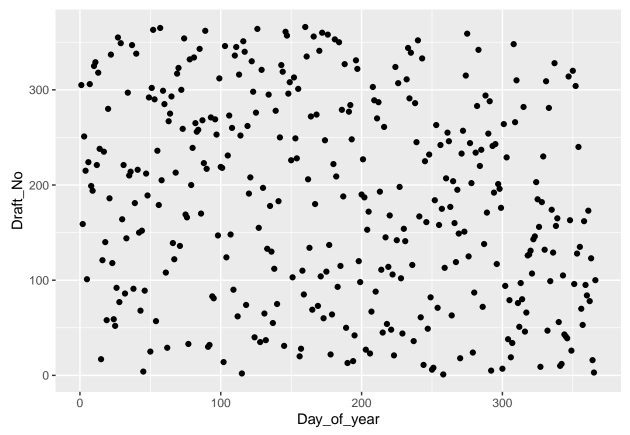
Computational Statistics Computer Lab 5 (Group 7)

Question 1: Hypothesis testing (Solved by Qinyuan Qi)

Answer:

(1.1) Make a scatter plot of Y versus X and conclude

The plot is as follows, and according to the plot, the pattern is not very clear. So it seems to be random distributed.



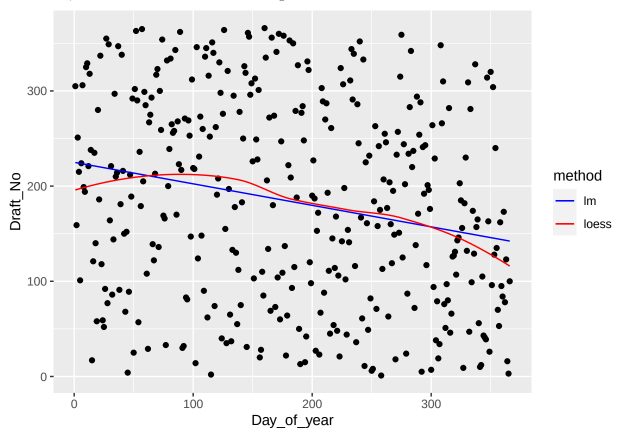
(1.2) Fit a curve to the data and conclude

Fitting the points using lm() and loess(), the plot as below.

According to the 2 curves, it seems that all the points are divided to 2 groups randomly. and we can guess that the points are randomly distributed.

According to the plot, we can not get the conclusion that which model(lm or loess) is better since only parameter (Day of year) is used. And according to the output of 2 models, output format is different, however,

lm version's residual standard error is 103.2 and loess version's residual standard error is 103. according to this value, we can know that there have no big difference between these 2 versions.



(1.3) Estimate S's distribution through a non-parametric bootstrap

The bootstrap value is generated using boot function. please check code in the appendix for detail.

We use the following formula to calc the p-value:

$$\hat{p} = \frac{\sum S \geqslant S0}{N}$$

The S value and p-value for lm and loess regression are generated by code.

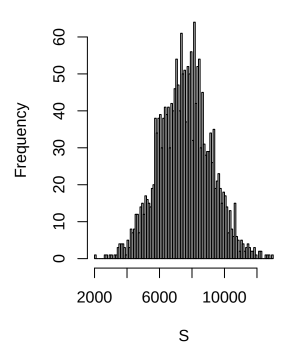
Because S value is not close to zero, then this indicates some trend in the data, we throws suspicion on the randomness of the lottery dataset.

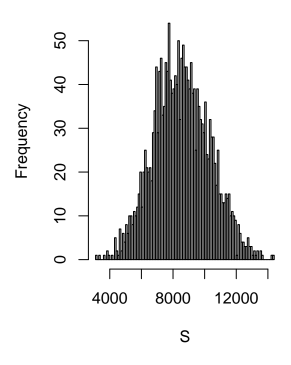
According to the plot, 2 plots are not symmetrical, also both of them not not follow the pattern of the known distributions, so we can say that it is not randomly distributed.

