

RWorksheet_Salinas#4b

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1. Using the for loop, create an R script that will display a 5x5 matrix as shown in Figure 1. It must contain vectorA = [1,2,3,4,5] and a 5 x 5 zero matrix.

```
vectorA <- c(1,2,3,4,5)

matrixA <- matrix(c(0,0,0,0,0), nrow = 5, ncol = 5)

for (i in 1:5){
  for(k in 1:5){
    if (i == k){
      matrixA[i,k] <- vectorA[i]
    }
  }
}
abs(matrixA)
```

```
##      [,1] [,2] [,3] [,4] [,5]
## [1,]    1    0    0    0    0
## [2,]    0    2    0    0    0
## [3,]    0    0    3    0    0
## [4,]    0    0    0    4    0
## [5,]    0    0    0    0    5
```

2. Print the string "*" using for() function. The output should be the same as shown in Figure 1.

```
for(i in 1:5) {
  numb <- rep("*", i)
  print(numb)
}
```

```
## [1] "*"
## [1] "*" "*"
## [1] "*" "*" "*"
## [1] "*" "*" "*" "*"
## [1] "*" "*" "*" "*" "*"
```

3. Get an input from the user to print the Fibonacci sequence starting from the 1st input up to 500. Use repeat and break statements. Write the R Scripts and its output.

```
start_num <- as.numeric(readline("Enter the starting number for the Fibonacci sequence: "))

## Enter the starting number for the Fibonacci sequence:
if (is.na(start_num)) {
  cat("Please enter a valid numeric starting number.\n")
} else {
  num1 <- 0
```

```

num2 <- 1

while (num2 <= 500) {
  if (!is.na(start_num) && num2 >= start_num) {
    cat(num2, " ")
  }

  fib_sum <- num1 + num2
  num1 <- num2
  num2 <- fib_sum
}

cat("\n")
}

```

Please enter a valid numeric starting number.

4 Import the dataset as shown in Figure 1 you have created previously.

4. A What is the R script for importing an excel or a csv file? Display the first 6 rows of the dataset? Show your codes and its result.

```

imprt <- read.csv("Household.csv")
head(imprt)

```

```

##   X ShoeSize Height Gender
## 1 1      6.5   66.0      F
## 2 2      9.0   68.0      F
## 3 3      8.5   64.5      F
## 4 4      8.5   65.0      F
## 5 5     10.5   70.0      M
## 6 6      7.0   64.0      F

```

4. B Create a subset for gender(female and male). How many observations are there in Male? How about in Female? Write the R scripts and its output.

```

library(readr)
Household <- read_csv("Household.csv")

```

```

## New names:
## Rows: 28 Columns: 4
## -- Column specification
## ----- Delimiter: "," chr
## (1): Gender dbl (3): ...1, ShoeSize, Height
## i Use `spec()` to retrieve the full column specification for this data. i
## Specify the column types or set `show_col_types = FALSE` to quiet this message.
## * `` -> `...1`

```

```

# Filter the data based on Gender
males <- Household[Household$Gender == "M",]
females <- Household[Household$Gender == "F",]

# Display the results
males

```

```

## # A tibble: 14 x 4
##   ...1 ShoeSize Height Gender
##   <dbl>   <dbl>   <dbl> <chr>

```

```
## 1      5      10.5    70    M
## 2      9      13      72    M
## 3     11     10.5    74.5  M
## 4     13      12      71    M
## 5     14     10.5    71    M
## 6     15      13      77    M
## 7     16     11.5    72    M
## 8     19      10      72    M
## 9     22       8.5    67    M
## 10    23     10.5    73    M
## 11    25     10.5    72    M
## 12    26      11      70    M
## 13    27       9      69    M
## 14    28      13      70    M
```

```
females
```

```
## # A tibble: 14 x 4
##   ...1 ShoeSize Height Gender
##   <dbl>    <dbl>  <dbl> <chr>
## 1      1      6.5    66    F
## 2      2       9     68    F
## 3      3      8.5   64.5  F
## 4      4      8.5    65    F
## 5      6       7     64    F
## 6      7      9.5    70    F
## 7      8       9     71    F
## 8     10      7.5    64    F
## 9     12      8.5    67    F
## 10    17      8.5    59    F
## 11    18       5     62    F
## 12    20      6.5    66    F
## 13    21      7.5    64    F
## 14    24      8.5    69    F
```

```
# Calculate the number of observations for each gender
```

```
observationF <- nrow(females)
```

```
observationM <- nrow(males)
```

```
# Display the number of observations
```

```
cat("Number of Female Observations:", observationF, "\n")
```

```
## Number of Female Observations: 14
```

```
cat("Number of Male Observations:", observationM, "\n")
```

