Project_part3

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DATA CLEANING AND PREP

```
#DATA CLEANING AND PREP
# Set the working directory and read in the files
setwd("C:/Users/AlexandraTreml/Desktop/MS/QBS103/Project/QBS103_proj")
genes <- read.csv("genes_GSE157103.csv")</pre>
participant <- read.csv("series_matrix_GSE157103.csv")</pre>
# Rename the first column in genes to 'gene'
genes <- genes %>%
  rename(gene = X) %>%
  filter(gene == "A2M")
# Pivot the genes dataframe
genes_long <- genes %>%
  tidyr::pivot_longer(cols = -gene, names_to = 'id', values_to = 'expression')
#rename id column in participant, select my categorical covariates and continuous variable (+age for ed
participant <- participant %>%
  filter(icu_status == ' yes') %>%
  rename(id = participant_id)
  #select(id, age, sex, mechanical_ventilation, lactate.mmol.l.)
#join 2 dataframes
df <- left_join(participant, genes_long, by = "id")</pre>
#fill lactate unknowns with NA
df$lactate.mmol.l.[df$lactate.mmol.l. == " unknown"] <- NA
#fill sex unknowns with NA
df$sex[df$sex == " unknown"] <- NA</pre>
#remove : from age, and fill with NA
df$age[df$age == " :"] <- NA
df <- na.omit(df) #get rid of na
```

Part 1: Latex table of summary stats for all variables.

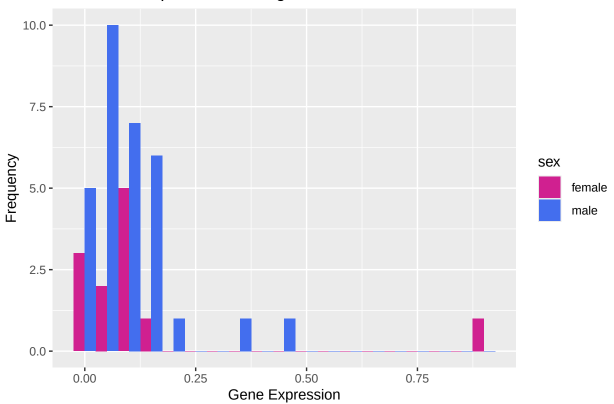
```
# Convert columns to appropriate types
df$fibrinogen <- as.numeric(df$fibrinogen)
```

Warning: NAs introduced by coercion

```
df$lactate.mmol.l. <- as.numeric(df$lactate.mmol.l.)</pre>
df$expression <- as.numeric(df$expression)</pre>
df$sex <- as.factor(df$sex)</pre>
df$mechanical_ventilation <- as.factor(df$mechanical_ventilation)</pre>
df$gene <- as.factor(df$gene)</pre>
df$crp.mg.l. <- as.numeric(df$crp.mg.l.)</pre>
## Warning: NAs introduced by coercion
# Summary of numeric variables
stargazer(df %% select(expression, lactate.mmol.l., fibrinogen, crp.mg.l.),
         type = "text",
         summary.stat = c("mean", "sd", "n"),
         digits = 2,
         out = "cont_var.tex")
##
## =============
             Mean St. Dev. N
## Statistic
## -----
## expression 0.11 0.13 54
## lactate.mmol.l. 1.64 1.39 54
## fibrinogen 501.82 216.97 45
## crp.mg.l.
                144.71 102.81 48
## -----
sex_counts <- df %>%
 group_by(sex) %>%
 summarize(
   count = n(),
   n_{percentage} = (n()/nrow(df)) *100)
gene_counts <- df %>%
 group_by(gene) %>%
 summarize(
   count = n(),
   n_{percentage} = (n()/nrow(df)) *100)
vent_counts <- df %>%
 group_by(mechanical_ventilation) %>%
 summarize(
   count = n(),
   n_{percentage} = (n()/nrow(df)) *100)
sex_counts <- sex_counts %>%
 mutate(variable = "Sex")
gene_counts <- gene_counts %>%
 mutate(variable = "Gene")
vent_counts <- vent_counts %>%
 mutate(variable = "Mechanical Ventilation")
```

```
cat_variables <- bind_rows(sex_counts, gene_counts, vent_counts)</pre>
cat_variables <- cat_variables%>%
 select(variable, count, n_percentage)
stargazer(cat_variables, type = "text", summary = FALSE, rownames = FALSE, out = "cat_variables.tex")
##
count n_percentage
## variable
## Sex
                         17 31.4814814814815
## Sex
                         37 68.5185185185
## Gene
                         54
                                    100
## Mechanical Ventilation 17 31.4814814815
## Mechanical Ventilation 37 68.5185185185
Part 2: Generate a publication quality histogram, boxplot, and scatterplot
df <- na.omit(df)</pre>
df <- df %>%
 select(id, gene, sex, age, mechanical_ventilation, expression, lactate.mmol.l.)
#head(df)
df$sex <- as.factor(df$sex)</pre>
levels(df$sex)
## [1] " female" " male"
ggplot(data = df, aes(x = expression, fill = sex)) +
 geom_histogram(binwidth = 0.05, position = 'dodge') +
 labs(x = 'Gene Expression', y = 'Frequency', title = 'A2M Gene Expression Among COVID ICU Patients')
 scale_fill_manual(values = c("violetred", "royalblue2"))
```

A2M Gene Expression Among COVID ICU Patients



```
# Convert lactate.mmol.l. to numeric
df$lactate.mmol.l. <- as.numeric(df$lactate.mmol.l.)</pre>
# Remove leading and trailing whitespaces from sex column
df$sex <- trimws(df$sex)</pre>
df$mechanical_ventilation <- trimws(df$mechanical_ventilation)</pre>
# Check unique levels
unique(df$sex)
                 "female"
```

