STATS 506 HW 1

Calder Moore

STATS 506 Homework 1

Problem 1 - Abalone Data

a)

```
#a)
#Set directory and load data
setwd("C:/Users/moore/Desktop/School/Master/STATS 506/STATS-506-HW-1/abalone")
abalone <- read.csv("abalone.data")

#Rename columns
colnames(abalone)[1:9] <- c("Sex", "Length", "Diameter", "Height", "Whole weight", "Shucked to the set of the set of
```

b)

```
#b)
table(abalone$Sex)
```

```
F I M
1307 1342 1527
```

There are 1,307 females, 1,526 males, and 1,342 infants.

c) #1

[1] 0.05960039

```
#c)
#1)
#Using covariance function between Rings and each weight
cov(abalone$Rings, abalone$`Whole weight`)
[1] 0.8549952
cov(abalone$Rings, abalone$`Shucked weight`)
[1] 0.3014396
cov(abalone$Rings, abalone$`Viscera weight`)
[1] 0.1781965
cov(abalone$Rings, abalone$`Shell weight`)
[1] 0.2818386
Whole weight has the greatest correlation with Rings, with a correlation of 0.855.
c) #2
#2)
cov(abalone$Sex == "F", abalone$`Whole weight`)
[1] 0.0681564
cov(abalone$Sex == "M", abalone$`Whole weight`)
```

```
cov(abalone$Sex == "I", abalone$`Whole weight`)
```

[1] -0.1277568

Being infant has the strongest correlation with Whole weight, with a correlation of -0.128. It is the greatest correlation in terms of magnitude, the correlation is more strongly negative than the others are positive.

c) #3

```
$`Whole weight`
[1] 1.8075

$`Shucked weight`
[1] 0.7055

$`Viscera weight`
[1] 0.3215

$`Shell weight`
[1] 0.475
```

Whole weight is 1.8075, Shucked weight is 0.7055, Viscera weight is 0.3215, Shell weight is 0.475

c) #4

```
#4)
#Count the observations with a viscera weight greater than shell weight, and divide by the tweight_prop <- sum(abalone$`Viscera weight` > abalone$`Shell weight`)/nrow(abalone)
weight_prop
```

[1] 0.0651341

Only ~6.5% of the observations have a Viscera weight larger than their Shell weight.

d)

```
#d)
#Pull out the different weight values by sex
cor_F <- c(cov(abalone$Sex == "F", abalone$`Whole weight`),</pre>
           cov(abalone$Sex == "F", abalone$`Shucked weight`),
           cov(abalone$Sex == "F", abalone$`Viscera weight`),
           cov(abalone$Sex == "F", abalone$`Shell weight`))
cor_M <- c(cov(abalone$Sex == "M", abalone$`Whole weight`),</pre>
           cov(abalone$Sex == "M", abalone$`Shucked weight`),
           cov(abalone$Sex == "M", abalone$`Viscera weight`),
           cov(abalone$Sex == "M", abalone$`Shell weight`))
cor_I <- c(cov(abalone$Sex == "I", abalone$`Whole weight`),</pre>
           cov(abalone$Sex == "I", abalone$`Shucked weight`),
           cov(abalone$Sex == "I", abalone$`Viscera weight`),
           cov(abalone$Sex == "I", abalone$`Shell weight`))
#Create the table with the weights split out by sex, and rename the rows and columns for eas
sex_weight_table <- rbind(cor_F, cor_M, cor_I)</pre>
colnames(sex_weight_table) <- c("Whole weight", "Shucked weight", "Viscera weight", "Shell w
rownames(sex_weight_table) <- c("Female", "Male", "Infant")</pre>
sex_weight_table
```

```
Whole weight Shucked weight Viscera weight Shell weight Female 0.06815640 0.02716934 0.01567647 0.01977180 Male 0.05960039 0.02694935 0.01280370 0.01580163 Infant -0.12775680 -0.05411869 -0.02848017 -0.03557343
```

```
abalone_FM <- subset(abalone, abalone$Sex %in% c("F", "M"))
abalone_FI <- subset(abalone, abalone$Sex %in% c("F", "I"))
abalone_MI <- subset(abalone, abalone$Sex %in% c("M", "I"))
ttest_FM <- t.test(abalone_FM$Rings ~ abalone_FM$Sex)</pre>
ttest_FI <- t.test(abalone_FI$Rings ~ abalone_FI$Sex)</pre>
ttest_MI <- t.test(abalone_MI$Rings ~ abalone_MI$Sex)</pre>
ttest_FM
    Welch Two Sample t-test
data: abalone_FM$Rings by abalone_FM$Sex
t = 3.69, df = 2741.8, p-value = 0.0002286
alternative hypothesis: true difference in means between group F and group M is not equal to
95 percent confidence interval:
0.1999174 0.6533201
sample estimates:
mean in group F mean in group M
       11.12930
                       10.70269
ttest_FI
    Welch Two Sample t-test
data: abalone_FI$Rings by abalone_FI$Sex
t = 29.477, df = 2508.9, p-value < 2.2e-16
alternative hypothesis: true difference in means between group F and group I is not equal to
95 percent confidence interval:
3.023380 3.454304
sample estimates:
mean in group F mean in group I
```

