My_Graphs

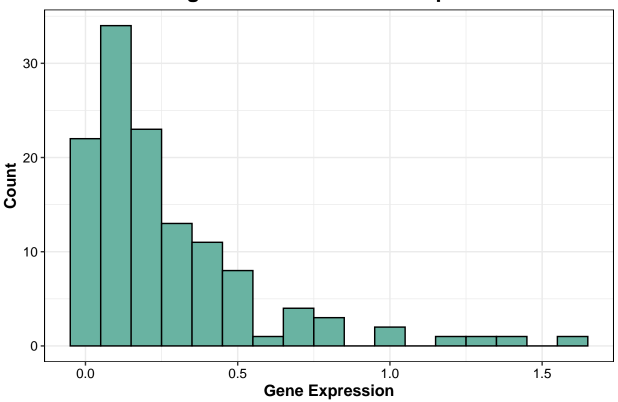
2024-07-25

```
# Load packages and tidyverse includes ggplot2
library(tidyverse)
## -- Attaching core tidyverse packages ----- tidyverse 2.0.0 --
## v dplyr 1.1.4
                       v readr
                                   2.1.5
## v forcats 1.0.0
                     v stringr 1.5.1
## v ggplot2 3.5.1
                      v tibble 3.2.1
## v lubridate 1.9.3
                                   1.3.1
                     v tidyr
## v purrr
              1.0.2
## -- Conflicts ----- tidyverse conflicts() --
## x dplyr::filter() masks stats::filter()
## x dplyr::lag()
                    masks stats::lag()
## i Use the conflicted package (<a href="http://conflicted.r-lib.org/">http://conflicted.r-lib.org/</a>) to force all conflicts to become error
# Pulling Data from the two provided files.
# row.names = 1 to indicate that the first column of the CSV file is names for the data frame.
gene_express_df <- read.csv("C:/Users/casha/Downloads/QBS103_GSE157103_genes.csv", row.names = 1)</pre>
metadata_df <- read.csv("C:/Users/casha/Downloads/QBS103_GSE157103_series_matrix.csv")</pre>
# Selecting specific columns of interest
metadata_relevant <- metadata_df %>%
 select(participant_id, geo_accession, age, disease_status, sex)
#Continuous Covariant = age
#Categorical Covariants = disease_status, sex
# Extract expression data for the ABCB4 gene
abcb4 expression <- gene express df %>%
 rownames_to_column("gene") %>%
 filter(gene == "ABCB4") %>% #Keep ONLY ABCB4 rows
#wide format to long format
 pivot_longer(cols = -gene, names_to = "participant_id", values_to = "expression") %>%
 select(-gene)
# Merge expression data with metadata
merged_df <- abcb4_expression %>%
#https://www.datacamp.com/tutorial/merging-data-r: How to merge 2 datasets in R
 merge(metadata_relevant, by = "participant_id")
# Ensure correct data types
merged_df <- merged_df %>%
 mutate(age = as.numeric(age), #make sure age is numeric
        sex = factor(sex),
```

disease_status = factor(disease_status))

```
## Warning: There was 1 warning in 'mutate()'.
## i In argument: 'age = as.numeric(age)'.
## Caused by warning:
## ! NAs introduced by coercion
# Histogram of ABCB4 gene expression
histogram <- ggplot(merged_df, aes(x = expression)) +
\verb|#https://www.datacamp.com/tutorial/make-histogram-basic-r|
  geom_histogram(binwidth = 0.1, fill = "#69b3a2", color = "black") + #binwidth is how wide we want eac
#changed colors to make it more like other plots I've seen in papers
#change labels
  labs(
   title = "Histogram of ABCB4 Gene Expression",
   x = "Gene Expression",
    y = "Count" #number which falls within THIS level of gene expression
  ) +
  theme_bw() +
  theme(
   plot.background = element_rect(fill = "white"),
    panel.background = element_rect(fill = "white"),
    panel.border = element_rect(color = "black", fill = NA),
   text = element_text(size = 12),
   plot.title = element_text(hjust = 0.5, face = "bold", size = 16),
   axis.title = element_text(face = "bold", size = 12),
    axis.text = element_text(color = "black", size = 10)
histogram
```

