

RWorksheet_Celestra#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-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.

```
# Number of rows
num_rows <- 5

for (i in 1:num_rows)
  cat(strrep(' *', i), "\n")
```

```
##  "*"
##  "*" "*"
##  "*" "*" "*"
##  "*" "*" "*" "*"
##  "*" "*" "*" "*" "*"
```

3.

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

start <- 1
a <- start
b <- 1

repeat {
  if (a >= start) {
```

```

    cat(a, " ")
  }

  next_number <- a + b
  a <- b
  b <- next_number

  if (a > 500) {
    break
  }
}

```

```
## 1  1  2  3  5  8 13 21 34 55 89 144 233 377
```

4. a

```

data <- read.csv("Shoe_sizes.csv")
head(data)

```

```

##   Show.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

```

maledata <- subset(data, Gender == "M")
femaledata <- subset(data, Gender == "F")

num_male <- nrow(maledata)
num_female <- nrow(femaledata)

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

```

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

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

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

c

```

# Sample data for Household Data with counts of males and females
gender_counts <- table(data$Gender)

```

```

barplot(
  gender_counts,
  main = "Number of Males and Females",
  col = c("blue", "pink"), # Colors for bars
  names.arg = c("Male", "Female"),
  xlab = "Gender",
  ylab = "Count",
)
legend(
  "topright",

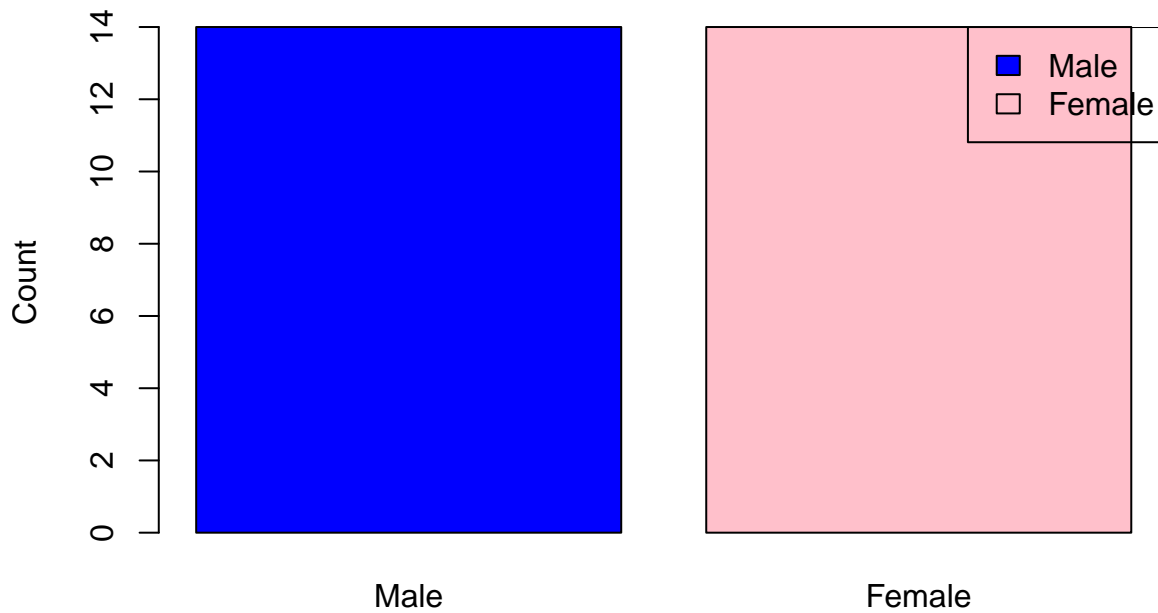
```

```

legend = c("Male", "Female"),
fill = c("blue", "pink")
)