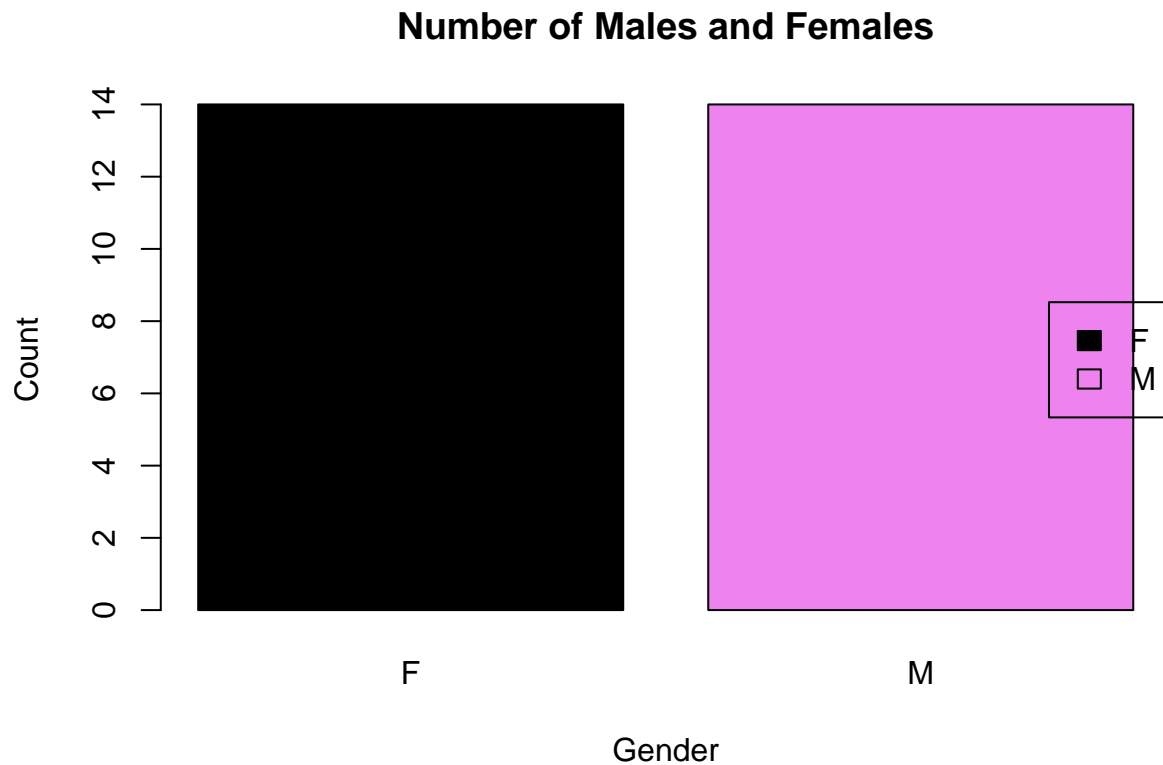
```
## Number of Male Observations: 14
```

4. C Create a graph for the number of males and females for Household Data. Use plot(), chart type = barplot. Make sure to place title, legends, and colors. Write the R scripts and its result.

```
total <- table(Household$Gender)
```

```
barplot(total,
```

```
  main = "Number of Males and Females", xlab = "Gender", ylab = "Count", col = c("black", "violet",
  legend("right", legend = rownames(total), fill = c("black", "violet"))
```



5. The monthly income of Dela Cruz family was spent on the following:

6. A Create a piechart that will include labels in percentage. Add some colors and title of the chart. Write the R scripts and show its output.