[1] "male"

```
unique(df$mechanical_ventilation)
```

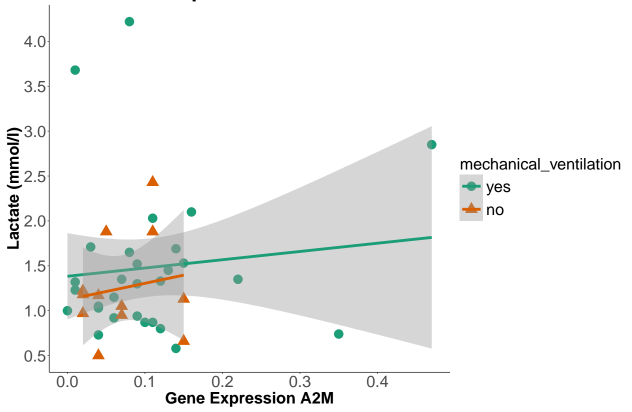
[1] "yes" "no"

```
# Set factor levels
df$sex <- factor(df$sex, levels = c("male", "female"))</pre>
df$mechanical_ventilation <- factor(df$mechanical_ventilation, levels = c("yes", "no"))
# Filter out values greater than 6.5 for the y-axis
df <- df %>% filter(lactate.mmol.1. <= 6.5)</pre>
```

```
df <- df %>% filter(expression <= 0.5)</pre>
# scatterplot vent
ggplot(df, aes(x = expression, y = lactate.mmol.l., color = mechanical_ventilation, shape = mechanical_
  geom_point(size = 3) +
  scale_y_continuous(breaks = seq(0, ceiling(max(df$lactate.mmol.l., na.rm = TRUE)), by = 0.5)) +
  geom_smooth(method = "lm", se = TRUE) +
  labs(x = 'Gene Expression A2M', y = 'Lactate (mmol/1)', title = ' A2M Gene Expression vs Lactate for
  theme_classic(base_size = 5) +
  scale_color_brewer(palette = 'Dark2') +
  #scale_color_manual(values = c("male" = "royalblue2", "female" = "violetred3")) +
  scale_shape_manual(values = c(16, 17)) +
   plot.title = element_text(size = 16, face = "bold"), # Center and bold title
   axis.title = element_text(size = 12, face = "bold"), # Bold axis titles
   axis.text = element_text(size = 12), # Increase axis text size
   legend.title = element_text(size = 12),
   legend.text = element_text(size = 12)
```

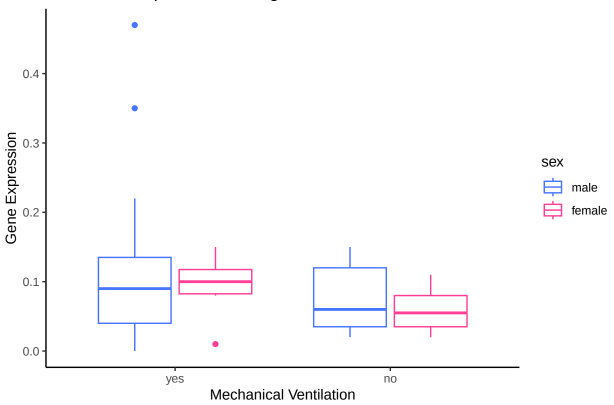
'geom_smooth()' using formula = 'y ~ x'

A2M Gene Expression vs Lactate for COVID ICU Patients



```
df$age <- as.numeric(df$age)</pre>
```

A2M Gene Expression Among Ventilated vs Non-Ventilated COVID ICU Pa



DATA PREP

```
genes_1 <- read.csv("genes_GSE157103.csv")
participant <- read.csv("series_matrix_GSE157103.csv")

genes_1 <- genes_1 %>%
    rename(gene = X)

# Pivot the genes dataframe
genes_long_1 <- genes_1 %>%
    tidyr::pivot_longer(cols = -gene, names_to = 'id', values_to = 'expression')
```

```
#rename id columm in participant, select my categorical covariates and continuous variable
participant <- participant %>%
    filter(icu_status == ' yes') %>%
    rename(id = participant_id)
    #select(id, age, sex, mechanical_ventilation, lactate.mmol.l.)

#join 2 dataframes
df1 <- left_join(participant, genes_long_1, by = "id")

#fill lactate unknowns with NA
df1$lactate.mmol.l.[df1$lactate.mmol.l. == " unknown"] <- NA
#fill sex unknowns with NA
df1$sex[df1$sex == " unknown"] <- NA
#remove : from age, and fill with NA
df1$age[df1$age == " :"] <- NA</pre>
df1 <- na.omit(df1) #get rid of na
```

Part 3: Generate a Heat Map

Aggregation function missing: defaulting to length



Part 4: A plot type we did not discuss in class

```
*percent of people in the ICU on mechanical ventilation
df$mechanical_ventilation <- trimws(df$mechanical_ventilation)</pre>
on_vent <- df %>%
  group_by(mechanical_ventilation) %>%
  summarize(count = n())
unique(df$mechanical_ventilation)
## [1] "yes" "no"
pie_chart <- ggplot(on_vent, aes(x = "", y = count, fill = mechanical_ventilation)) +</pre>
  geom_bar(width = 1, stat = "identity") +
  coord_polar("y") +
  labs(title = "Mechanical Ventilation Count Among ICU Patients") +
  theme_void() +
  scale_fill_manual(values = c("yes" = "darkorange1", "no" = "cornflowerblue")) +
   geom_text(aes(label = count),
            position = position_stack(vjust = 0.5))
print(pie_chart)
```

Mechanical Ventilation Count Among ICU Patients

