

# RWorksheet\_asenjo#4b

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1.

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

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

for (i in 1:5) {
  for (j in 1:5) {
    matrixA[i, j] <- abs(i - vectorA[j])
  }
}

print(matrixA)
```

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

2.

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

for(i in v){
  cat(rep("*", i), "\n")
}
```

```
## *
## * *
## * * *
## * * * *
## * * * * *
```

3.

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

## Enter the starting number for the Fibonacci sequence:

a <- 0
b <- 1
if (!is.na(start_num) < 0) {
  1
```

```

cat("Please enter a non-negative starting number.\n")
} else {
repeat {
if (!is.na(start_num) && a >= start_num) {
cat(a, "\n")
}
temp <- a + b
a <- b
b <- temp
if (!is.na(start_num) && a > 500) {
break
}
}
}
}

```

```
## Please enter a non-negative starting number.
```

4. a.

```

Shoesizes <- read.csv("/cloud/project/Worksheet 4/shoesizes.csv")
head(Shoesizes)

```

```

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

```

b.

```

male_data <- subset(Shoesizes, Gender == "M")
female_data <- subset(Shoesizes, Gender == "F")

num_males <- nrow(male_data)
num_females <- nrow(female_data)

cat("Number of observations for Male: ", num_males, "\n")

```

```
## Number of observations for Male: 14
```

```
cat("Number of observations for Female: ", num_females, "\n")
```

```
## Number of observations for Female: 14
```

c.

```

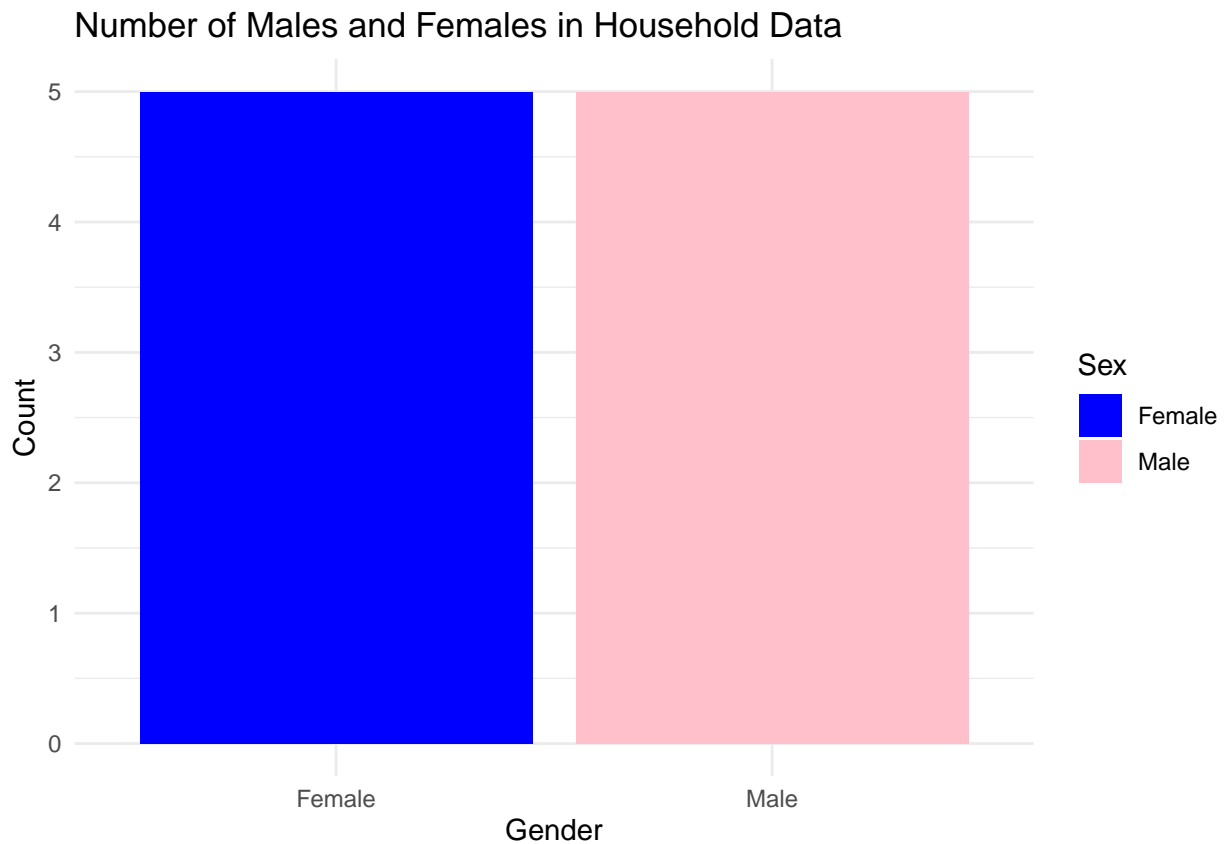
library(ggplot2)