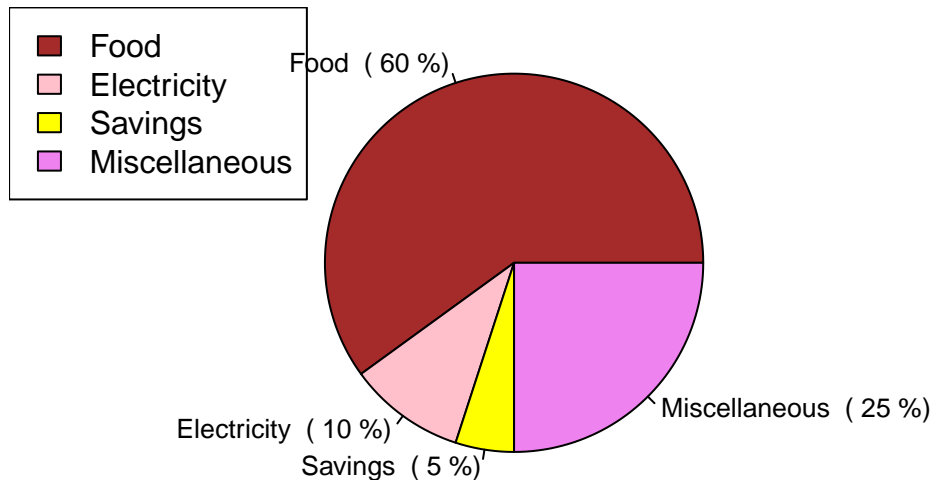
```
spend <- data.frame(
  Category = c("Food", "Electricity", "Savings", "Miscellaneous"),
  Value = c(60, 10, 5, 25)
)

spend$Percentage <- spend$Value / sum(spend$Value) * 100
colors <- c("brown", "pink", "yellow", "violet")

# Adjust the font size with the cex parameter
pie(spend$Value,
  labels = paste(spend$Category, " (", spend$Percentage, "%)",
  col = colors,
  main = "The Monthly Income Spending of Dela Cruz Family", cex = 0.8)

legend("topleft", spend$Category, fill = colors)
```

The Monthly Income Spending of Dela Cruz Family



6. Use the iris dataset.

A. Check for the structure of the dataset using the `str()` function. Describe what you have seen in the output.

```
str(iris)
```

```
## 'data.frame':  150 obs. of  5 variables:
## $ Sepal.Length: num  5.1 4.9 4.7 4.6 5 5.4 4.6 5 4.4 4.9 ...
## $ Sepal.Width : num  3.5 3 3.2 3.1 3.6 3.9 3.4 3.4 2.9 3.1 ...
## $ Petal.Length: num  1.4 1.4 1.3 1.5 1.4 1.7 1.4 1.5 1.4 1.5 ...
## $ Petal.Width : num  0.2 0.2 0.2 0.2 0.2 0.4 0.3 0.2 0.2 0.1 ...
## $ Species      : Factor w/ 3 levels "setosa","versicolor",...: 1 1 1 1 1 1 1 1 1 1 ...
```

I was curious and made a research about this. The output of the code provides information about the iris dataset, which is renowned for its use in data analysis and machine learning. It reveals the following details:

1. The dataset comprises 150 observations and 5 variables.
2. 'Sepal.Length' represents the sepal length of iris flowers.
3. 'Sepal.Width' represents the sepal width of iris flowers.
4. 'Petal.Length' signifies the petal length of iris flowers.
5. 'Petal.Width' signifies the petal width of iris flowers.
6. 'Species' This is the categorized variables.

B. Create an R object that will contain the mean of the sepal.length, sepal.width,petal.length,and petal.width. What is the R script and its result?

```
value_of_means <- c(
  lsepal <- mean(iris$Sepal.Length),
  wsepal <- mean(iris$Sepal.Width) ,
  lpetal <- mean(iris$Petal.Length),
  wpetal <- mean(iris$Petal.Width)
)
value_of_means
```

```
## [1] 5.843333 3.057333 3.758000 1.199333
```

C.Create a pie chart for the Species distribution. Add title, legends, and colors. Write the R script and its result.

```
# Assuming 'iris' is the name of your dataset
species <- table(iris$Species)
```

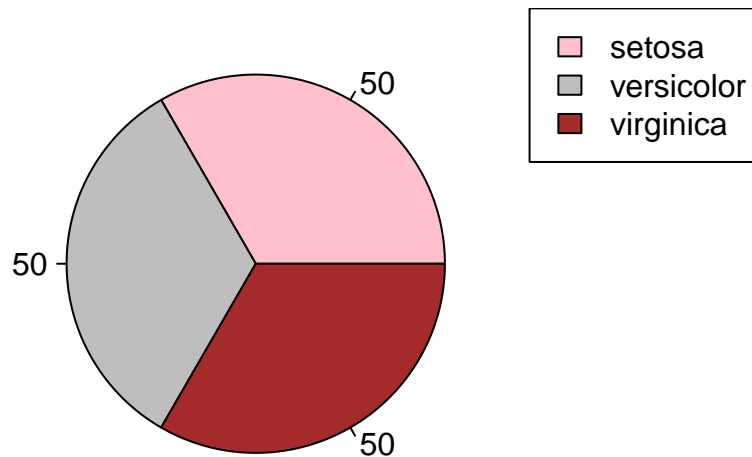
```

colors <- c("pink", "gray", "brown")

# Create the pie chart
pie(species, col = colors, labels = species )
legend("topright", legend = levels(iris$Species), fill = colors)
title("Species Distribution")