- ## mean of bootstrap sample statistics(lm) = 7551.223
- ## mean of bootstrap sample statistics(loess) = 8544.389
- ## p-value for linear regression = 0.492
- ## p-value for loess regression = 0.5525

Bootstrap S Distribution(Im)

Bootstrap S Distribution(loess)





(1.4)

The following code will generate the value of S and its p-value, based on 2000 bootstrap samples.

```
test_hypothesis <- function(data, B = 2000) {</pre>
  # bootstrap samples
  bootstrap_samples_lm <- boot(data = data, statistic = fun_lm_s, R = B,
                  formula=Draft_No ~ Day_of_year,parallel = "multicore")
  bootstrap_samples_loess <- boot(data = data, statistic = fun_loess_s, R = B,
                  formula=Draft_No ~ Day_of_year,parallel = "multicore")
  # p-values
  pvalue_lm = sum(boot_samples_lm$t >= boot_samples_lm$t0)/length(boot_samples_lm$t)
  pvalue_loess = sum(boot_samples_loess$t >= boot_samples_loess$t0)/length(boot_samples_loess$t)
  mean_boot_samples_lm <- mean(boot_samples_lm$t)</pre>
  mean_boot_samples_loess <- mean(boot_samples_loess$t)</pre>
  result <- list(
    s_lm_value = mean_boot_samples_lm,
    p_lm_value = pvalue_lm,
    s_loess_value = mean_boot_samples_loess,
    p_loess_value = pvalue_loess
  return(result)
}
```

(1.5)

We will generate a dataset of the same dimensions as the original data.

If we set k=9, and alpha = 0.05, compare pvalue and alpha we can accept or reject hypothesis.

According to the result, in the case of lm, we fail to reject H0 hypothesis, which is Lottery is random.

P-value(loess): 0.5525

If we set k=10 to 50, and skip all k %% 3 != 0,we got 13 Fail to Reject H0 Hypothesis, and not got any Reject case.

```
#max_k = dim(data)[1]
max_k = 50

for(k in 10:max_k){
   if (k %% 3 != 0) {
      next
   }
   data_random <- gen_df_1_5(k)

res <- test_hypothesis(data = data_random)
   # we only test p_loess_value here.
   if(result$p_loess_value <= alpha ){
      cat("Reject HO Hypothesis","\n")
   }else{
      cat("Fail to Reject HO Hypothesis","\n")
}</pre>
```

```
## Fail to Reject HO Hypothesis
```

```
## Fail to Reject HO Hypothesis
```

According to the above, we know that Lottery is random.

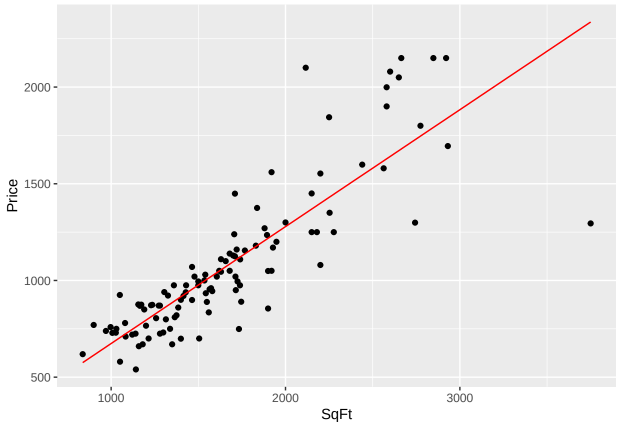
Question 2: Bootstrap, jackknife and confidence intervals (Solved by Satya Sai Naga Jaya Koushik Pilla)

Answer:

(2.1)

we create a scatter plot of SqFt versus Price, and fit a line. Price = 69.30663 + 0.60457 * SqFt

```
data <- read.csv("prices1.csv", sep = ";")</pre>
g_2_1 <- ggplot(data, aes(x = SqFt, y = Price)) + geom_point()</pre>
# fit a linear regression model
fit2_1_lm <- lm(Price ~ SqFt, data = data)</pre>
summary(fit2 1 lm)
##
## Call:
## lm(formula = Price ~ SqFt, data = data)
## Residuals:
       Min
                 1Q
                      Median
                                   3Q
                                          Max
## -1041.43
            -99.12
                        1.99
                                59.26
                                       751.43
##
## Coefficients:
              Estimate Std. Error t value Pr(>|t|)
## (Intercept) 69.30663
                         65.13461
                                  1.064
                                             0.29
               0.60457
                          0.03713 16.282
                                           <2e-16 ***
## SqFt
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
## Residual standard error: 206 on 108 degrees of freedom
## Multiple R-squared: 0.7105, Adjusted R-squared: 0.7079
## F-statistic: 265.1 on 1 and 108 DF, p-value: < 2.2e-16
minimal x <- min(data$SqFt)</pre>
maximum_x <- max(data$SqFt)</pre>
x_2_1_lm <- seq(minimal_x, maximum_x, 0.1)</pre>
y_2_1_lm <- predict(fit2_1_lm, newdata = data.frame(SqFt = x_2_1_lm))</pre>
data_2_1 \leftarrow data.frame(SqFt = x_2_1_lm, Price = y_2_1_lm)
g_2_1 + geom_line(data = data_2_1, aes(x = x_2_1_lm, y = y_2_1_lm), color = "red")
```



