### HW4

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2024-10-13

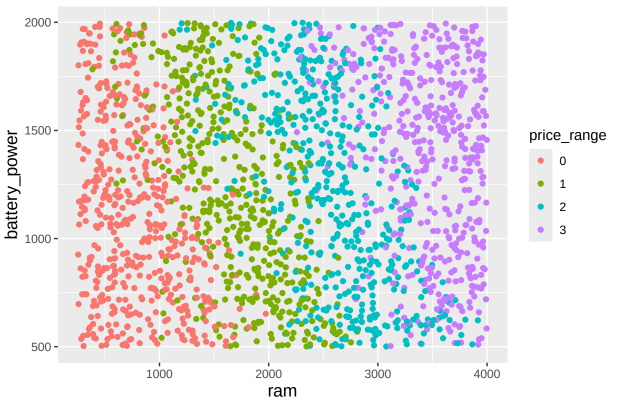
```
# Importing Necessary Libraries:
library(ggplot2)
library(tidyr)
library(UsingR)
## Loading required package: MASS
## Loading required package: HistData
## Loading required package: Hmisc
##
## Attaching package: 'Hmisc'
## The following objects are masked from 'package:base':
##
       format.pval, units
library(readr)
library(dplyr)
##
## Attaching package: 'dplyr'
## The following objects are masked from 'package:Hmisc':
##
##
       src, summarize
## The following object is masked from 'package:MASS':
##
##
       select
## The following objects are masked from 'package:stats':
##
##
       filter, lag
## The following objects are masked from 'package:base':
##
       intersect, setdiff, setequal, union
##
# Get Path to folder:
path <- "C:\\Users\\fakoy\\OneDrive - Houston Community College\\UTD_Courses\\Fall2024\\Data_Analysis_w</pre>
setwd(path)
```

#### Problem 1

(a.)

```
# Read the file:
df <- read.csv(file = "./HW4_Data/train.csv", header = T)</pre>
# 1a: scatterplot between battery_power vs ram
df$price_range <- factor(df$price_range)</pre>
base_plot <- df %>%
 ggplot(
    data = .,
    mapping =
      aes(x = ram, color = price_range)
scatter_plot <- base_plot +</pre>
  geom_point(mapping = aes(y = battery_power)) +
  ggtitle("Scatter Plot of Battery Power vs RAM") +
  theme(
    plot.title = element_text(hjust = 0.5, size = rel(1.2)),
    axis.title = element_text(size = rel(1.2)))
scatter_plot
```

# Scatter Plot of Battery Power vs RAM

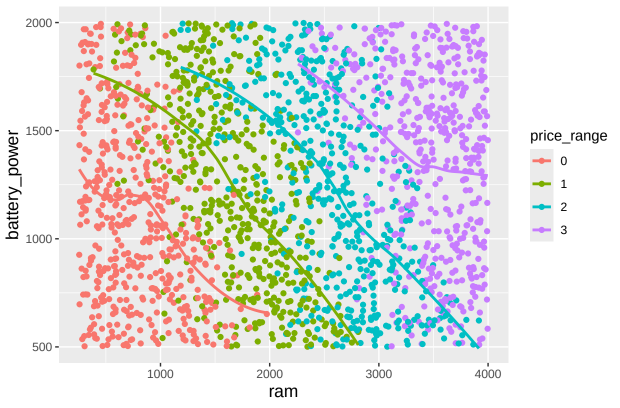


(b)

```
# 1b: Recreate plot from (a), and add trend lines for each price separately
scatter_plot_w_trend <- scatter_plot +
geom_smooth(
   mapping = aes(y = battery_power),
   method = "loess",</pre>
```

```
formula = "y ~ x",
  fill = NA
) +
coord_cartesian(
  ylim =
      c(
      round(min(df$battery_power), 2),
      round(max(df$battery_power), 2)
    )
) +
theme(
  plot.title = element_text(hjust = 0.5),
  legend.location = "panel"
) +
  ggtitle("Scatter Plot of Battery Power vs RAM based on Price Range.")
scatter_plot_w_trend
```