```

Species Distribution



D. Subset the species into setosa, versicolor, and virginica. Write the R scripts and show the last six (6) rows of each species.

```

#subset

setosa_subset <- iris[iris$Species == "setosa" ,]
versicolor_subset <- iris[iris$Species == "versicolor",]
virginica_subset <- iris[iris$Species == "virginica",]

#last 6 row each

tail(setosa_subset, 6)

```

```

##      Sepal.Length Sepal.Width Petal.Length Petal.Width Species
## 45          5.1         3.8         1.9         0.4   setosa
## 46          4.8         3.0         1.4         0.3   setosa
## 47          5.1         3.8         1.6         0.2   setosa
## 48          4.6         3.2         1.4         0.2   setosa
## 49          5.3         3.7         1.5         0.2   setosa
## 50          5.0         3.3         1.4         0.2   setosa

```

```

tail(versicolor_subset, 6)

```

```

##      Sepal.Length Sepal.Width Petal.Length Petal.Width   Species
## 95          5.6         2.7         4.2         1.3 versicolor
## 96          5.7         3.0         4.2         1.2 versicolor
## 97          5.7         2.9         4.2         1.3 versicolor
## 98          6.2         2.9         4.3         1.3 versicolor
## 99          5.1         2.5         3.0         1.1 versicolor
## 100         5.7         2.8         4.1         1.3 versicolor

```

```
tail(virginica_subset, 6)
```

```
##      Sepal.Length Sepal.Width Petal.Length Petal.Width  Species
## 145           6.7         3.3         5.7         2.5 virginica
## 146           6.7         3.0         5.2         2.3 virginica
## 147           6.3         2.5         5.0         1.9 virginica
## 148           6.5         3.0         5.2         2.0 virginica
## 149           6.2         3.4         5.4         2.3 virginica
## 150           5.9         3.0         5.1         1.8 virginica
```

E. Create a scatterplot of the sepal.length and sepal.width using the different species(setosa,versicolor,virginica). Add a title = “Iris Dataset”, subtitle = “Sepal width and length, labels for the x and y axis, the pch symbol and colors should be based on the species.

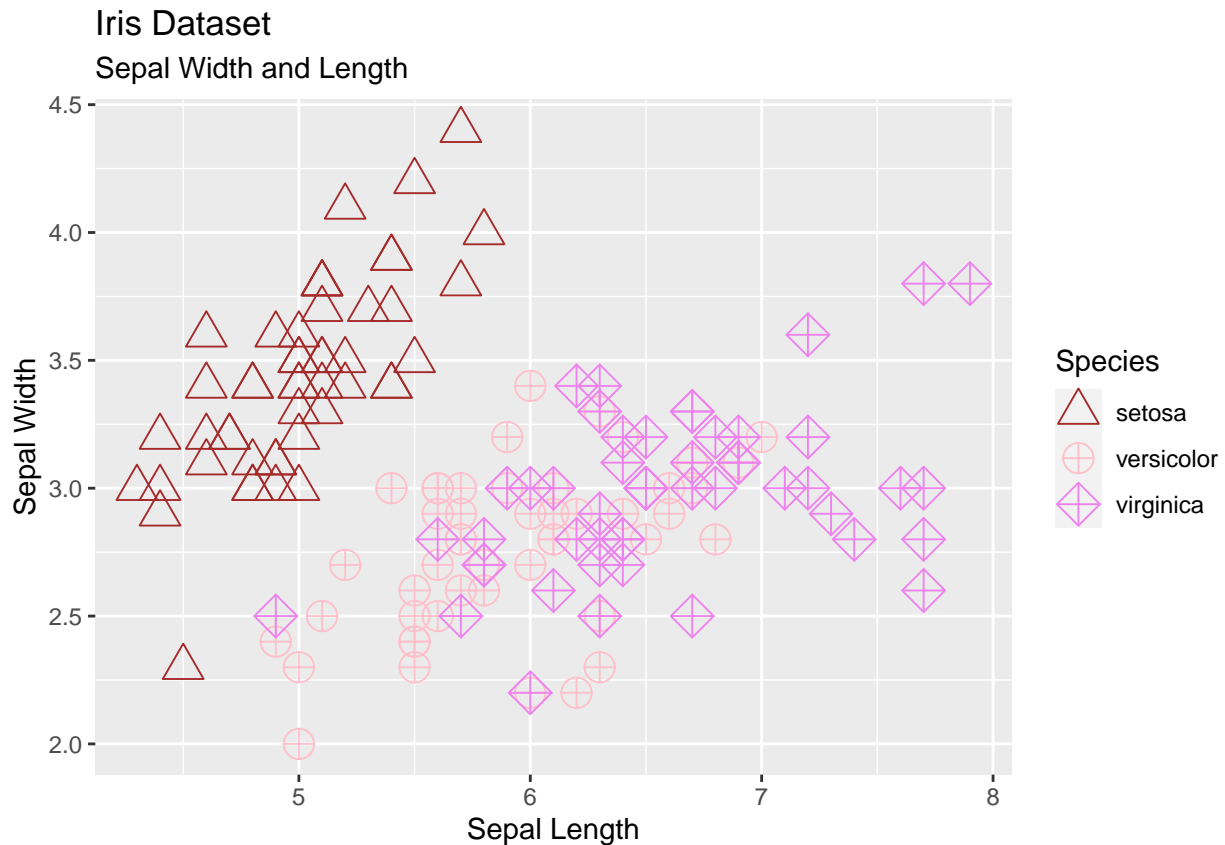
Hint: Need to convert to factors the species to store categorical variables.

```
library(ggplot2)

# factor
iris$Species <- as.factor(iris$Species)

# Create a scatterplot
scatterplot <- ggplot(iris, aes(x = Sepal.Length, y = Sepal.Width, color = Species, shape = Species)) +
  geom_point(size = 5) +
  labs(
    title = "Iris Dataset",
    subtitle = "Sepal Width and Length",
    x = "Sepal Length",
    y = "Sepal Width"
  ) +
  scale_color_manual(values = c("setosa" = "brown", "versicolor" = "pink", "virginica" = "violet")) +
  scale_shape_manual(values = c("setosa" = 2, "versicolor" = 10, "virginica" = 9))

print(scatterplot)
```



