external vector in selections was deprecated in tidyselect 1.1.0.
## i Please use 'all_of()' or 'any_of()' instead.
##
     # Was:
##
     data %>% select(genes_list)
##
##
    # Now:
##
     data %>% select(all_of(genes_list))
## See <https://tidyselect.r-lib.org/reference/faq-external-vector.html>.
## This warning is displayed once every 8 hours.
## Call 'lifecycle::last_lifecycle_warnings()' to see where this warning was
## generated.
```

## Distribution of ABI1 expression across samples



```
## Warning: 'aes_string()' was deprecated in ggplot2 3.0.0.
```

<sup>##</sup> i Please use tidy evaluation idioms with 'aes()'.

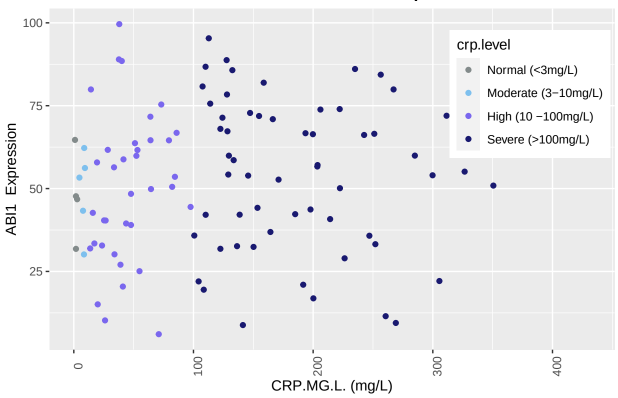
<sup>##</sup> i See also 'vignette("ggplot2-in-packages")' for more information.

<sup>##</sup> This warning is displayed once every 8 hours.

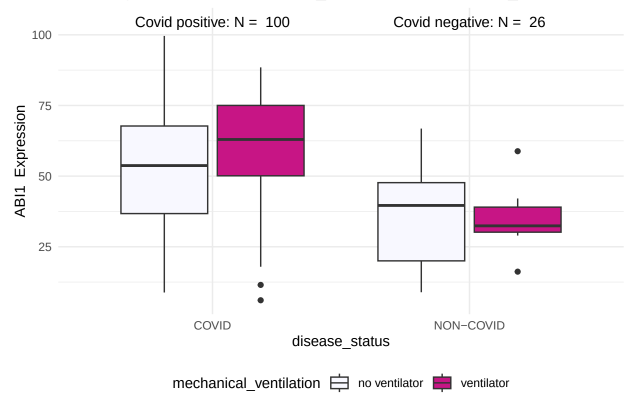
<sup>##</sup> Call 'lifecycle::last\_lifecycle\_warnings()' to see where this warning was

<sup>##</sup> generated.

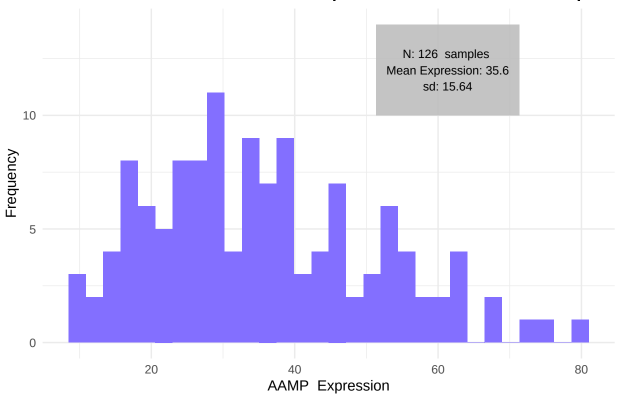
## CRP.MG.L. vs ABI1 Expression



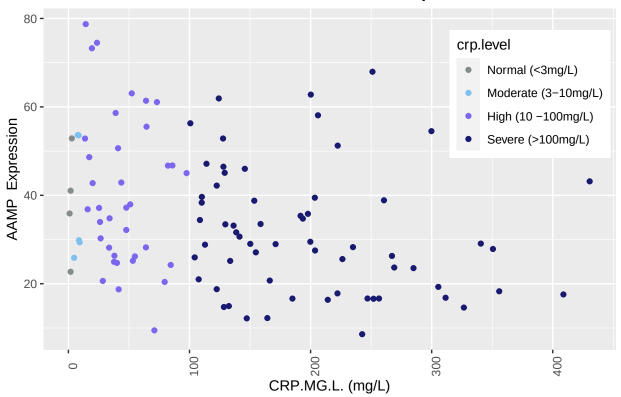
ABI1 Expression across disease\_status and mechanical\_ventilation



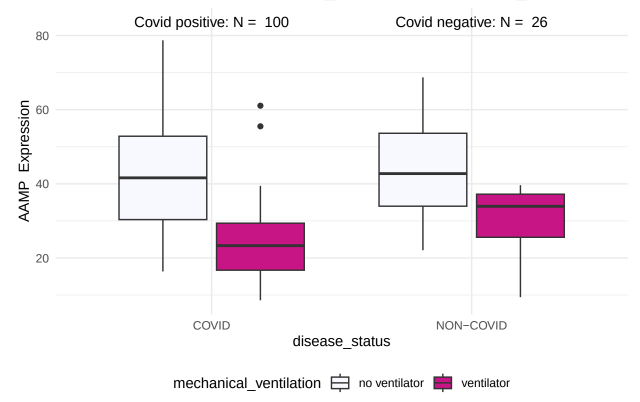
## Distribution of AAMP expression across samples



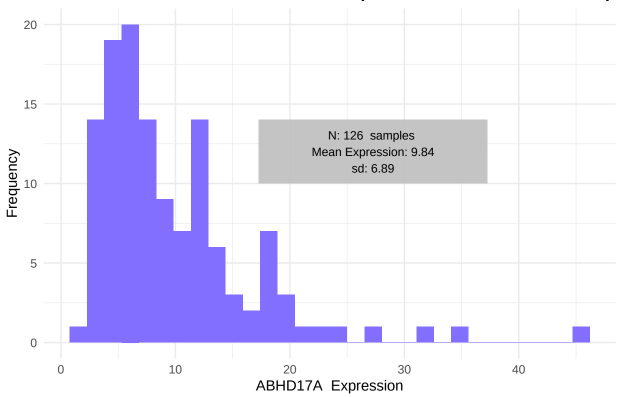
# CRP.MG.L. vs AAMP Expression



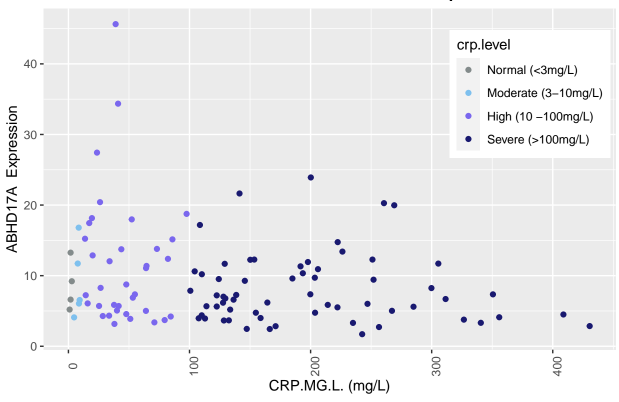
AAMP Expression across disease\_status and mechanical\_ventilation



## Distribution of ABHD17A expression across sampl



## CRP.MG.L. vs ABHD17A Expression



ABHD17A Expression across disease\_status and mechanical\_ventilation