# Scatter Plot of Battery Power vs RAM based on Price Range.

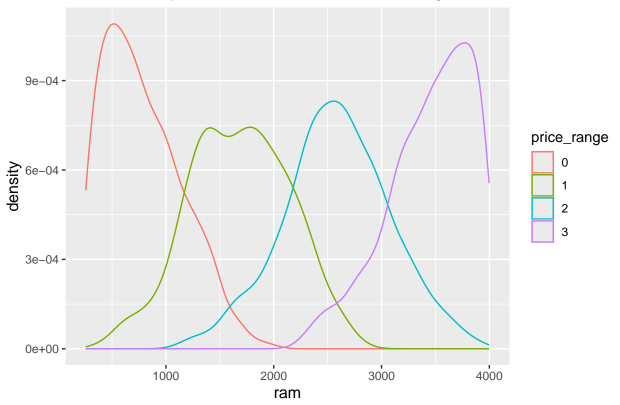


(c)

```
# 1c: density curves in one plot
density_curve <- base_plot +
  geom_density() +
  theme(
    plot.title = element_text(hjust = 0.5, size = rel(1.2)),
    axis.title = element_text(size = rel(1.1))
) +
  ggtitle("Density Plot of RAM based on Price Range")</pre>
```

#### density\_curve

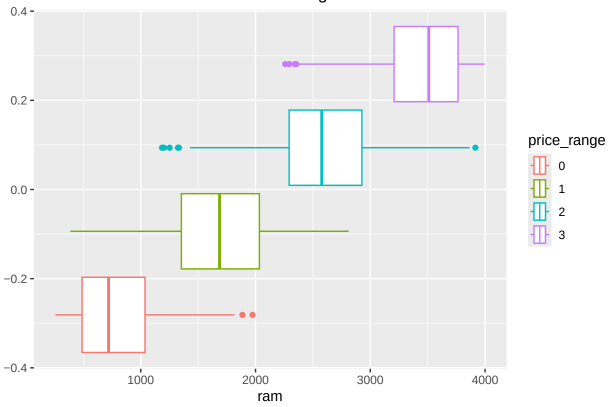
# Density Plot of RAM based on Price Range



(d)

```
# 1d: boxplots in one plot
boxplots <- base_plot +
  geom_boxplot() +
  ggtitle("Box Plot of RAM based on Price Range")
boxplots</pre>
```

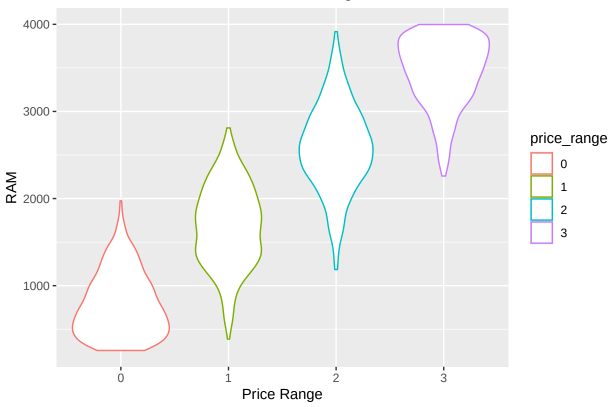
## Box Plot of RAM based on Price Range



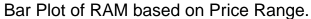
```
(e)
```

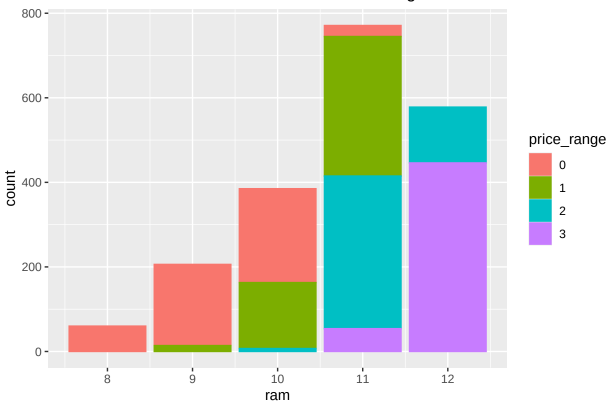
```
# 1e: violin plot
violin <- base_plot +
  geom_violin(mapping = aes(x = price_range, y = ram)) +
  xlab("Price Range") +
  ylab("RAM") +
  ggtitle("RAM vs Price Range") +
  theme(plot.title = element_text(hjust = 0.5))</pre>
```