F. Interpret the result.

The plot shows a visual representation of the Sepal Length and Sepal Width for each Iris flower species. Each species is represented by a different color and shape.

Setosa flowers are brown and have a cross-like shape. Versicolor flowers are pink and have a circle shape. Virginica flowers are violet and have a diamond shape.

This plot allows you to easily compare and differentiate between different species based on their sepal length and sepal width, offering a comprehensive and visually appealing representation of the data.

7

```
library(readxl)
alexa_file <- read_excel("alexa_file.xlsx")
alexa_file
```

```
## # A tibble: 3,150 x 5
##   rating date          variation verified_reviews feedback
##   <dbl> <dtm>          <chr>          <chr>          <dbl>
## 1     5 2018-07-31 00:00:00 Charcoal Fabric Love my Echo!         1
## 2     5 2018-07-31 00:00:00 Charcoal Fabric Loved it!             1
## 3     4 2018-07-31 00:00:00 Walnut Finish   Sometimes while play~ 1
## 4     5 2018-07-31 00:00:00 Charcoal Fabric I have had a lot of ~ 1
## 5     5 2018-07-31 00:00:00 Charcoal Fabric Music                 1
## 6     5 2018-07-31 00:00:00 Heather Gray Fabric I received the echo ~ 1
## 7     3 2018-07-31 00:00:00 Sandstone Fabric Without having a cel~ 1
## 8     5 2018-07-31 00:00:00 Charcoal Fabric I think this is the ~ 1
## 9     5 2018-07-30 00:00:00 Heather Gray Fabric looks great         1
## 10    5 2018-07-30 00:00:00 Heather Gray Fabric Love it! I've listen~ 1
```



```
## # i 3,140 more rows
```

7 A. Rename the white and black variants by using `gsub()` function.

```
alexa_file$variation <- gsub("White Dot", "WhiteDot", alexa_file$variation)
alexa_file$variation <- gsub("White Show", "WhiteShow", alexa_file$variation)
alexa_file$variation <- gsub("White Plus", "WhitePlus", alexa_file$variation)
alexa_file$variation <- gsub("White Spot", "WhiteSpot", alexa_file$variation)

alexa_file$variation <- gsub("Black Dot", "BlacDot", alexa_file$variation)
alexa_file$variation <- gsub("Black Show", "BlackShow", alexa_file$variation)
alexa_file$variation <- gsub("Black Plus", "BlackPlus", alexa_file$variation)
alexa_file$variation <- gsub("Black Spot", "BlackSpot", alexa_file$variation)

alexa_file
```

```
## # A tibble: 3,150 x 5
```

```
##   rating date          variation      verified_reviews      feedback
##   <dbl> <dtm>          <chr>          <chr>          <dbl>
## 1     5 2018-07-31 00:00:00 Charcoal Fabric Love my Echo!          1
## 2     5 2018-07-31 00:00:00 Charcoal Fabric Loved it!              1
## 3     4 2018-07-31 00:00:00 Walnut Finish   Sometimes while play~   1
## 4     5 2018-07-31 00:00:00 Charcoal Fabric I have had a lot of ~   1
## 5     5 2018-07-31 00:00:00 Charcoal Fabric Music                  1
## 6     5 2018-07-31 00:00:00 Heather Gray Fabric I received the echo ~   1
## 7     3 2018-07-31 00:00:00 Sandstone Fabric Without having a cel~   1
## 8     5 2018-07-31 00:00:00 Charcoal Fabric I think this is the ~   1
## 9     5 2018-07-30 00:00:00 Heather Gray Fabric looks great          1
## 10    5 2018-07-30 00:00:00 Heather Gray Fabric Love it! I've listen~ 1
## # i 3,140 more rows
```