Histogram of ABCB4 Gene Expression



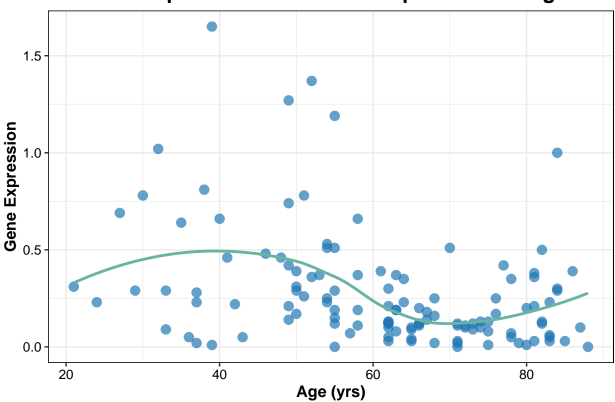
```
# Scatterplot of ABCB4 expression vs. age
scatterplot <- ggplot(merged_df, aes(x = age, y = expression)) +</pre>
     geom_point(color = "#1F77B4", size = 3, alpha = 0.7) + #using a clear color and changing the thickne
     geom_smooth(method = "loess", color = "#69b3a2", se = FALSE, size = 1) + #keeping the color scheme si
     # Add a regression line https://stackoverflow.com/questions/15633714/adding-a-regression-line-on-a-qq
# (decided to go with a loess instead of lm since a loess line shows the higher average gene expression
    labs(
         title = "Scatterplot of ABCB4 Gene Expression vs Age",
         x = "Age (yrs)", #continuous covariate
         y = "Gene Expression"
     ) +
     theme_bw() + #same theme for all graphs
         plot.background = element_rect(fill = "white"),
         panel.background = element_rect(fill = "white"),
         text = element_text(size = 12),
         plot.title = element_text(hjust = 0.5, face = "bold", size = 16), #keeping sizing consistent betwee
         axis.title = element_text(face = "bold", size = 12),
         axis.text = element_text(color = "black", size = 10),
          panel.border = element_rect(color = "black", fill = NA)
\#https://www.geeksforgeeks.org/add-panel-border-to-ggplot2-plot-in-r/ggplot2-plot-in-r/ggplot2-plot-in-r/ggplot2-plot-in-r/ggplot2-plot-in-r/ggplot2-plot-in-r/ggplot2-plot-in-r/ggplot2-plot-in-r/ggplot2-plot-in-r/ggplot2-plot-in-r/ggplot2-plot-in-r/ggplot2-plot-in-r/ggplot2-plot-in-r/ggplot2-plot-in-r/ggplot2-plot-in-r/ggplot2-plot-in-r/ggplot2-plot-in-r/ggplot2-plot-in-r/ggplot2-plot-in-r/ggplot2-plot-in-r/ggplot2-plot-in-r/ggplot2-plot-in-r/ggplot2-plot-in-r/ggplot2-plot-in-r/ggplot2-plot-in-r/ggplot2-plot-in-r/ggplot2-plot-in-r/ggplot2-plot-in-r/ggplot2-plot-in-r/ggplot2-plot-in-r/ggplot2-plot-in-r