# Final project code

### Group F

2023-12-07

#### Load data

```
library(ggplot2)
library(dplyr)
## Attaching package: 'dplyr'
## The following objects are masked from 'package:stats':
##
##
      filter, lag
## The following objects are masked from 'package:base':
##
      intersect, setdiff, setequal, union
##
library(tidyr)
library(pvclust)
library(cluster)
library(factoextra)
## Welcome! Want to learn more? See two factoextra-related books at https://goo.gl/ve3WBa
data1 <- read.csv("data_1.csv", header=TRUE)</pre>
data1<- data1[,-1]
# Data source and structure description
cat("Data Structure Description:\n")
## Data Structure Description:
str(data1)
## 'data.frame':
                 14728 obs. of 15 variables:
## $ M2ID
                : int 10005 10005 10005 10005 10005 10005 10005 10005 10015 10015 ...
## $ B1SPWBU2 : num 48 48 48 48 48 48 48 48 38 38 ...
              : int 0 0 0 0 0 0 0 0 126250 126250 ...
## $ B1STINC1
## $ B1PAGE_M2.x: int 80 80 80 80 80 80 80 53 53 ...
## $ B1PGENDER : int 2 2 2 2 2 2 2 2 2 2 ...
## $ B2DN_STR : int 0 0 0 0 0 0 0 0 0 ...
## $ race
                : int 1 1 1 1 1 1 1 1 1 1 ...
                : int 000000011...
## $ marital
## $ B1SQ2
              : int 10 10 10 10 10 10 10 10 8 8 ...
## $ B1SQ1
               : int 10 10 10 10 10 10 10 10 7 7 ...
## $ B1SQ3
                : int 10 10 10 10 10 10 10 9 9 ...
```

```
## $ B2DNEGAV : num 0 0 0 0.07 0 0 0 0.14 0 ...
## $ meanNA : num 0.00875 0.00875 0.00875 0.00875 0.00875 ...
              : num 0.0247 0.0247 0.0247 0.0247 0.0247 ...
## $ sdNA
## $ gender
               : int 00000000000...
```

#### EDA

```
# Load necessary libraries for visualization
library(ggplot2)
# EDA
cat("Exploratory Data Analysis with Descriptive Statistics and Visualizations:\n")
```

The descriptive statistical analysis aims to understand sample characteristics

labs(title = "Age Distribution", x = "Age", y = "Frequency")

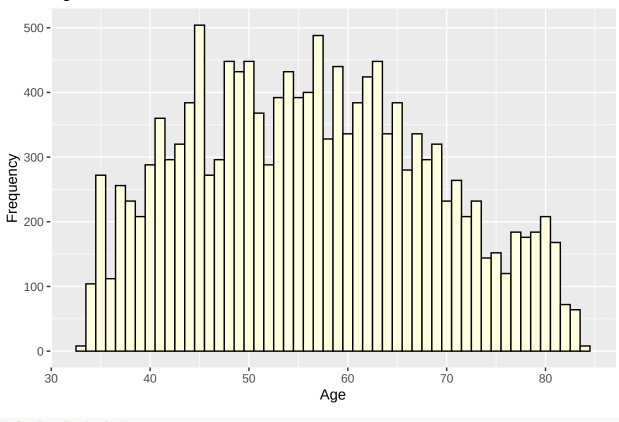
## Exploratory Data Analysis with Descriptive Statistics and Visualizations:

```
# Age Distribution
cat("Age Distribution:\n")
```