## RAM vs Price Range



```
(f)
```





#### Problem 2

```
(a)
# Get data:
df <- UScereal
# 2a: replace levels of the factor
# variable mfr to their full names
levels(df$mfr) <- c(</pre>
  "General Mills",
 "Kellogs",
 "Nabisco",
  "Post",
  "Quaker Oats",
 "Ralston Purina")
 (b)
# 2b: turn variable shelf to a factor variable
df <- df %>%
 mutate(shelf = factor(shelf))
# 2c: Create new variable Product for the product name
rows <- rownames(df)</pre>
df <- df %>%
```

```
mutate(product = rows)
 (d)
# d: Calculate the Pearson Correlation coefficient between calories and each seven nutrition facts
pearson <- lapply(</pre>
  df[
    c(
      "protein",
      "fat",
      "sodium",
      "fibre",
      "carbo",
      "sugars",
      "potassium"
    )
  ],
  FUN = cor,
  x = df$calories,
  method = "pearson"
) %>%
  data.frame()
rownames(pearson) <- "calories"</pre>
table <- knitr::kable(pearson,</pre>
  caption = "Pearson Coefficients",
  align = "lcccc"
)
```

Table 1: Pearson Coefficients

	protein	fat	sodium	fibre	carbo	sugars	potassium
calories	0.7060105	0.5901757	0.5286552	0.3882179	0.7887227	0.4952942	0.4765955

```
(e)
# 2e: make a bar plot of the resulting correlations in (a)
# and arrange the nutrition facts in decreasing order in terms of
# their correlation with calories
sorted_pearson <- pearson %>%
    t() %>%
    data.frame() %>%
    arrange(desc(calories))

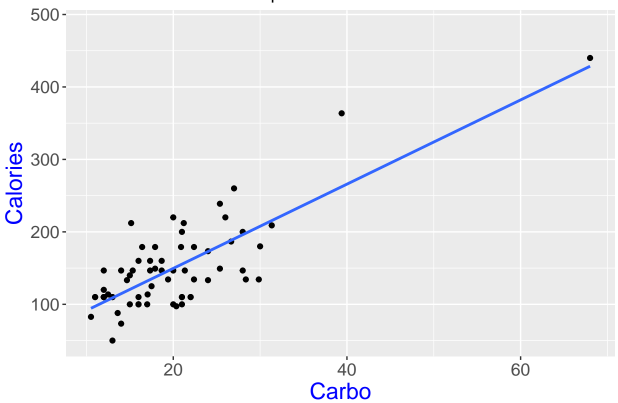
# which nutrition fact has the highest values: ans: carbo
max_pearson <- c(pearson[which(max(pearson) == pearson)])</pre>
```

The nutrition fact that has the highest values is carbo with value of 0.788722682963849

(f)

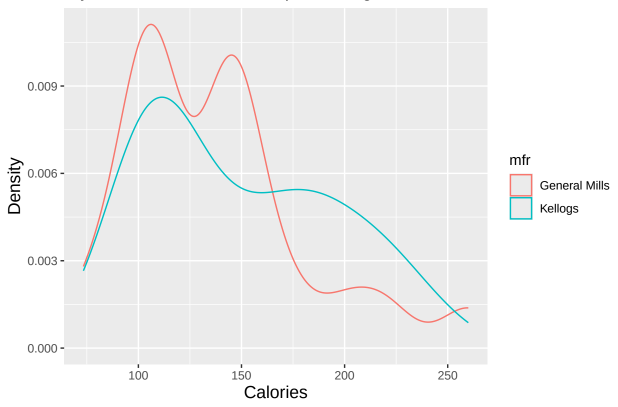
```
\# 2f: scatter plot where y represents calories and x represents
# the nutrition fact with the largest pearson correlation
# coefficient to calories
scatter_plot_w_trend <- ggplot(</pre>
  data = df,
 mapping = aes(
   x =
     df[, names(max_pearson)],
    y = calories)) +
  geom_point() +
  geom_smooth(
    method = "lm",
   formula = "y \sim x",
   fill = NA) +
  ggtitle(paste("Scatterplot of Calories vs ",
    capitalize(names(max_pearson)))) +
  theme(
   plot.title = element_text(hjust = 0.5),
   axis.title = element_text(
     colour = "BLUE",
     size = rel(1.5)),
    axis.text = element_text(size = rel(1.2))) +
    x = capitalize(names(max_pearson)),
    y = "Calories")
scatter_plot_w_trend
```