7 B. Get the total number of each variations and save it into another object. Save the object as `variations.RData`. Write the R scripts. What is its result?

```
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
```

```
# Perform the desired operations
```

```
variations <- alexa_file %>%
  count(alexa_file$variation)
```

```
variations
```

```
## # A tibble: 16 x 2
##   `alexa_file$variation`      n
##   <chr>          <int>
## 1 Black          261
```

```
## 2 Black Dot 516
## 3 Black Plus 270
## 4 Black Show 265
## 5 Black Spot 241
## 6 Charcoal Fabric 430
## 7 Configuration: Fire TV Stick 350
## 8 Heather Gray Fabric 157
## 9 Oak Finish 14
## 10 Sandstone Fabric 90
## 11 Walnut Finish 9
## 12 White 91
## 13 White Dot 184
## 14 White Plus 78
## 15 White Show 85
## 16 White Spot 109
```

```
save(variations, file = "variations.RData")
```

7 C. From the variations.RData, create a barplot(). Complete the details of the chart which include the title, color, labels of each bar.

```
# Load the variations data
load("variations.RData")
```

```
# Extract the variation names
```

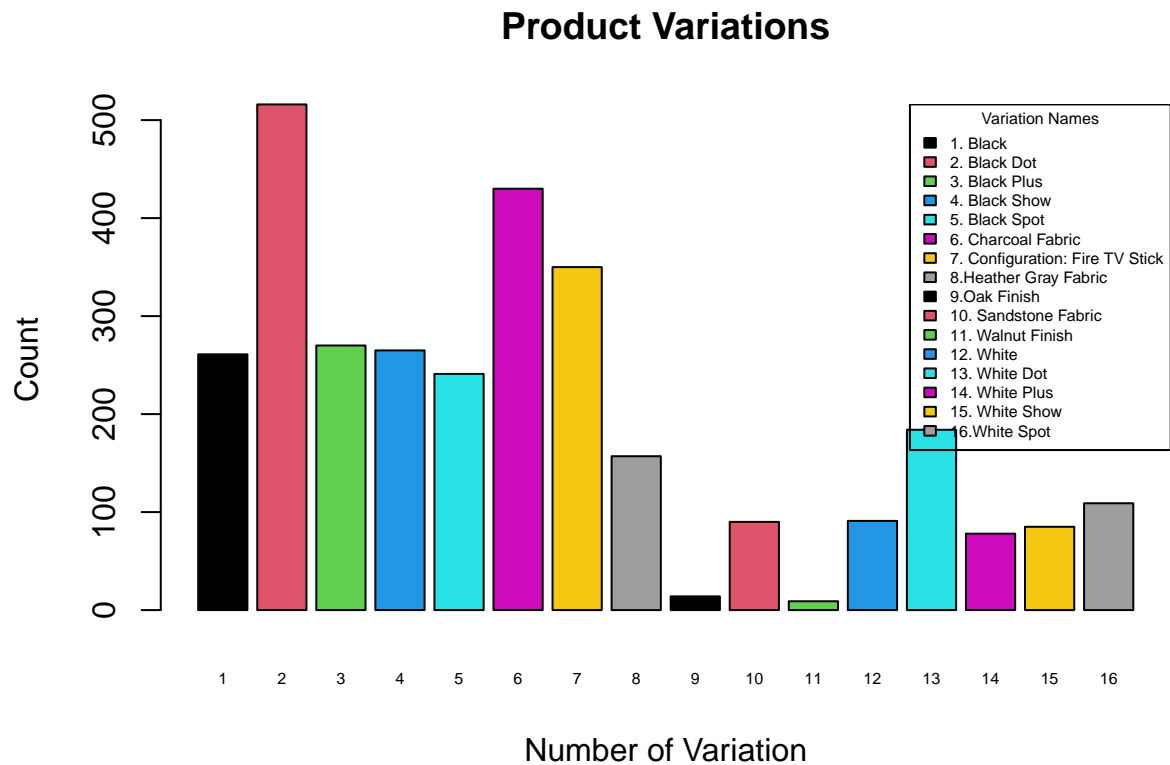
```
Vnames <- c(
  "1. Black", "2. Black Dot", "3. Black Plus", "4. Black Show",
  "5. Black Spot", "6. Charcoal Fabric", "7. Configuration: Fire TV Stick",
  "8.Heather Gray Fabric", "9.Oak Finish", "10. Sandstone Fabric",
  "11. Walnut Finish", "12. White", "13. White Dot", "14. White Plus", "15. White Show", "16.White Spot"
)
```

```
# Barplot
```