## Age Distribution:

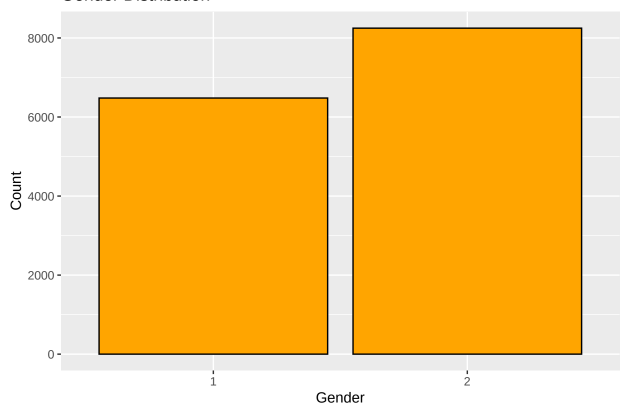
```
summary(data1$B1PAGE_M2.x, na.rm = TRUE)
##
     Min. 1st Qu. Median
                              Mean 3rd Qu.
                                              Max.
            47.00
                    56.00
                                             84.00
##
                             56.51
                                     65.00
sd(data1$B1PAGE_M2.x, na.rm = TRUE)
## [1] 12.2322
ggplot(data1, aes(x = B1PAGE_M2.x)) +
  geom_histogram(binwidth = 1, fill = "lightyellow", color = "black") +
```





```
# Gender Distribution
cat("Gender Distribution:\n")
## Gender Distribution:
summary(data1$B1PGENDER, na.rm = TRUE)
     Min. 1st Qu. Median
                              Mean 3rd Qu.
##
                                              Max.
##
      1.00
              1.00
                      2.00
                              1.56
                                      2.00
                                              2.00
ggplot(data1, aes(x = factor(B1PGENDER))) +
  geom_bar(fill = "orange", color = "black") +
  labs(title = "Gender Distribution", x = "Gender", y = "Count")
```

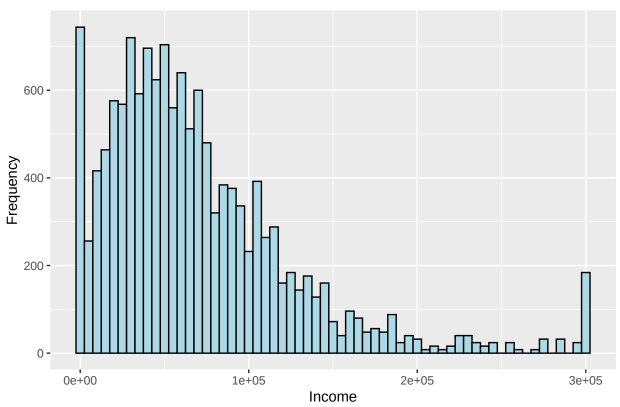
### **Gender Distribution**



```
# Income Distribution
cat("Income Distribution:\n")
## Income Distribution:
summary(data1$B1STINC1, na.rm = TRUE)
##
      Min. 1st Qu. Median
                              Mean 3rd Qu.
                                              Max.
                                                      NA's
         0
            30178
                    57500
##
                             70508
                                     93750 300000
                                                       904
sd(data1$B1STINC1, na.rm = TRUE)
## [1] 57837.37
ggplot(data1, aes(x = B1STINC1)) +
  geom_histogram(binwidth = 5000, fill = "lightblue", color = "black") +
  labs(title = "Income Distribution", x = "Income", y = "Frequency")
```

## Warning: Removed 904 rows containing non-finite values (`stat\_bin()`).

## **Income Distribution**



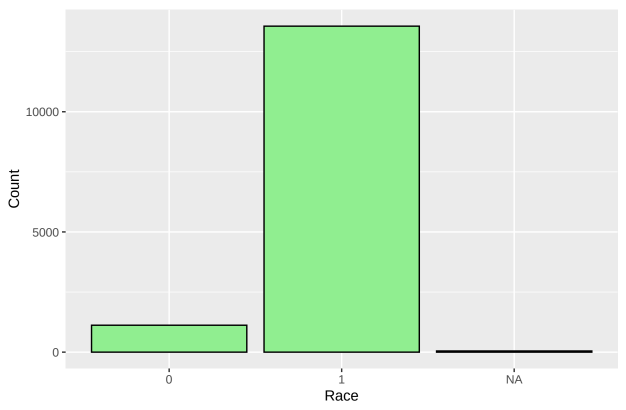
```
# Race Distribution
cat("Race Distribution:\n")

## Race Distribution:
summary(data1$race, na.rm = TRUE)

## Min. 1st Qu. Median Mean 3rd Qu. Max. NA's
## 0.0000 1.0000 1.0000 0.9237 1.0000 1.0000 48

ggplot(data1, aes(x = factor(race))) +
   geom_bar(fill = "lightgreen", color = "black") +
   labs(title = "Race Distribution", x = "Race", y = "Count")
```