```

Number of Males and Females



Gender

5. a

```

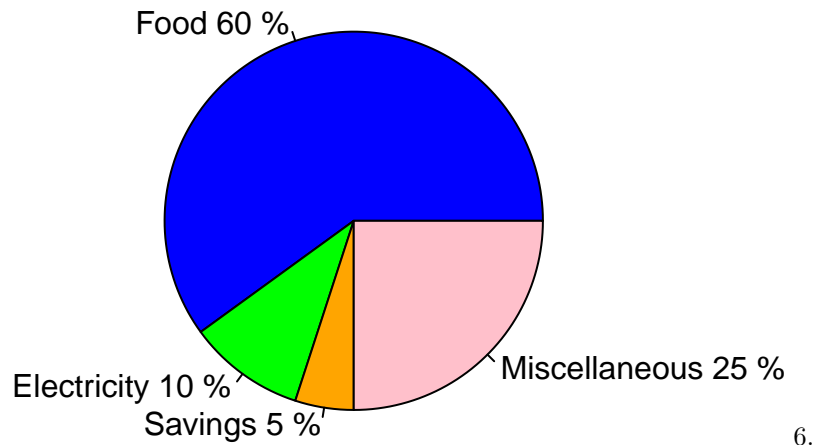
expenses <- c(Food = 60, Electricity = 10, Savings = 5, Miscellaneous = 25)

percent_labels <- paste(names(expenses), round(expenses / sum(expenses) * 100, 1), "%")

pie(
  expenses,
  labels = percent_labels,
  col = c("blue", "green", "orange", "pink"),
  main = "Dela Cruz Family Monthly Expenses"
)

```

Dela Cruz Family Monthly Expenses



6.

```
data(iris)
```

a

```
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 ...
```

The Sepal.Length representing the length of the sepal (in centimeters). Sepal.Width representing the width of the sepal. Petal.Length representing the length of the petal. Petal.Width representing the width of the petal. Species A factor variable with 3 levels: “setosa”, “versicolor”, and “virginica”, representing three species of iris flowers.

b

```
mean_values <- colMeans(iris[, 1:4])
```

```
mean_values
```

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

c

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

```
percent_labels <- paste(names(species_counts), round(species_counts / sum(species_counts) * 100, 1), "%")
```

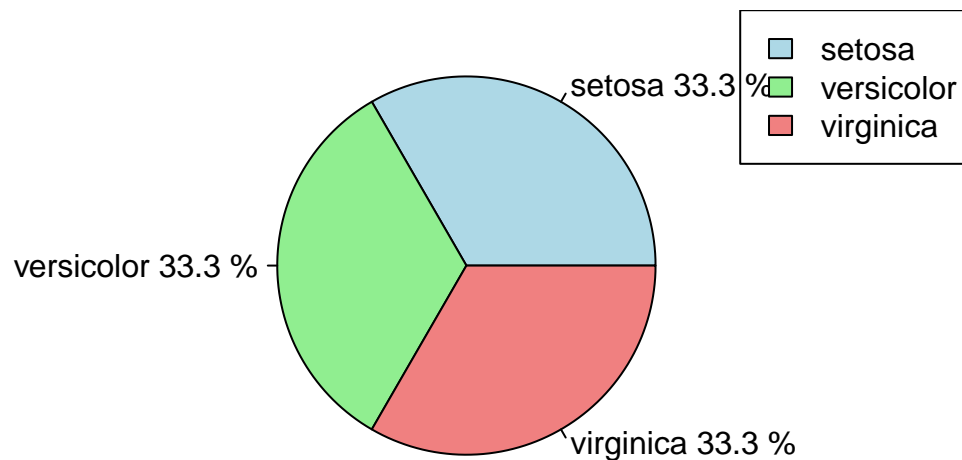
```
pie(
  species_counts,
  labels = percent_labels,
  col = c("lightblue", "lightgreen", "lightcoral"), # Colors for each species
  main = "Species Distribution in the Iris Dataset" # Title
)
```

```

legend(
  "topright",
  legend = names(species_counts),
  fill = c("lightblue", "lightgreen", "lightcoral")
)

```

Species Distribution in the Iris Dataset



d

```

setosa_subset <- subset(iris, Species == "setosa")
versicolor_subset <- subset(iris, Species == "versicolor")
virginica_subset <- subset(iris, Species == "virginica")

print(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