#Create subsets of the data that only include the certain sex pairs so that we can run the t

7.890462

11.129304

```
Welch Two Sample t-test
```

The difference in rings between females and males was the smallest. Both females and males had significantly different numbers of rings from infants.

Problem 2 - Food Expenditure Data

a)

```
#Load data
setwd("C:/Users/moore/Desktop/School/Master/STATS 506/STATS-506-HW-1")
food <- read.csv("food_expenditure.csv")</pre>
```

b)

```
#Rename variables colnames(food) <- c("ID", "Age", "Household Members", "State", "Currency", "Total Expenditure
```

c)

```
#Count observations before currency restriction
nrow(food)
```

[1] 262

```
#Restrict to only USD
food_USD <- subset(food, food$Currency %in% c("USD"))

#Recheck observation count
nrow(food_USD)</pre>
```

[1] 230

Number of observations decreased by 32.

d-h)

```
#d) Remove under 18 since they probably don't have a great sense of home expenditures usually food_clean <- subset(food_USD, food_USD$Age %in% c(18:100))

#e) Remove the observations with a state of XX food_clean <- subset(food_clean, !food_clean$State %in% c("XX"))

#f) In total expenditures, there is an observation with "-350" rather than just 350. There as food_clean$`Total Expenditures` [which(food_clean$`Total Expenditures` == "-350")] <- 350 food_clean <- subset(food_clean, food_clean$`Total Expenditures` > 0) food_clean <- subset(food_clean, food_clean$`Grocery Store Expenditures` > 0) food_clean <- subset(food_clean, food_clean$`Dining Out Expenditures` > 0) food_clean <- subset(food_clean, food_clean$`Misc Expenditures` > 0)

#g) There are some entries in dine out count that seem excessive. Eating out 30 times in a way food_clean <- subset(food_clean, food_clean$`Dine Out Count` < 20)

#h) nrow(food_clean)
```

[1] 116

Only 116 observations left after cleaning compared to an original 262.

Problem 3 - Collatz Conjecture

a)

```
#' Function to find the next value in the Collatz Conjecture sequence
# '
#' Oparam integer
#' Oreturns the next number in the Collatz Conjecture sequence. Gives errors requesting numer
#' @examples an input of 5 returns 16, of 16 returns 8 according to the rule
nextCollatz <- function(integer){</pre>
  #Check for numbers
  if(is.numeric(integer)){
    #check for positive numbers
    if(integer > 0){
      #Check for evens
      if(round(integer) %% 2 == 0){
        next_int <- round(integer)/2</pre>
      }
      #Check for odds
      else if(round(integer) %% 2 != 0){
        next_int <- 3*round(integer) + 1</pre>
      }
      return(next_int)
    }
    else{
      return("Input must be positive")
    }
  }
  else{
    return("Input must be numeric")
}
#Test
nextCollatz(5)
```

[1] 16

```
nextCollatz(16)
[1] 8
b)
collatzSequence <- function(integer){</pre>
  seq <- c(integer)</pre>
  next_int <- nextCollatz(integer)</pre>
  while(next_int > 1){
    seq <- c(seq, next_int)</pre>
    next_int <- nextCollatz(next_int)</pre>
    #Making sure it includes the final "1" to end the sequence
    if(next_int == 1){
      seq <- c(seq, next_int)</pre>
      break
    }
  }
  return(seq)
#Test
collatzSequence(5)
[1] 5 16 8 4 2 1
collatzSequence(19)
 [1] 19 58 29 88 44 22 11 34 17 52 26 13 40 20 10 5 16 8 4 2 1
c)
```

```
collatzList <- c()
for(i in 100:500){
  collatzList <- c(collatzList, length(collatzSequence(i)))
}</pre>
```

```
names(collatzList) <- c(100:500)

collatzList[which(collatzList == max(collatzList))]

327
144

collatzList[which(collatzList == min(collatzList))]

128
8</pre>
```

gives the longest sequence with a length of 144, and 128 has the shortest sequence, with a length of 8.