## Race Distribution



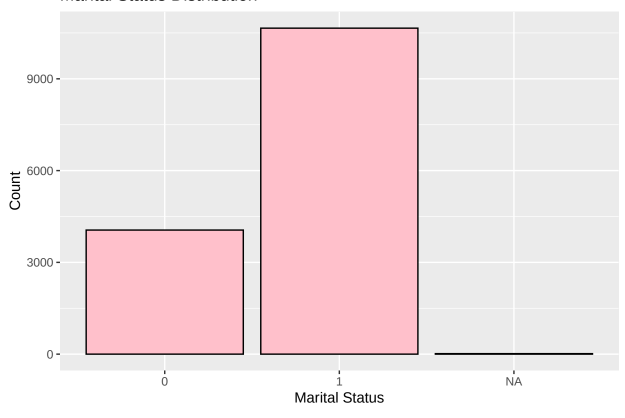
```
# Marital Status Distribution:\n")

## Marital Status Distribution:
summary(data1$marital, na.rm = TRUE)

## Min. 1st Qu. Median Mean 3rd Qu. Max. NA's
## 0.0000 0.0000 1.0000 0.7243 1.0000 1.0000 16

ggplot(data1, aes(x = factor(marital))) +
   geom_bar(fill = "pink", color = "black") +
   labs(title = "Marital Status Distribution", x = "Marital Status", y = "Count")
```

### Marital Status Distribution

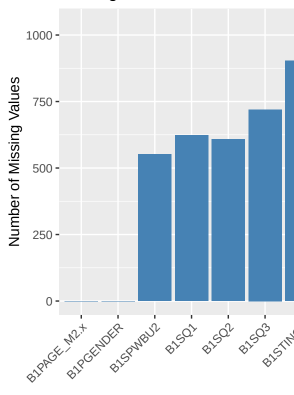


### Missing value Handling

```
# Create a data frame with the number of missing values for each variable
na_counts <- data1 %>% summarise_all(~sum(is.na(.))) %>% gather(key = "Variable", value = "NA_Count")

# Draw a bar chart of missing values
ggplot(na_counts, aes(x = Variable, y = NA_Count)) +
    geom_bar(stat = "identity", fill = "steelblue") +
    theme(axis.text.x = element_text(angle = 45, hjust = 1)) +
    labs(title = "Missing Values in Each Variable", x = "Variable", y = "Number of Missing Values")
```

### Missing Values in Each Variable



First, visually display the missing values, and then process them.

```
# Function to calculate the mode
get_mode <- function(v) {</pre>
  uniqv <- unique(na.omit(v))</pre>
  uniqv[which.max(tabulate(match(v, uniqv)))]
# Missing value handling
# Use median to fill missing values for continuous variables
data1$B1SPWBU2[is.na(data1$B1SPWBU2)] <- median(data1$B1SPWBU2, na.rm = TRUE)
data1$B1STINC1[is.na(data1$B1STINC1)] <- median(data1$B1STINC1, na.rm = TRUE)
data1$B2DNEGAV[is.na(data1$B2DNEGAV)] <- median(data1$B2DNEGAV, na.rm = TRUE)
# Use mode to fill missing values for categorical variables
data1$B2DN_STR[is.na(data1$B2DN_STR)] <- get_mode(data1$B2DN_STR)</pre>
data1$race[is.na(data1$race)] <- get_mode(data1$race)</pre>
data1$marital[is.na(data1$marital)] <- get_mode(data1$marital)</pre>
# For rating variables (assuming a 1-10 scale), use median or mode
data1$B1SQ2[is.na(data1$B1SQ2)] <- median(data1$B1SQ2, na.rm = TRUE)</pre>
data1$B1SQ1[is.na(data1$B1SQ1)] <- median(data1$B1SQ1, na.rm = TRUE)</pre>
data1$B1SQ3[is.na(data1$B1SQ3)] <- median(data1$B1SQ3, na.rm = TRUE)</pre>
# Print the updated data to check
sum(is.na(data1))
```

