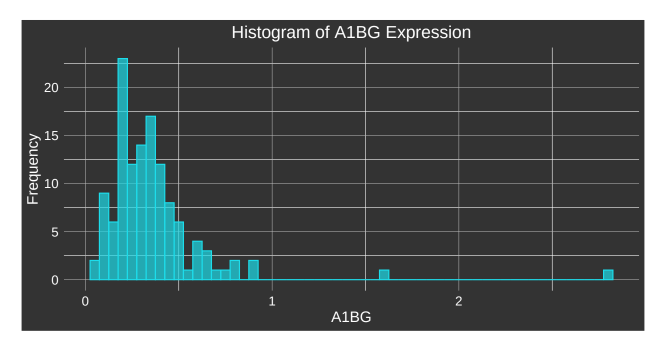
## final\_submission

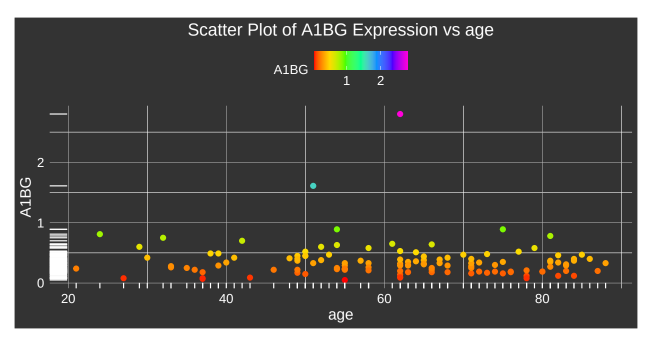
## 2024-08-29

```
# Create a function that calculate the mean and std
summary_continuous <- function(var, group) {</pre>
  data_frame %>%
   group_by(!!sym(group)) %>%
   summarise(
     Mean = mean(!!sym(var), na.rm = TRUE),
     SD = sd(!!sym(var), na.rm = TRUE),
   )
}
age_summary <- summary_continuous("age", "icu_status")</pre>
ferritin_summary <- summary_continuous("ferritin", "icu_status")</pre>
crp_summary <- summary_continuous("crp", "icu_status")</pre>
age_summary
## # A tibble: 2 x 3
## icu_status Mean
##
   58.7 17.8
## 1 no
## 2 yes
              63.2 13.9
ferritin_summary
## # A tibble: 2 x 3
##
   icu_status Mean
   <chr> <dbl> <dbl>
##
               716. 1068.
## 1 no
## 2 yes
              935. 1019.
crp_summary
## # A tibble: 2 x 3
## icu status Mean
## <chr> <dbl> <dbl>
                109. 94.4
## 1 no
               150. 106.
## 2 yes
# Create a function that calculate the count and percentage
summary_categorical <- function(var, group) {</pre>
 data frame %>%
   group_by(!!sym(group), !!sym(var)) %>%
 summarise(n = n()) \%>\%
```

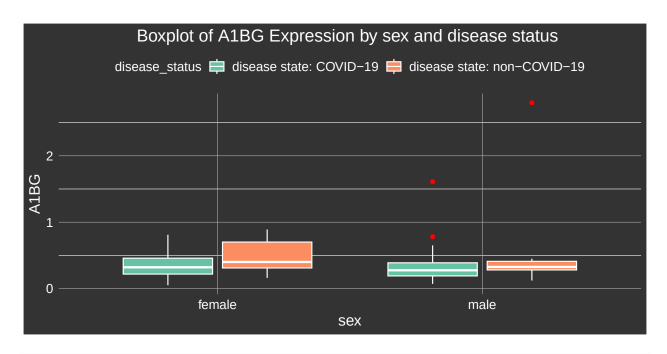
```
mutate(percentage = n / sum(n) * 100)
}
sex_summary <- summary_categorical("sex", "icu_status")</pre>
## 'summarise()' has grouped output by 'icu_status'. You can override using the
## '.groups' argument.
charlson summary <- summary categorical("disease status", "icu status")