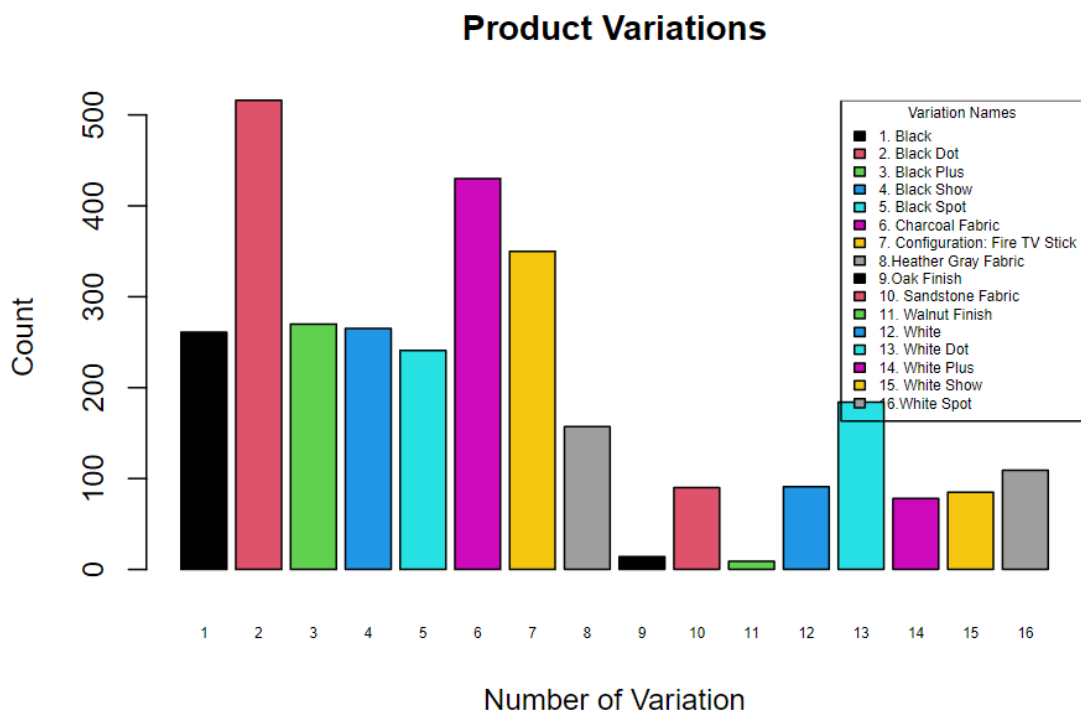
```
CompletePlot <- barplot(variations$n,
  names.arg = 1:16,
  col = 1:16,
  main = "Product Variations",
  xlab = "Number of Variation",
  ylab = "Count",
  las = 0.0,
  cex.names = 0.5,
  space = 0.2
)
```

```
# Add legend
```

```
legend("topright", legend = Vnames, fill = 1:16, title = "Variation Names", cex = 0.5)
```



```
knitr::include_graphics("/cloud/project/Worksheet#4/FINALVARIATIONDATA.png")
```



7 D. Create a `barplot()` for the black and white variations. Plot it in 1 frame, side by side. Complete the details of the chart.

```

```r
Load the required libraries
library(ggplot2)
library(magrittr)

Assuming your data frame is named variations
Replace "alexa_file$variation" with the actual column name in your data frame

Subset the data for Black variations
Blackplot <- variations[variations$`alexa_file$variation` %in% c("Black", "BlackDot", "BlackShow", "BlackShowDot"),]