HouseHoldData <- read.csv("/cloud/project/Worksheet 4/HouseholdData.csv")

ggplot(HouseHoldData, aes(x=Sex, fill=Sex)) +
  geom_bar() +
  ggtitle("Number of Males and Females in Household Data") +
  xlab("Gender") +

```

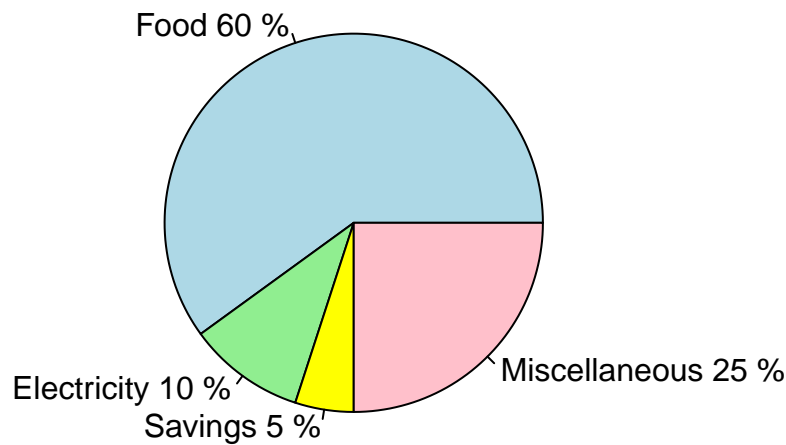
```
ylab("Count") +
scale_fill_manual(values = c("blue", "pink")) +
theme_minimal()
```



5.

```
expenses <- c(Food = 60, Electricity = 10, Savings = 5, Miscellaneous = 25)
percentages <- round(100 * expenses / sum(expenses), 1)
labels <- paste(names(expenses), percentages, "%")
colors <- c("lightblue", "lightgreen", "yellow", "pink")
pie(expenses, labels = labels, col = colors, main = "Dela Cruz Family Monthly Expenses")
```

## Dela Cruz Family Monthly Expenses



6.

```
data(iris)
```

a. The output shows number of variables and objects, and rows and columns

```
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 ...
```

b.

```
means <- colMeans(iris[, c("Sepal.Length", "Sepal.Width", "Petal.Length", "Petal.Width")])
```

```
means
```

```
## Sepal.Length Sepal.Width Petal.Length Petal.Width
##      5.843333      3.057333      3.758000      1.199333
```

c.

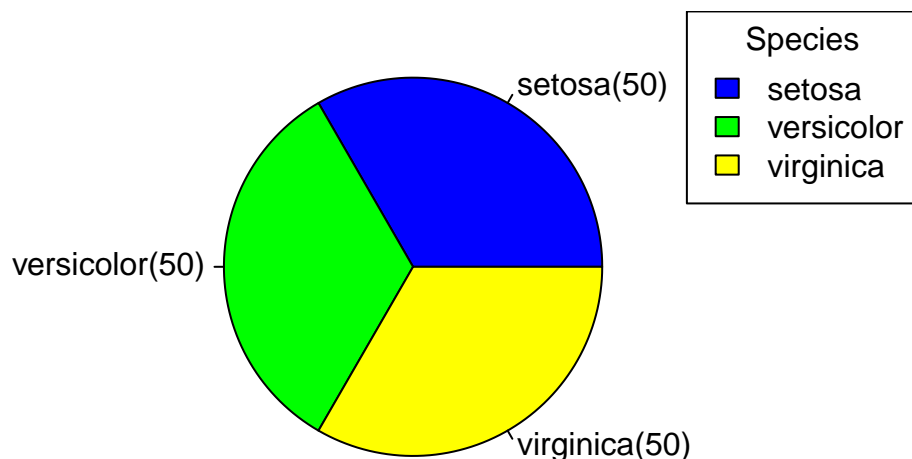
```
species_distribution <- table(iris$Species)
```

```
colors <- c("blue", "green", "yellow")
```

```
pie(species_distribution,
     main = "Distribution of Iris Species",
     col = colors,
     labels = paste(names(species_distribution), "(", species_distribution, ")", sep=""))
```

```
legend("topright", legend = names(species_distribution), fill = colors, title = "Species")
```

## Distribution of Iris Species



d.