## 'summarise()' has grouped output by 'icu status'. You can override using the
## '.groups' argument.
sex_summary
## # A tibble: 4 x 4
## # Groups: icu_status [2]
## icu_status sex n percentage
   <chr> <chr> <int>
                                 <dbl>
## 1 no
                                  45
              female 27
## 2 no
              \mathtt{male}
                        33
                                  55
## 3 yes
              female 24
                                  36.9
## 4 yes
              male 41
                                63.1
charlson summary
## # A tibble: 4 x 4
## # Groups: icu status [2]
    icu_status disease_status
##
                                               n percentage
##
    <chr>
             <chr>
                                           <int>
                                                      <dbl>
## 1 no
              disease state: COVID-19
                                              50
                                                       83.3
## 2 no
              disease state: non-COVID-19
                                              10
                                                       16.7
                                              50
## 3 yes
              disease state: COVID-19
                                                       76.9
## 4 yes
              disease state: non-COVID-19
                                              15
                                                       23.1
# Define a custom theme
custom_theme <- theme(</pre>
    panel.background = element_rect(fill = '#333333'),
   plot.title = element_text(hjust = 0.5, color = 'white', size = 16),
   axis.title = element_text(size = 14, color = 'white'),
   axis.text = element_text(size = 12, color = 'white'),
   plot.background = element_rect(fill = '#333333'),
   panel.grid.major = element_line(color = "gray", size = 0.1),
   legend.position = 'top',
   legend.background = element_rect(fill = "#333333"),
   legend.text = element_text(size = 12, color = 'white'),
   legend.title = element_text(size = 12, color = 'white'),
    legend.key = element_rect(fill = "#333333")
)
```



```
# Scatter Plot with rainbow colors
scatter_plot <- ggplot(data_frame,</pre>
                       aes(x = age,
                           y = A1BG,
                           color = A1BG)) +
   geom_point(size = 2) +
   geom_rug(color = "white") + # Add a rug plot
    scale_color_gradientn(colors = rainbow(7)) + # Apply rainbow colors
   labs(x = 'age', y = 'A1BG',
         title = paste("Scatter Plot of A1BG Expression vs age")) +
    custom_theme
# Save the scatter plot
ggsave(paste0("plots/plots_A1BG_scatter_plot.png"),
       plot = scatter_plot, width = 10, height = 6)
# Print the scatter plot
print(scatter_plot)
```



```
# Define labels for the legend
labels <- unique(data_frame[["disease_status"]])</pre>
# Create the palette dynamically based on the labels
color <- c("#66c2a5", "#fc8d62")</pre>
palette <- setNames(color, labels)</pre>
# https://www.statology.org/setnames-r/
# Boxplot
boxplot <- ggplot(data_frame, aes(x = sex,</pre>
                                   y = A1BG,
                                   fill = disease_status)) +
    geom_boxplot(outlier.color = "red", color = "white") +
    labs(x = "sex", y = "A1BG", fill = "disease_status" ,
         title = paste("Boxplot of A1BG Expression by sex and disease status ")) +
    scale_fill_manual(values = palette, labels = labels) +
    custom_theme
# Save the boxplot
ggsave(paste0("plots/plots_A1BG_boxplot.png"),
       plot = boxplot, width = 10, height = 6)
# Print the boxplot
print(boxplot)
```



```
genes_heat = as.data.frame(genes_df)
rownames(genes_heat) = genes_data$Genes
# filter out genes with O value
# such that we dont get -inf during log2 normalization
genes_heat <- genes_heat[rowSums(genes_heat == 0) == 0, ]</pre>
# select 15 genes
genes_heat <- genes_heat[1:15, ]</pre>
# perform log2 normalization
log2.genes_heat = log2(genes_heat)
# extract sex and icu_status from datast
annotation_data <- data_frame[, c("sex", "icu_status")]</pre>
# create heatmap with pheatmap
heat_map <- pheatmap(log2.genes_heat,</pre>
clustering_distance_cols = 'euclidean',
clustering_distance_rows = 'euclidean',
annotation_col = annotation_data,
fontsize_col = 4)
```