/ggplot2-plot-in-r/ggplot2-plot-in-r/ggplot2-plot-in-r/ggplot2-plot-in-r/ggplot2-plot-in-r/ggplot2-plot-in-r/ggplot2-plot-in-r/ggplot2-plot-in-r/ggplot2-plot-in-r/ggplot2-plot-in-r/ggplot2-plot-in-r/ggplot2-plot-in-r/ggplot2-plot-in-r/ggplot2-plot-in-r/ggplot2-plot-in-r/ggplot2-plot-in-r/ggplot2-plot-in-r/ggplot2-plot-in-r/ggplot2-plot-in-r/ggplot2-plot-in-r/ggplot-in-r/ggplot-in-r/ggplot-in-r/ggplot-in-r/ggplot-in-r/ggplot-in-r/ggplot-in-r/ggplot-in-r/ggplot-in-r/ggplot-in-r/ggplot-in-r/ggplot-in-r/ggplot-in-r/ggplot-in-r/ggplot-in-r/ggplot-in-r/ggplot-in-r/ggplot-in-r/ggplot-in-r/ggplot-in-r/ggplot-in-r/ggplot-in-r/ggplot-in-r/ggplot-in-r/ggplot-in-r/ggplot-in-r/ggplot-in-r/ggplot-in-r/ggplot-in-r/ggplot-in-r/ggplot-in-r/ggplot-in-r/ggplot-in-r/ggplot-in-r/ggplot-in-r/ggplot-in-r/ggplot-in-r/ggplot-in-r/ggplot-in-r/ggplot-in-r/ggplot-in-r/ggplot-in-r/ggplot-in-r/ggplot-in-r/ggplot-in-r/ggplot-in-r/ggplot-in-r/ggplot-in-r/ggplot-in-r/ggplot-in-r/ggplot-in-r/ggplot-in-r/ggplot-in-r/ggplot-in-r/ggplot-in-r/ggplot-in-r/ggplot-in-r/ggplot-in-r/ggplot-in-r/ggplot-in-r/ggplot-in-r/ggplot-in-r/ggplot-in-r/ggplot-in-r/ggplot-in-r/ggplot-in-r/ggplot-in-r/ggplot-in-r/ggplot-in-r/ggplot-in-r/ggplot-in-r/ggplot-in-r/ggplot-in-r/ggplot-in-r/ggplot-in-r/ggplot-in-r/ggplot-in-r/ggplot-in-r/ggplot-in-r/ggplot-in-r/ggplot-in-r/ggplot-in-r/ggplot-in-r/ggplot-in-r/ggplot-in-r-ggplot-in-r-ggplot-in-r-ggplot-in-r-ggplot-in-r-ggplot-
#don't fill border or no graph appears
            panel.grid = element_line(color = "grey") : decided to remove as did not help but learned a new te
#https://r-charts.com/ggplot2/grid/
```

```
## Warning: Using 'size' aesthetic for lines was deprecated in ggplot2 3.4.0.
## i Please use 'linewidth' instead.
## This warning is displayed once every 8 hours.
## Call 'lifecycle::last_lifecycle_warnings()' to see where this warning was
## generated.
```