```
setosa <- subset(iris, Species == "setosa")
versicolor <- subset(iris, Species == "versicolor")
virginica <- subset(iris, Species == "virginica")
```

```
tail(setosa)
```

##	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)
```

##	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)
```

##	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

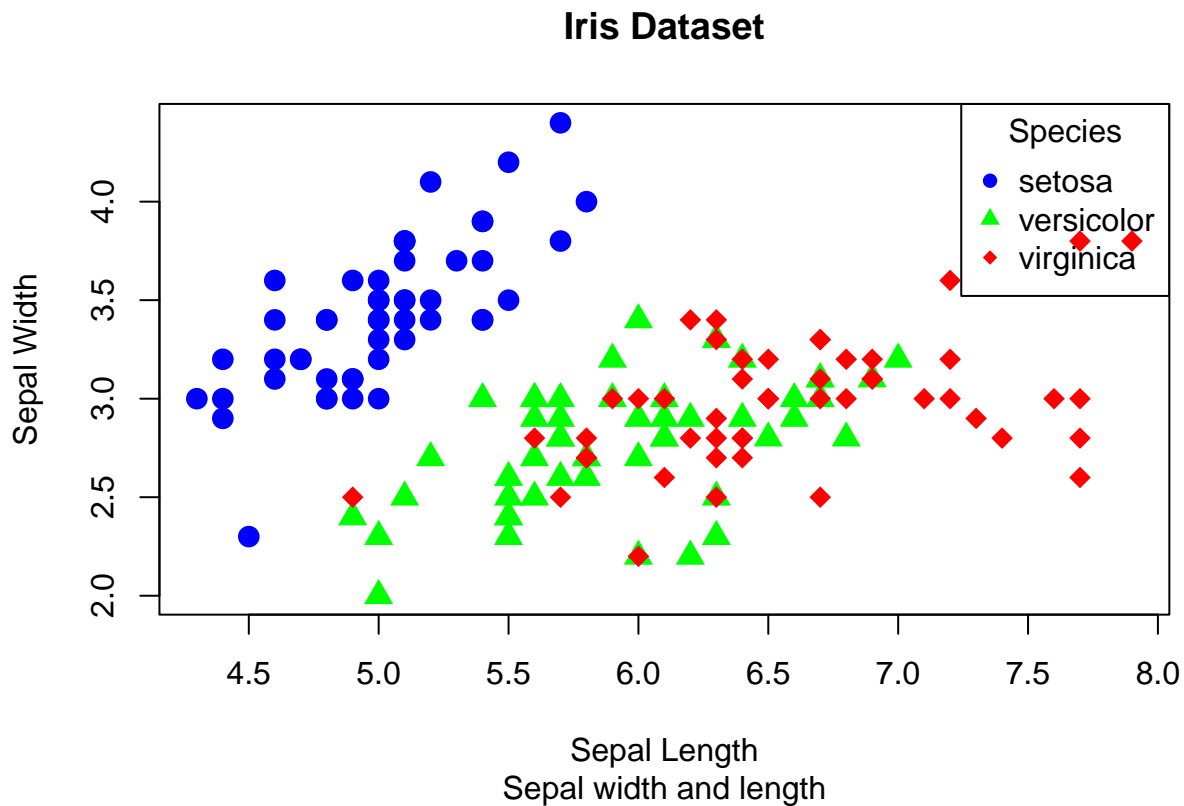
e.

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

colors <- c("setosa" = "blue", "versicolor" = "green", "virginica" = "red")
symbols <- c("setosa" = 16, "versicolor" = 17, "virginica" = 18)

plot(iris$Sepal.Length, iris$Sepal.Width,
     col = colors[iris$Species],
     pch = symbols[iris$Species],
     xlab = "Sepal Length",
     ylab = "Sepal Width",
     main = "Iris Dataset",
     sub = "Sepal width and length",
     cex = 1.5)

legend("topright", legend = levels(iris$Species),
     col = c("blue", "green", "red"),
     pch = c(16, 17, 18),
     title = "Species")
```



f. The scatterplot shows clear separation between setosa and the other two species based on Sepal Length and Sepal Width. Setosa has distinctively shorter and wider sepals, forming a separate cluster. Versicolor and virginica overlap more, particularly in sepal width, but virginica tends to have longer sepals. Overall, there is a slight negative correlation, where longer sepals tend to be narrower, especially in virginica.