```

```
print(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

```

```
print(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

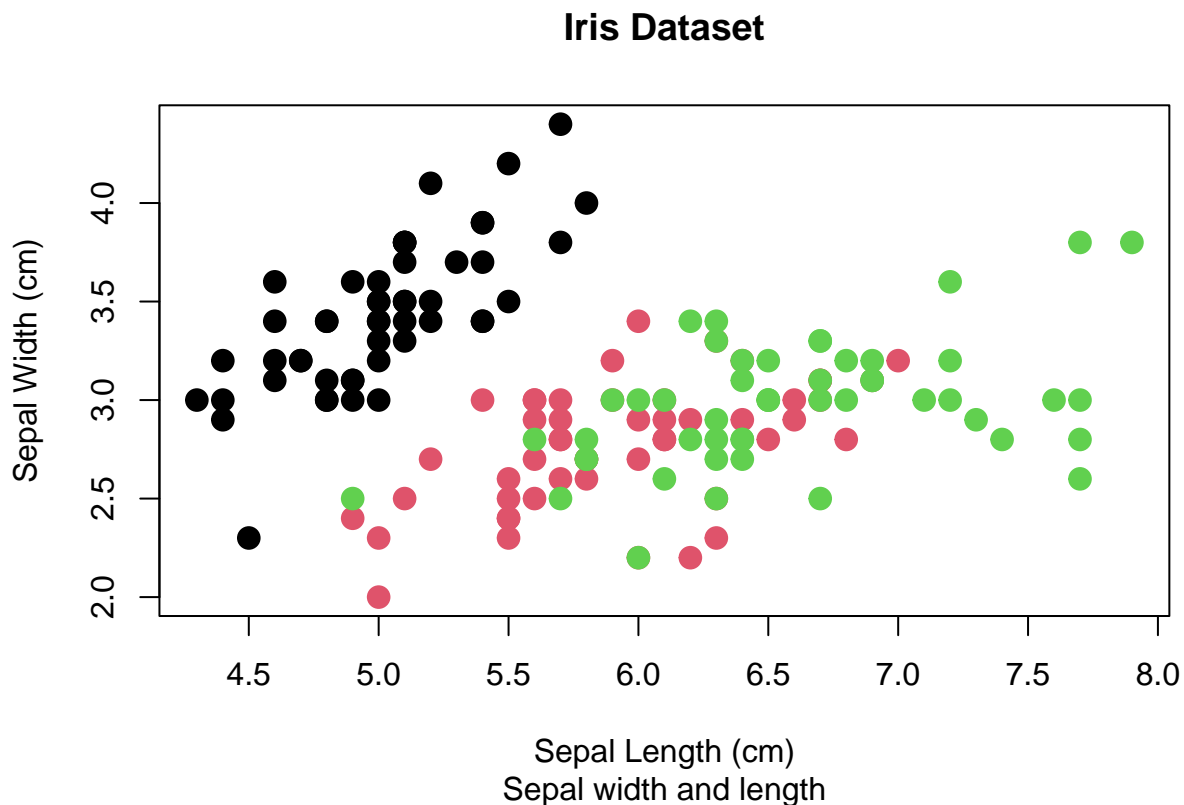
```

```
## 149      6.2      3.4      5.4      2.3 virginica
## 150      5.9      3.0      5.1      1.8 virginica
```

e

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

```
plot(
  iris$Sepal.Length,
  iris$Sepal.Width,
  main = "Iris Dataset",
  sub = "Sepal width and length",
  xlab = "Sepal Length (cm)",
  ylab = "Sepal Width (cm)",
  pch = 21,
  col = iris$Species,
  bg = iris$Species,
  cex = 1.5
)
```



f The scatterplot depicts the relationship between Sepal.Length and Sepal.Width for three iris species: setosa, versicolor, and virginica. Setosa is clearly clustered in the lower left quadrant, with smaller dimensions, distinguishing it from the other species. Versicolor exhibits a broader range of sizes, overlapping with virginica, which generally has larger measurements. 7. a

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

alexa_data$variation <- gsub("Black Dot", "BlackDot", alexa_data$variation)
alexa_data$variation <- gsub("Black Plus", "BlackPlus", alexa_data$variation)
```

```

alexa_data$variation <- gsub("Black Show", "BlackShow", alexa_data$variation)
alexa_data$variation <- gsub("Black Spot", "BlackSpot", alexa_data$variation)

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

knitr::include_graphics("/cloud/project/Worksheet4/Screenshot 2024-11-02 153147.png")

```

R 4.4.1 · /cloud/project/Worksheet4/ ↻

```

[673] "BlackShow" "WhiteShow" "WhiteShow" "BlackShow"
[677] "WhiteShow" "BlackShow" "WhiteShow" "BlackShow"
[681] "BlackShow" "BlackShow" "BlackShow" "BlackShow"
[685] "BlackShow" "WhiteShow" "BlackShow" "BlackShow"
[689] "BlackShow" "WhiteShow" "WhiteShow" "BlackShow"
[693] "BlackShow" "BlackShow" "BlackShow" "BlackShow"
[697] "BlackShow" "BlackShow" "BlackShow" "BlackPlus"
[701] "BlackPlus" "WhitePlus" "BlackPlus" "BlackPlus"
[705] "WhitePlus" "WhitePlus" "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 <- alexa_data %>%
  count(alexa_data$variation)