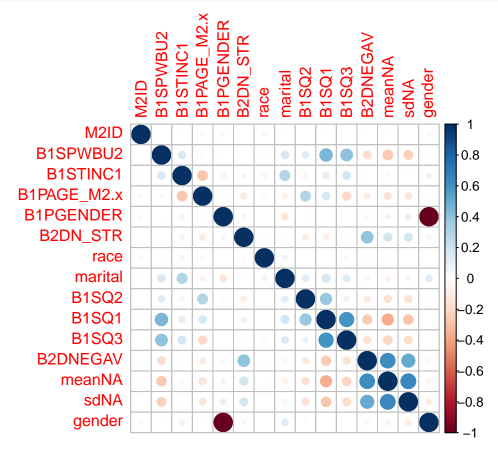
**##** [1] 0

#### More dimensions of data visualization

```
# correlation matrix - Correlation between variables
library(corrplot)

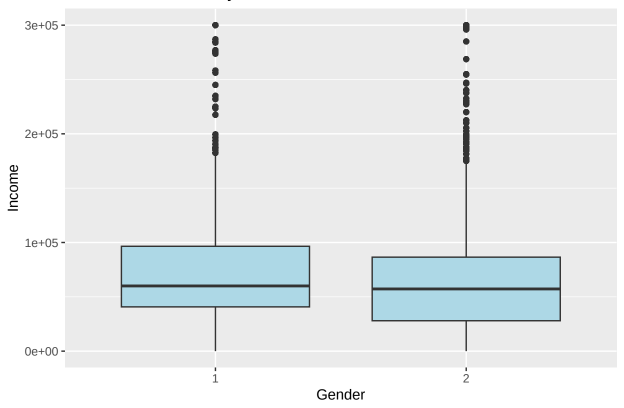
## corrplot 0.92 loaded

M <- cor(data1, use = "complete.obs")
corrplot(M, method = "circle")</pre>
```



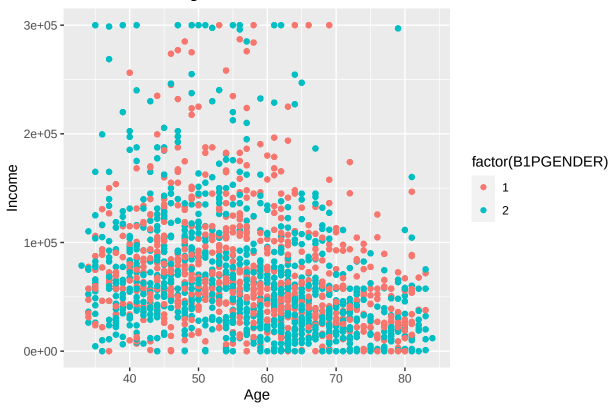
```
# Comparative analysis - Income distribution by sex
ggplot(data1, aes(x = factor(B1PGENDER), y = B1STINC1)) +
  geom_boxplot(fill = "lightblue") +
  labs(title = "Income Distribution by Gender", x = "Gender", y = "Income")
```

# Income Distribution by Gender



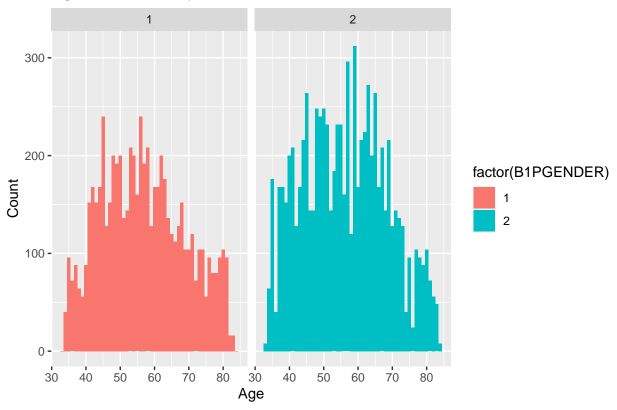
```
# Correlation analysis - Scatter plot of age and income
ggplot(data1, aes(x = B1PAGE_M2.x, y = B1STINC1)) +
  geom_point(aes(color = factor(B1PGENDER))) +
  labs(title = "Scatter Plot of Age vs Income", x = "Age", y = "Income")
```