scatterplot

```
## 'geom_smooth()' using formula = 'y ~ x'
## Warning: Removed 2 rows containing non-finite outside the scale range
## ('stat_smooth()').
## Warning: Removed 2 rows containing missing values or values outside the scale range
## ('geom_point()').
```

Scatterplot of ABCB4 Gene Expression vs Age



```
#Boxplot
boxplot <- ggplot(merged_df,aes(x = disease_status, y = expression, fill = sex)) +
#Add box plot
geom_boxplot() +
#Define colors: Have to use three colors because sex is female, male, and unknown
scale_fill_manual(values = c('#1F77B4', '#69b3a2', 'yellow')) +
#using the same color scheme
labs(</pre>
```

```
title = "Boxplot of ABCB4 Expression by Disease Status and Sex"
  ) +
  theme bw() #same theme
  theme(
   text = element_text(size = 12),
    plot.title = element_text(hjust = 0.5, face = "bold", size = 16), #keeping sizing consistent betwee
    axis.title = element_text(face = "bold", size = 12),
    axis.text = element_text(color = "black", size = 10),
    panel.border = element_rect(color = "black", fill = NA)
## List of 5
                 :List of 11
## $ text
    ..$ family
                   : NULL
##
     ..$ face
                     : NULL
##
     ..$ colour
                     : NULL
##
     ..$ size
                    : num 12
##
     ..$ hjust
                    : NULL
                    : NULL
##
     ..$ vjust
##
     ..$ angle
                     : NULL
##
     ..$ lineheight : NULL
##
     ..$ margin
                    : NULL
##
                    : NULL
     ..$ debug
     ..$ inherit.blank: logi FALSE
##
##
    ..- attr(*, "class")= chr [1:2] "element_text" "element"
## $ axis.title :List of 11
##
    ..$ family
                    : NULL
##
    ..$ face
                    : chr "bold"
##
    ..$ colour
                    : NULL
##
     ..$ size
                    : num 12
     ..$ hjust
                     : NULL
##
##
     ..$ vjust
                    : NULL
##
     ..$ angle
                    : NULL
##
     ..$ lineheight : NULL
##
     ..$ margin
                     : NULL
##
                    : NULL
     ..$ debug
##
     ..$ inherit.blank: logi FALSE
##
     ..- attr(*, "class")= chr [1:2] "element_text" "element"
##
    $ axis.text
                :List of 11
##
    ..$ family
                  : NULL
                    : NULL
     ..$ face
##
                     : chr "black"
##
     ..$ colour
                     : num 10
##
     ..$ size
##
     ..$ hjust
                    : NULL
##
     ..$ vjust
                     : NULL
                     : NULL
##
     ..$ angle
##
     ..$ lineheight
                    : NULL
##
     ..$ margin
                   : NULL
##
                    : NULL
     ..$ debug
    ..$ inherit.blank: logi FALSE
##
##
   ..- attr(*, "class")= chr [1:2] "element_text" "element"
## $ panel.border:List of 5
```

##

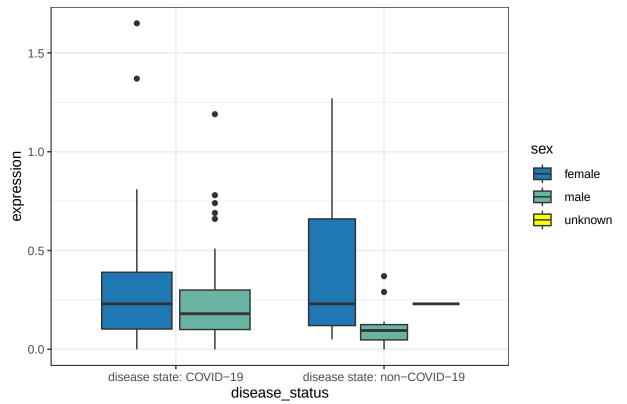
..\$ fill

: logi NA

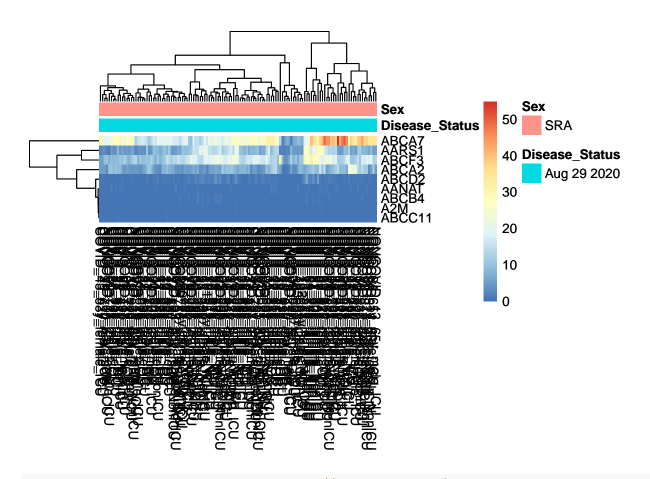
```
..$ colour
                 : chr "black"
##
     ..$ linewidth
                    : NULL
##
     ..$ linetype
                    : NULL
##
##
     ..$ inherit.blank: logi FALSE
     ..- attr(*, "class")= chr [1:2] "element_rect" "element"
##
##
    $ plot.title :List of 11
##
     ..$ family
                     : NULL
     ..$ face
                      : chr "bold"
##
##
     ..$ colour
                     : NULL
     ..$ size
##
                     : num 16
     ..$ hjust
                     : num 0.5
##
     ..$ vjust
                     : NULL
                     : NULL
##
     ..$ angle
     ..$ lineheight
                    : NULL
##
##
     ..$ margin
                     : NULL
                     : NULL
##
     ..$ debug
##
     ..$ inherit.blank: logi FALSE
   ..- attr(*, "class")= chr [1:2] "element_text" "element"
## - attr(*, "class")= chr [1:2] "theme" "gg"
## - attr(*, "complete") = logi FALSE
  - attr(*, "validate")= logi TRUE
```