According to the plotted line, it does not seem like a good fit. since the left bottom of the plot is very dense, but top right corner is very sparse.

(2.2)

According to the question, we need to min RSS as follows

$$RSS = \sum (y_i - f(f(x_i)))^2$$

And the code implementation as follows.

```
rss_fun <- function(par,SqFt,Price) {
    a1 <- par[1]
    a2 <- par[2]
    b <- par[3]
    c <- par[4]

    predicted <- ifelse(SqFt > c, b + a1 * SqFt + a2 * (SqFt - c), b + a1 * SqFt)

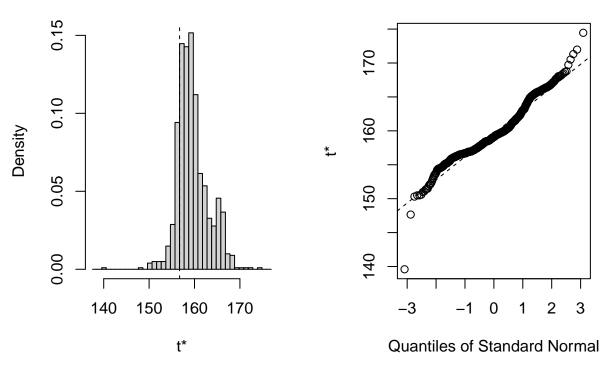
    rss <- sum((Price - predicted)^2)
    return(rss)
}

c_func <- function(c, SqFt, Price) {
    # Init
    init_pars <- c(2, 0.1, 50,c)</pre>
```

```
# Optimize using optim()
  result <- optim(par = init_pars, rss_fun, SqFt = SqFt, Price = Price)
  opt_pars <- result$par</pre>
  rss <- result$value
 return(list(c = opt_pars, params = opt_pars, rss = rss))
c <- 150
results <- c_func(c, SqFt = data$SqFt, Price = data$Price)</pre>
results
## $c
## [1]
        0.68946673 -0.08460581 55.69699121 156.71143820
##
## $params
        0.68946673 -0.08460581 55.69699121 156.71143820
## [1]
## $rss
## [1] 4584296
(2.3)
We estimate the distribution of c using bootstrap
stat1 <- function(data, vn){</pre>
    data = as.data.frame(data[vn,])
    res <- c_func(c, SqFt = data$SqFt, Price = data$Price)</pre>
    return(res$param[4])
}
set.seed(12345)
res1 = boot(data, stat1, R=1000)
res1
## ORDINARY NONPARAMETRIC BOOTSTRAP
##
##
## Call:
## boot(data = data, statistic = stat1, R = 1000)
##
## Bootstrap Statistics :
                          std. error
       original bias
## t1* 156.7114 2.875368
                             3.410341
We calc the 95% confidence interval
print(boot.ci(res1))
## BOOTSTRAP CONFIDENCE INTERVAL CALCULATIONS
## Based on 1000 bootstrap replicates
```

```
## CALL :
## boot.ci(boot.out = res1)
##
## Intervals :
## Level
              Normal
                                  Basic
## 95%
         (147.2, 160.5)
                            (146.7, 159.2)
##
             Percentile
## Level
                                    BCa
## 95%
         (154.3, 166.8)
                           (139.6, 159.0)
## Calculations and Intervals on Original Scale
## Warning : BCa Intervals used Extreme Quantiles
## Some BCa intervals may be unstable
# plot
plot(res1)
```

Histogram of t



(2.4)