# Scatter Plot of Age vs Income



```
# multivariate analysis - gender difference of income
ggplot(data1, aes(x = B1PAGE_M2.x, fill = factor(B1PGENDER))) +
  geom_histogram(binwidth = 1) +
  facet_wrap(~ B1PGENDER) +
  labs(title = "Age Distribution by Gender", x = "Age", y = "Count")
```

## Age Distribution by Gender



```
# save data into original path
write.csv(data1, "cleaned_data.csv", row.names = FALSE)
```

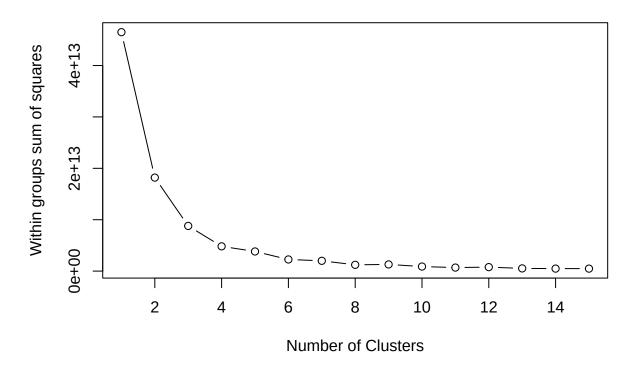
#### Cluster Analysis

```
library(cluster)
life_satisfaction <- na.exclude(data1[, c("B1SQ2", "B1SQ1", "B1SQ3")])
d <- dist(life_satisfaction, method = "euclidean")
fit_hc <- hclust(d, method="ward.D2")
clusters <- cutree(fit_hc, k=3)
data1$cluster <- clusters

# Perform K-means clustering

# Determine the optimal number of clusters
set.seed(123)
wss <- (nrow(data1)-1)*sum(apply(data1,2,var))
for (i in 2:15) wss[i] <- sum(kmeans(data1, centers=i)$withinss)

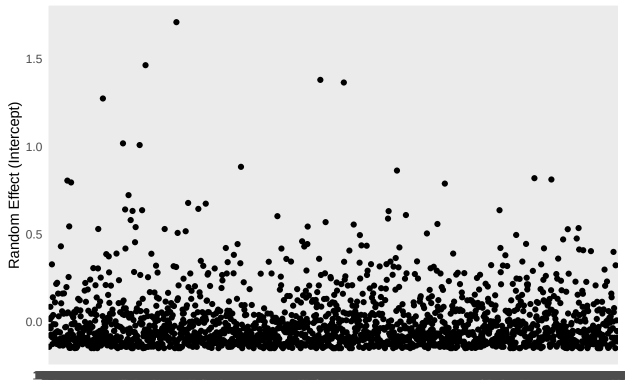
# Plot elbow method
plot(1:15, wss, type="b", xlab="Number of Clusters", ylab="Within groups sum of squares")</pre>
```



```
# Perform K-means with an appropriate number of clusters
set.seed(123)
kmeans_result <- kmeans(data1, centers=3)</pre>
data1$cluster <- as.factor(kmeans_result$cluster)</pre>
# Analyze the cluster results
table(data1$cluster)
##
##
      1
           2
                 3
## 4560 9360
Multilevel Analysis
library(lme4)
## Loading required package: Matrix
##
## Attaching package: 'Matrix'
## The following objects are masked from 'package:tidyr':
       expand, pack, unpack
##
library(ggplot2)
model_mcrm <- lmer(B2DNEGAV ~ cluster + (1|M2ID), data=data1)</pre>
summary(model_mcrm)
```

```
## Linear mixed model fit by REML ['lmerMod']
## Formula: B2DNEGAV ~ cluster + (1 | M2ID)
     Data: data1
##
## REML criterion at convergence: 1480.9
##
## Scaled residuals:
##
      Min
             1Q Median
                                3Q
                                       Max
## -6.5149 -0.4007 -0.1748 0.1698 10.5657
##
## Random effects:
## Groups Name
                         Variance Std.Dev.
             (Intercept) 0.03598 0.1897
## M2ID
## Residual
                         0.05103 0.2259
## Number of obs: 14728, groups: M2ID, 1841
##
## Fixed effects:
              Estimate Std. Error t value
## (Intercept) 0.17564
                           0.00862 20.375
## cluster2
              0.00143
                           0.01051
                                    0.136
## cluster3 -0.02502
                           0.02222 -1.126
##
## Correlation of Fixed Effects:
            (Intr) clstr2
## cluster2 -0.820
## cluster3 -0.388 0.318
# Extracting random effects for M2ID
rand_eff <- ranef(model_mcrm)$M2ID</pre>
rand_eff_df <- as.data.frame(rand_eff)</pre>
rand_eff_df$M2ID <- rownames(rand_eff_df)</pre>
# Plotting random effects
ggplot(rand_eff_df, aes(x=M2ID, y=`(Intercept)`)) +
  geom_point() +
 theme_minimal() +
 labs(title="Random Effects (Intercepts) for Each M2ID",
      x="M2ID",
      y="Random Effect (Intercept)")
```