boxplot

Boxplot of ABCB4 Expression by Disease Status and Sex



```
library(pheatmap)
# MANY ways to make a heatmap. Googled anything I was unsure about and linked the results in the docume
#https://davetang.org/muse/2018/05/15/making-a-heatmap-in-r-with-the-pheatmap-package/ - How to make a
# Calling the data under different names for the sake of simplicity
genes_df <- read.csv("C:/Users/casha/Downloads/QBS103_GSE157103_genes.csv", row.names = 1)</pre>
metadata df <- read.csv("C:/Users/casha/Downloads/QBS103 GSE157103 series matrix.csv", sep = "\t", head
# Pick 10 genes of interest
genes_of_interest <- c("A2M", "ABCB4", "AANAT", "AARS1", "ABCA2", "ABCB4", "ABCC11", "ABCD2", "ABCA7",
filtered_genes_df <- genes_df[rownames(genes_df) %in% genes_of_interest, ]
# Separate the metadata
metadata_processed <- metadata_df %>%
  separate(V1, into = c("Sample", "GSM", "Public", "Source_Name", "Disease_Status", "Sex", "Additional_
  select(Sample, Disease_Status, Sex)
## Warning: Expected 7 pieces. Additional pieces discarded in 127 rows [1, 2, 3, 4, 5, 6,
## 7, 8, 9, 10, 11, 12, 13, 14, 15, 16, 17, 18, 19, 20, ...].
#SYSTEM OF CHECKS
# Remove rows with NA values in essential columns or will cause errors
metadata_processed <- metadata_processed %>% drop_na(Sample, Disease_Status, Sex)
# Check and match the number of columns
metadata_processed <- metadata_processed %>%
  filter(Sample %in% colnames(filtered_genes_df))
# Ensure that the sample names in metadata match the column names in the gene expression data
filtered_genes_df <- filtered_genes_df[, metadata_processed$Sample]</pre>
# Create annotation data
annotation <- data.frame(</pre>
  Disease_Status = metadata_processed$Disease_Status,
  Sex = metadata_processed$Sex
rownames(annotation) <- metadata_processed$Sample</pre>
# Generate heatmap using pheatmap
heatmap <- pheatmap(</pre>
  as.matrix(filtered_genes_df),
  annotation_col = annotation,
# How to cluster rows and columns in a heatmap https://www.geeksforgeeks.org/draw-heatmap-with-clusters
  clustering_distance_rows = "euclidean",
  clustering_distance_cols = "euclidean",
  show_rownames = TRUE,
  show_colnames = TRUE
heatmap
```



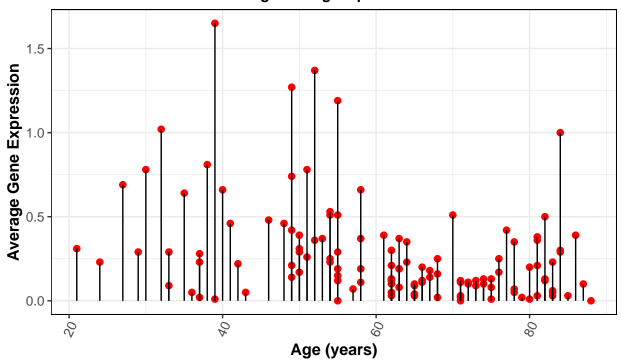
List of possible qqplot2 plots? - https://r-statistics.co/Top50-Gqplot2-Visualizations-MasterList-R-C theme_set(theme_bw()) # Plot lollipop <- ggplot(merged_df, aes(x = age, y = expression)) +</pre> geom_point(size=2, color = "red") + #most resembles a lollipop color geom_segment(aes(x=age, xend=age, y=0, yend=expression)) + labs(title="Lollipop Chart", x = "Age (years)",y = "Average Gene Expression", subtitle="Age Vs Avg. Expression", caption="source:QBS103_GSE157103") + #a way to cite where the info came from theme(axis.text.x = element_text(angle=65, vjust=0.6), plot.background = element_rect(fill = "white"), panel.background = element_rect(fill = "white"), panel.border = element_rect(color = "black", fill = NA), text = element_text(size = 12), plot.title = element_text(hjust = 0.5, face = "bold", size = 16), plot.subtitle = element_text(hjust = 0.5, face = "bold", size = 11), axis.title = element_text(face = "bold", size = 12),) lollipop