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

print(variations.RData)

```

```

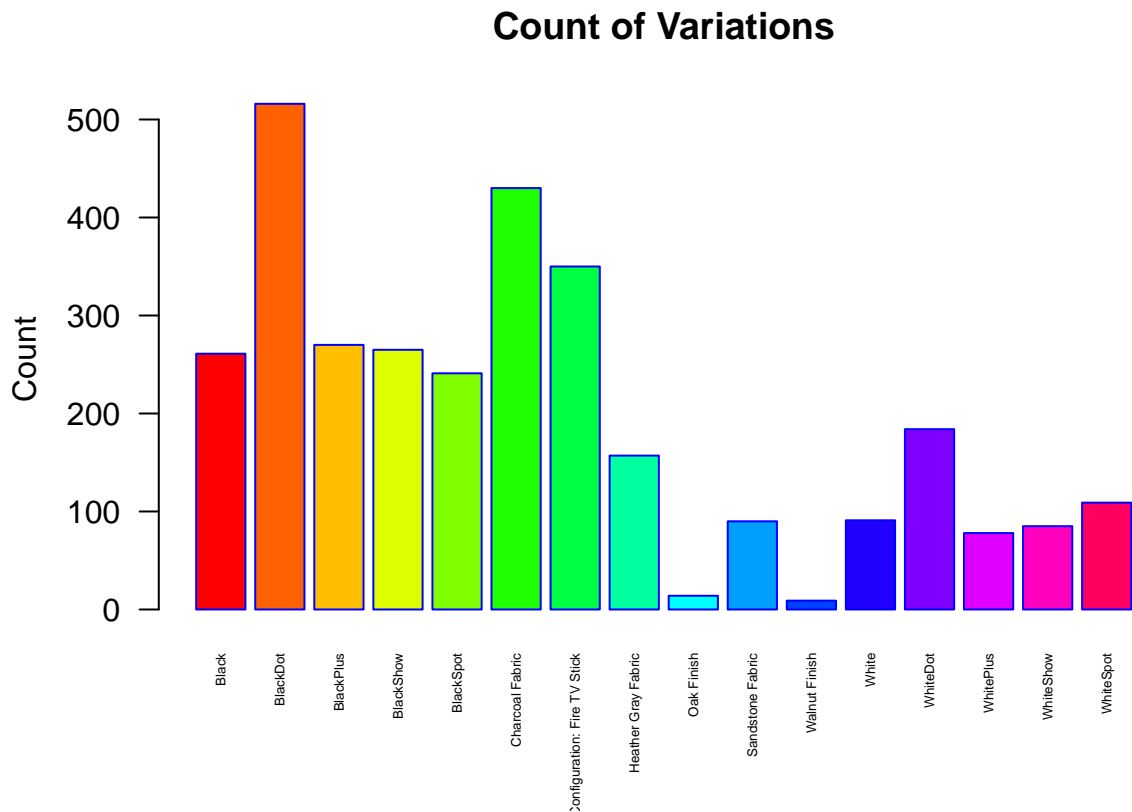
## # A tibble: 16 x 2
##   `alexa_data$variation`      n
##   <chr>                  <int>
## 1 Black                  261
## 2 BlackDot               516
## 3 BlackPlus              270
## 4 BlackShow              265
## 5 BlackSpot              241
## 6 Charcoal Fabric        430

```

```
## 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_data$variation`,
  las = 2,
  cex.names = 0.4,
  main = "Count of Variations",
  ylab = "Count",
  col = rainbow(length(variations.RData$n)),
  border = "blue"
)
```



d

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

par(mfrow = c(1, 2))
```



```
barplot(
  bw_variations$bw_variations$`alexa_data$variation` %in% c("Black", "BlackPlus", "BlackShow", "BlackSpot"),
  names.arg = bw_variations$`alexa_data$variation`[bw_variations$`alexa_data$variation` %in% c("Black", "BlackPlus", "BlackShow", "BlackSpot"),
  las = 2,
  cex.names = 1,
  main = "Black Variations",
  ylab = "Count",
  col = "black",
  border = "blue"
)
```

```
barplot(
  bw_variations$bw_variations$`alexa_data$variation` %in% c("White", "WhitePlus", "WhiteShow", "WhiteSpot"),
  names.arg = bw_variations$`alexa_data$variation`[bw_variations$`alexa_data$variation` %in% c("White", "WhitePlus", "WhiteShow", "WhiteSpot"),
  las = 2,
  cex.names = 1,
  main = "White Variations",
  ylab = "Count",
  col = "white",
  border = "blue"
)
```