## Random Effects (Intercepts) for Each M2ID



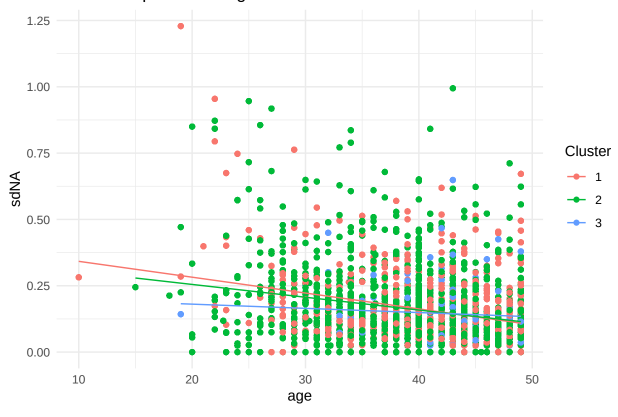
#### M2ID

```
Regression Analysis 1
```

```
model_ols <- lm(sdNA ~ cluster * B1SPWBU2, data=data1)
summary(model_ols)</pre>
```

```
##
## Call:
## lm(formula = sdNA ~ cluster * B1SPWBU2, data = data1)
##
## Residuals:
##
       Min
                1Q
                     Median
## -0.25501 -0.09272 -0.03306 0.05492 0.94055
##
##
  Coefficients:
                     Estimate Std. Error t value Pr(>|t|)
##
                    0.4018484 0.0130192 30.866 < 2e-16 ***
## (Intercept)
## cluster2
                   -0.0502626  0.0155249  -3.238  0.00121 **
## cluster3
                   ## B1SPWBU2
                              0.0003213 -18.603 < 2e-16 ***
                   -0.0059778
## cluster2:B1SPWBU2 0.0011490
                              0.0003882
                                          2.959 0.00309 **
## cluster3:B1SPWBU2 0.0043922
                              0.0008990
                                          4.886 1.04e-06 ***
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
##
## Residual standard error: 0.1422 on 14722 degrees of freedom
## Multiple R-squared: 0.05498, Adjusted R-squared: 0.05466
## F-statistic: 171.3 on 5 and 14722 DF, p-value: < 2.2e-16
```