Warning: Removed 2 rows containing missing values or values outside the scale range
('geom_point()').

Warning: Removed 2 rows containing missing values or values outside the scale range
('geom_segment()').

Lollipop Chart

Age Vs Avg. Expression



source:QBS103_GSE157103

```
# Install and load packages
# install.packages("knitr")
# install.packages("kableExtra")
# install.packages("dplyr")

#load libraries
# https://bookdown.org/yihui/rmarkdown-cookbook/kable.html
library(dplyr)
library(knitr)
library(kableExtra) # Ensure kableExtra is loaded for kable_styling()

##
## Attaching package: 'kableExtra'

## The following object is masked from 'package:dplyr':
##
## group_rows
```

```
# Load necessary libraries
library(dplyr)
library(knitr)
series_matrix_df <- read.csv("C:/Users/casha/Downloads/QBS103_GSE157103_series_matrix.csv", stringsAsFa</pre>
#colnames(series_matrix_df) checking names because of errors
# Convert relevant columns to numeric as necessary
series_matrix_df$age <- as.numeric(gsub("[^0-9]", "", series_matrix_df$age))</pre>
series_matrix_df$ferritin.ng.ml <- as.numeric(series_matrix_df$`ferritin.ng.ml.`)</pre>
## Warning: NAs introduced by coercion
series_matrix_df$ddimer.mg.l_feu <- as.numeric(series_matrix_df$^ddimer.mg.l_feu.^)</pre>
## Warning: NAs introduced by coercion
# View summary to check for any issues
# summary(series_matrix_df)
# Group by sex and calculate summary statistics
summary_stats <- series_matrix_df %>%
  group_by(sex) %>%
  summarise(
   age_mean = mean(age, na.rm = TRUE),
   age_sd = sd(age, na.rm = TRUE),
   ferritin_median = median(ferritin.ng.ml, na.rm = TRUE),
   ferritin_iqr = IQR(ferritin.ng.ml, na.rm = TRUE),
   ddimer_median = median(ddimer.mg.l_feu, na.rm = TRUE),
   ddimer_iqr = IQR(ddimer.mg.l_feu, na.rm = TRUE),
   disease_status_n = n(),
   mechanical_ventilation_n = n()
  )
# Add percentages for categorical variables (n%)
summary_stats <- summary_stats %>%
 mutate(
   disease_status_percent = (disease_status_n / sum(disease_status_n)) * 100,
   mechanical_ventilation_percent = (mechanical_ventilation_n / sum(mechanical_ventilation_n)) * 100
  )
# View the summary statistics
# print(summary_stats)
# Render the summary statistics table using kable
\# How to construct a nice summary table in r? - https://rdrr.io/github/grayclhn/dbframe-R-library/man/b
kable(summary_stats, format = "latex", booktabs = TRUE, caption = "Summary Statistics Stratified by Sex
 kable_styling(latex_options = c("striped", "hold_position"))
```

Table 1: Summary Statistics Stratified by Sex

sex	age_mean	age_sd	$ferritin_median$	$ferritin_iqr$	${\rm ddimer_median}$	$\operatorname{ddimer_iqr}$	disease_status_n ı
female	59.88235	18.22158	318	547	1.37	5.86	51
male	62.64384	14.64901	755	849	2.21	10.55	74
unknown	83.00000	NA	NA	NA	NA	NA	1