## Scatterplot of Calories vs Carbo



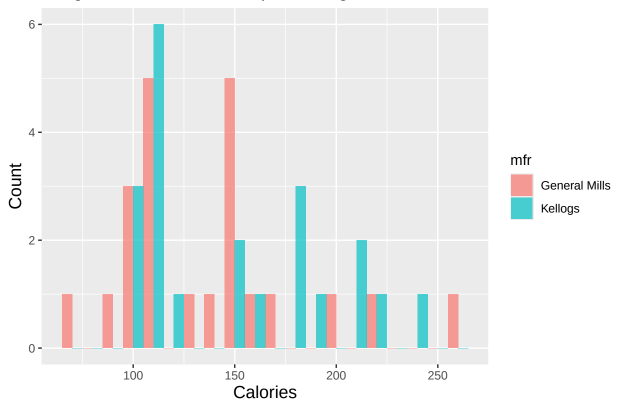
```
(g)
# 2g: Plot a density curve of Calories
density_curve <- df %>%
  filter(mfr %in% c("General Mills", "Kellogs")) %>%
  ggplot(
    data = .,
    mapping = aes(x = calories, color = mfr)
  geom_density() +
  ggtitle("Density Curve of Calories to compare Kellogs and General Mills") +
    plot.title = element_text(hjust = 0.5, size = rel(1.2)),
   axis.title = element_text(size = rel(1.2))
  ) +
  labs(
   x = "Calories",
   y = "Density"
density_curve
```

#### Density Curve of Calories to compare Kellogs and General Mills



```
(h)
# 2h: Plot Calories and Manufactures in a Histogram Plot
# to see the distribution of calories for General Mills and Kellogs:
histogram_plot <- df %>%
  filter(mfr %in% c("General Mills", "Kellogs")) %>%
  ggplot(mapping = aes(
    x = calories,
    fill = mfr)) +
  geom_histogram(
   binwidth = 10,
    position = "dodge",
    alpha = 0.7) + # Adjust binwidth as needed
  ggtitle("Histogram of Calories to Compare Kellogs and General Mills") +
  theme(
    plot.title = element_text(hjust = 0.5, size = rel(1.2)),
    axis.title = element_text(size = rel(1.2))) +
  labs(
    x = "Calories",
    y = "Count")
histogram_plot
```

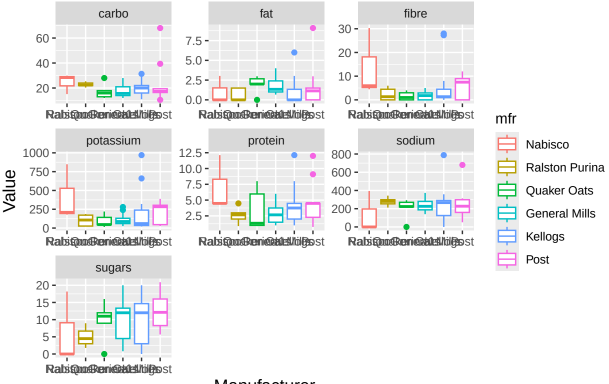
#### Histogram of Calories to Compare Kellogs and General Mills



(i) # 2i: Seven side-by-side Boxplots to compare each of the seven nutrition facts among the six mfr: long\_df <- df %>% select(mfr, calories, protein, fat, sodium, fibre, carbo, sugars, potassium) %>% pivot\_longer(cols = c(protein, fat, sodium, fibre, carbo, sugars, potassium), names\_to = "Seven\_Nutri\_Facts", values\_to = "value") # To order the boxplots according to the median value, median\_values <- long\_df %>% group\_by(mfr) %>% summarise(median\_value = median(value)) %>% arrange(median\_value) # We can re-order the mfr factor levels based on # the median values calculated long\_df\$mfr <- factor(long\_df\$mfr, levels = median\_values\$mfr)</pre> # Create the boxplot with the reordered mfr levels boxplot\_nutrition <- ggplot(long\_df,</pre> aes(x = mfr, y = value, color = mfr)) +geom\_boxplot() + facet\_wrap(~Seven\_Nutri\_Facts, scales = "free") + theme(plot.title = element\_text(hjust = 0.5, size = rel(1.2)), axis.title = element text(size = rel(1.2))) + labs(title = "Comparison of Seven Nutrition Facts Among Six MFR",

```
x = "Manufacturer",
y = "Value")
boxplot_nutrition
```

### Comparison of Seven Nutrition Facts Among Six MFR



#### Manufacturer

```
(j)
```

```
# 2j : Stacked Bar plot to show the relationship between manufacturer
# and shelf placement:
stacked_barplot <- df %>%
select(mfr, shelf) %>%
ggplot(
   data = .,
   mapping = aes(
        x = shelf,
        color = "black", fill = mfr
   )
) +
geom_bar(position = "stack") +
ggtitle("Staked Bar Plot of Shelf Placement") +
theme(plot.title = element_text(hjust = 0.5))
```