## Relationship between age and sdNA across Clusters



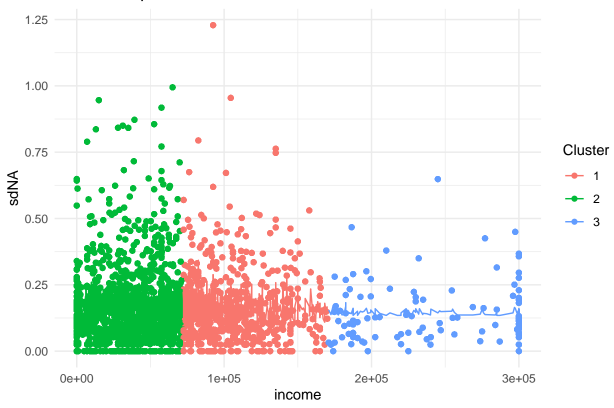
Regression Analysis 2

```
model_ols2 <- lm(sdNA ~ cluster * B1STINC1, data=data1)
summary(model_ols)</pre>
```

```
##
## Call:
## lm(formula = sdNA ~ cluster * B1SPWBU2, data = data1)
##
## Residuals:
## Min 1Q Median 3Q Max
## -0.25501 -0.09272 -0.03306 0.05492 0.94055
##
```

```
## Coefficients:
                     Estimate Std. Error t value Pr(>|t|)
##
                   0.4018484 0.0130192 30.866 < 2e-16 ***
## (Intercept)
## cluster2
                    -0.0502626 0.0155249 -3.238 0.00121 **
                    -0.1894491 0.0373259 -5.076 3.91e-07 ***
## cluster3
## B1SPWBU2
                    -0.0059778  0.0003213  -18.603  < 2e-16 ***
## cluster2:B1SPWBU2 0.0011490 0.0003882 2.959 0.00309 **
## cluster3:B1SPWBU2 0.0043922 0.0008990 4.886 1.04e-06 ***
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
## Residual standard error: 0.1422 on 14722 degrees of freedom
## Multiple R-squared: 0.05498, Adjusted R-squared: 0.05466
## F-statistic: 171.3 on 5 and 14722 DF, p-value: < 2.2e-16
# Create a new data frame for plotting
plot_data <- data1</pre>
plot_data$predicted_sdNA <- predict(model_ols, newdata = data1)</pre>
# Plotting
ggplot(plot_data, aes(x=B1STINC1, y=sdNA, color=factor(cluster))) +
 geom_point() + # Actual data points
 geom_line(aes(y=predicted_sdNA)) + # Regression lines
 theme minimal() +
 labs(title="Relationship between income and sdNA across Clusters",
      x="income",
      y="sdNA",
      color="Cluster")
```





#### Regression Analysis 3

```
model_ols3 <- lm(sdNA ~ cluster * B1PGENDER, data=data1)
summary(model_ols)</pre>
```

```
##
## Call:
## lm(formula = sdNA ~ cluster * B1SPWBU2, data = data1)
##
## Residuals:
##
        Min
                  1Q
                      Median
                                            Max
                                    3Q
## -0.25501 -0.09272 -0.03306 0.05492 0.94055
##
## Coefficients:
                      Estimate Std. Error t value Pr(>|t|)
##
                     0.4018484 0.0130192 30.866 < 2e-16 ***
## (Intercept)
## cluster2
                    -0.0502626  0.0155249  -3.238  0.00121 **
## cluster3
                     -0.1894491
                                0.0373259 -5.076 3.91e-07 ***
## B1SPWBU2
                                0.0003213 -18.603 < 2e-16 ***
                     -0.0059778
## cluster2:B1SPWBU2 0.0011490
                                0.0003882
                                            2.959 0.00309 **
## cluster3:B1SPWBU2 0.0043922
                                0.0008990
                                            4.886 1.04e-06 ***
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
##
## Residual standard error: 0.1422 on 14722 degrees of freedom
## Multiple R-squared: 0.05498, Adjusted R-squared: 0.05466
## F-statistic: 171.3 on 5 and 14722 DF, p-value: < 2.2e-16
```

# Relationship between gender and sdNA across Clusters