8.

a.

```
library(readxl)
```

```

alexla <- read_excel("/cloud/project/Worksheet 4/alexla_file.xlsx")

alexla$variation <- gsub("Black Show", "BlackShow", alexla$variation)

alexla$variation <- gsub("Black Dot", "BlackDot", alexla$variation)

alexla$variation <- gsub("Black Plus", "BlackPlus", alexla$variation)

alexla$variation <- gsub("Black Spot", "BlackSpot", alexla$variation)

alexla$variation <- gsub("White Show", "WhiteShow", alexla$variation)

alexla$variation <- gsub("White Dot", "WhiteDot", alexla$variation)

alexla$variation <- gsub("White Plus", "WhitePlus", alexla$variation)

alexla$variation <- gsub("White Spot", "WhiteSpot", alexla$variation)

knitr::include_graphics("/cloud/project/Worksheet 4/Screenshot 2024-11-03 231530.png")

```

```

R 4.4.1 . /cloud/project/Worksheet 4/ ↗
[1] "BlackShow" "WhiteShow" "WhiteShow" "BlackShow" "BlackShow" "BlackShow" "BlackShow" "BlackShow"
[9] "BlackShow" "BlackShow" "BlackShow" "BlackPlus" "BlackPlus" "WhitePlus" "BlackPlus" "BlackPlus"
[17] "WhitePlus" "WhitePlus" "BlackPlus" "BlackPlus" "WhitePlus" "WhitePlus" "BlackPlus" "BlackPlus"
[25] "BlackPlus" "BlackPlus" "WhitePlus" "BlackPlus" "BlackPlus" "BlackPlus" "BlackPlus" "BlackPlus"
[33] "WhitePlus" "BlackPlus" "BlackPlus" "BlackPlus" "BlackPlus" "WhitePlus" "BlackPlus" "BlackPlus"
[41] "BlackPlus" "BlackPlus" "BlackPlus" "BlackPlus" "BlackPlus" "BlackPlus" "BlackPlus" "BlackPlus"
[49] "BlackPlus" "BlackPlus" "BlackPlus" "BlackPlus" "BlackPlus" "BlackPlus" "BlackPlus" "BlackPlus"
[57] "BlackPlus" "BlackPlus" "BlackPlus" "BlackPlus" "BlackPlus" "BlackPlus" "BlackPlus" "WhitePlus"
[65] "BlackPlus" "BlackPlus" "WhitePlus" "BlackPlus" "BlackPlus" "WhitePlus" "BlackPlus" "WhitePlus"
[73] "WhitePlus" "BlackPlus" "BlackPlus" "BlackPlus" "BlackPlus" "WhitePlus" "BlackPlus" "BlackPlus"
[81] "BlackPlus" "BlackPlus" "BlackPlus" "BlackPlus" "BlackPlus" "BlackPlus" "BlackPlus" "WhitePlus"
[89] "BlackPlus" "BlackPlus" "BlackPlus" "BlackPlus" "BlackPlus" "BlackPlus" "BlackPlus" "BlackPlus"
[97] "BlackPlus" "BlackPlus" "BlackPlus" "BlackPlus" "BlackPlus" "BlackPlus" "BlackPlus" "BlackPlus"
[105] "BlackPlus" "BlackPlus" "BlackPlus" "BlackPlus" "BlackPlus" "BlackPlus" "BlackPlus" "BlackPlus"
[113] "BlackPlus" "BlackPlus" "WhitePlus" "BlackPlus" "BlackPlus" "BlackPlus" "BlackPlus" "BlackPlus"

```

b.

```

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

variations.RData <- alexla %>%
  count (alexla$variation)

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