Subset the data for White variations
Whiteplot <- variations[variations$`alexa_file$variation` %in% c("White", "WhiteDot", "WhiteShow", "WhiteShowDot"),]

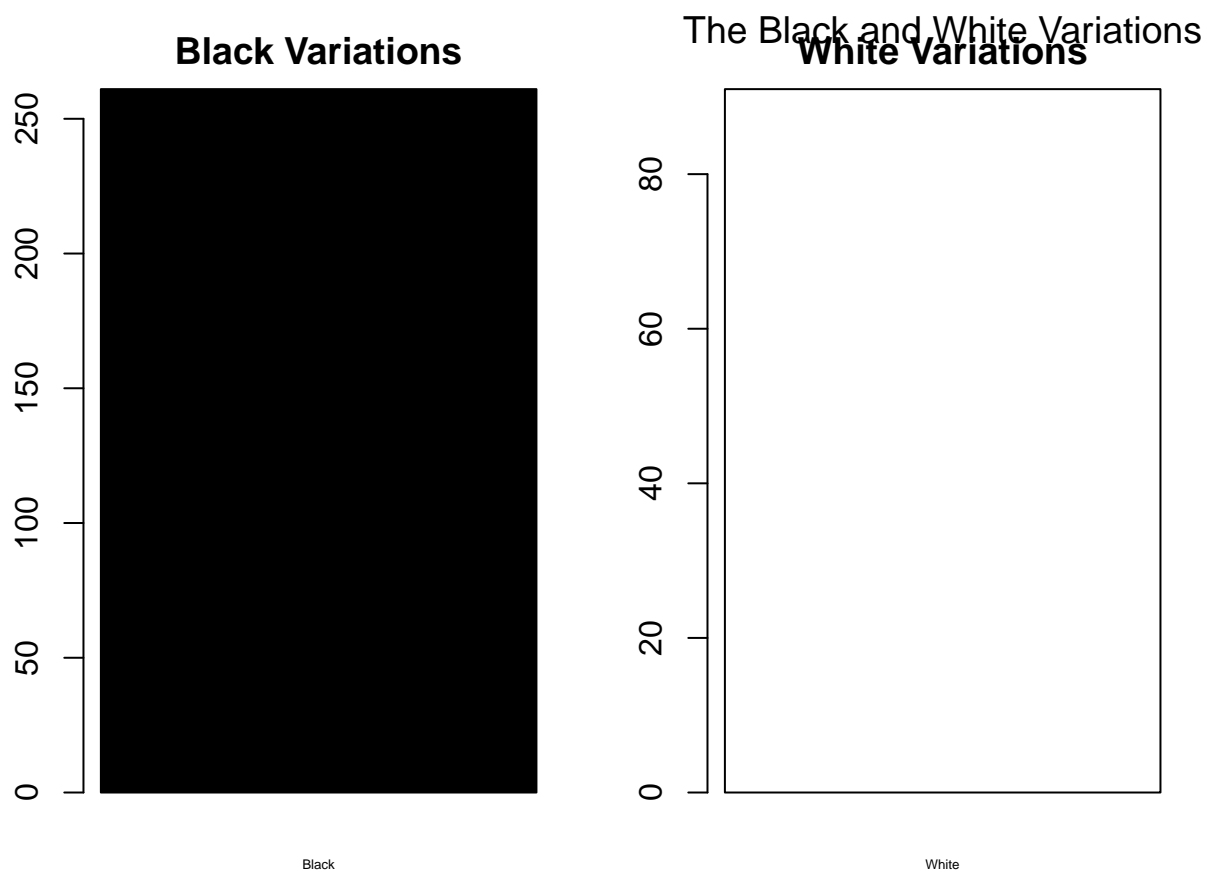
#layout one frame
par(mfrow = c(1, 2), mar = c(2, 2, 2, 2))

Barplot for Black variations
Black <- barplot(height = Blackplot$n,
 names.arg = Blackplot$`alexa_file$variation`,
 col = "Black",
 main = "Black Variations",
 xlab = "Number of Variation",
 ylab = "Count",
 las = 0.0,
 cex.names = 0.4,
 space = 0.2
)

Barplot for White variations
White <- barplot(height = Whiteplot$n,
 names.arg = Whiteplot$`alexa_file$variation`,
 col = "White",
 main = "White Variations",
 xlab = "Number of Variation",
 ylab = "Count",
 las = 0.0,
 cex.names = 0.4,
 space = 0.2
)

mtext("The Black and White Variations", side = 3, line = 1, cex = 1.2)

```



```
knitr::include_graphics("/cloud/project/Worksheet#4/BLACKandWHITE.png")
```