```

```
print(variations.RData)
```

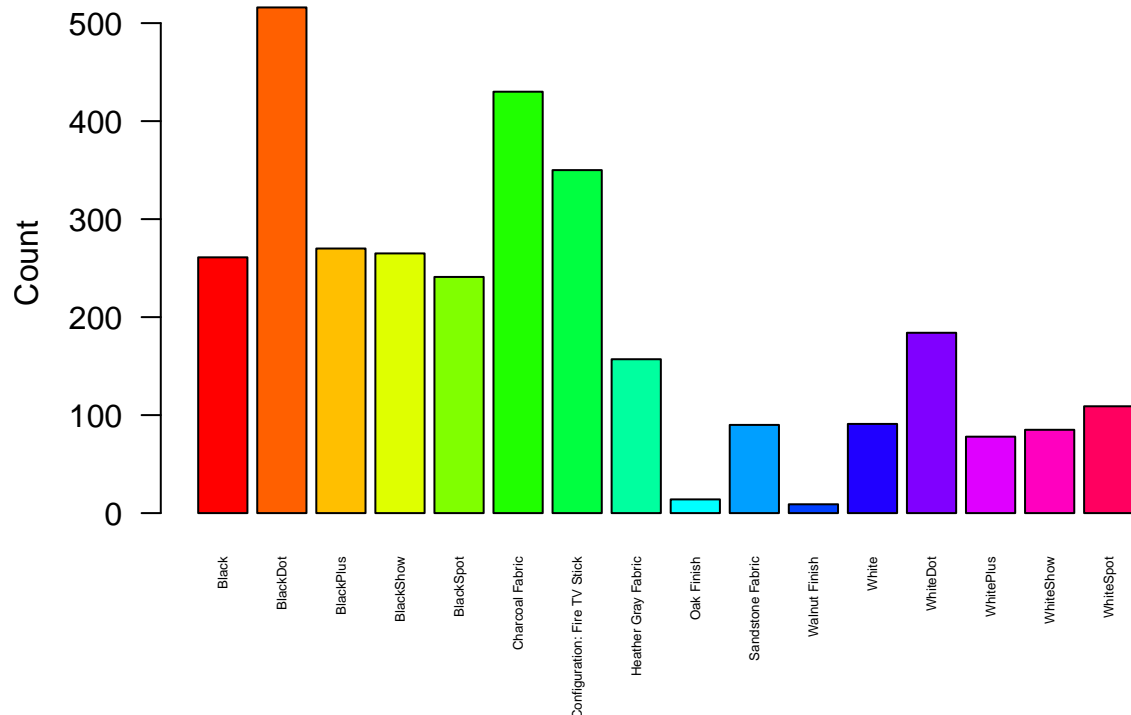
```
## # A tibble: 16 x 2
##   `alexa$variation`      n
##   <chr>                <int>
## 1 Black                261
## 2 BlackDot             516
## 3 BlackPlus            270
## 4 BlackShow            265
## 5 BlackSpot            241
## 6 Charcoal Fabric      430
## 7 Configuration: Fire TV Stick 350
## 8 Heather Gray Fabric  157
## 9 Oak Finish            14
## 10 Sandstone Fabric     90
## 11 Walnut Finish         9
## 12 White                91
## 13 WhiteDot             184
## 14 WhitePlus            78
## 15 WhiteShow            85
## 16 WhiteSpot           109
```

c.

```
barplot(
  variations.RData$n,
  names.arg = variations.RData$`alexa$variation`,
  cex.names = 0.4,
  main = "Count of Variations",
  ylab = "Count",
  col = rainbow(length(variations.RData$n)),
  border = "black",
  las = 2
)
```



## Count of Variations



d.

```

bv <- variations.RData %>%
  filter(grepl("^Black|^White", `alexa$variation`))

par(mfrow = c(1, 2))

barplot(
  bv$`bv$`alexa$variation` %in% c("Black", "BlackPlus", "BlackShow", "BlackSpot", "BlackDot")],
  names.arg = bv$`alexa$variation`[bv$`alexa$variation` %in% c("Black", "BlackPlus", "BlackShow", "BlackSpot", "BlackDot")],
  las = 3,
  cex.names = 1,
  main = "Black Variations",
  ylab = "Count",
  col = "black",
  border = "black"
)

barplot(
  bv$`bv$`alexa$variation` %in% c("White", "WhitePlus", "WhiteShow", "WhiteSpot", "WhiteDot")],
  names.arg = bv$`alexa$variation`[bv$`alexa$variation` %in% c("White", "WhitePlus", "WhiteShow", "WhiteSpot", "WhiteDot")],
  las = 2,
  cex.names = 1,
  main = "White Variations",
  ylab = "Count",
  col = "white",
  border = "black"
)

```