Estimate the variance of c using the jackknife:

```
# Jackknife Function
jackknife <- function(data, fun) {
    n <- length(data)
    indices <- 1:n
    estimates <- numeric(n)

for (i in 1:n) {
    jackknife_sample <- data[-i]
    estimates[i] <- fun(jackknife_sample)
}</pre>
```

```
bias_correction <- (n - 1) * mean(estimates) - fun(data)

jackknife_var <- ((n - 1) / n) * sum((estimates - mean(estimates) - bias_correction)^2)

return(list(estimates = estimates, jackknife_var = jackknife_var))

# not working here,c_values

jackknife_results <- jackknife(c_values, function(c_val) {
    c_func(c_val, SqFt, Price)*params[1]
})

# Print Jackknife Results
cat("Jackknife Estimates of c:", jackknife_results*estimates, "\n")
cat("Jackknife Variance of c:", jackknife_results*jackknife_var, "\n")</pre>
```

(2.5)

N/A

Appendix: Code for this report

```
rm(list = ls())
library(ggplot2)
library(boot)
set.seed(12345)
data <- read.csv("lottery.csv", sep = ";")</pre>
# Remove Month column since there is a Mo. Number column which is same as Month
data <- data[,4:5]
g_1_1 <- ggplot(data, aes(x = Day_of_year, y =Draft_No )) + geom_point()</pre>
g_{1}1_{1}
minimal_x <- min(data$Day_of_year)</pre>
maximum_x <- max(data$Day_of_year)</pre>
x_1_2 <- seq(minimal_x, maximum_x, 0.1)</pre>
# fit the data using linear regression
fit1_2_lm <- lm(Draft_No ~ Day_of_year, data = data)
#summary(fit1 2 lm)
y_1_2_lm <- predict(fit1_2_lm, newdata = data.frame(Day_of_year = x_1_2))</pre>
# fit using loess function
fit1_2_loess <- loess( Draft_No ~ Day_of_year, data = data, se = TRUE)</pre>
#summary(fit1_2_loess)
y_1_2_loess <- predict(fit1_2_loess, newdata = data.frame(Day_of_year = x_1_2))</pre>
# plot 2 predicted lines
data_1_2 <- data.frame(Day_of_year = x_1_2, y_lm = y_1_2_lm, y_loess = y_1_2_loess)
g 1 1 + geom line(data = data 1 2, aes(x = Day of year, y = y lm,colour="lm")) +
geom_line(data = data_1_2, aes(x = Day_of_year, y = y_loess, colour = "loess")) +
scale_color_manual(name = "method", values = c("lm" = "blue", "loess" = "red"))
# boot function for lm
fun_lm_s <- function(formula,data,indices){</pre>
 d <- data[indices, ]</pre>
 lm_mod <- lm(formula, data = d)</pre>
 lm_val <- predict(lm_mod,</pre>
                 newdata = data.frame(Day_of_year = d$Day_of_year))
 s_lm <- sum(abs(lm_val - mean(d$Draft_No)))</pre>
 return(s_lm)
# boot function for loess
```

```
fun_loess_s <- function(formula,data,indices){</pre>
  d <- data[indices, ]</pre>
  loess_mod <- loess(formula, data = d, se = TRUE)</pre>
  loess_val <- predict(loess_mod,</pre>
                    newdata = data.frame(Day_of_year = d$Day_of_year))
  s_loess <- sum(abs(loess_val - mean(d$Draft_No)))</pre>
 return(s_loess)
bootstrap sample number <- 2000
# bootstrap for linear regression
boot_samples_lm <- boot(data, statistic = fun_lm_s, R = bootstrap_sample_number, formula=Draft_No ~ Da
boot_samples_loess <- boot(data, statistic = fun_loess_s, R = bootstrap_sample_number, formula=Draft_No
# calculate mean of bootstrap sample statistics
mean_boot_samples_lm <- mean(boot_samples_lm$t)</pre>
cat("mean of bootstrap sample statistics(lm) = ", mean_boot_samples_lm, "\n")
#mean of bootstrap sample statistics(lm) = 7646.339
mean_boot_samples_loess <- mean(boot_samples_loess$t)</pre>
cat("mean of bootstrap sample statistics(loess) = ", mean_boot_samples_loess, "\n")
pvalue_lm = sum(boot_samples_lm$t >= boot_samples_lm$t0)/length(boot_samples_lm$t)
pvalue_loess = sum(boot_samples_loess$t >= boot_samples_loess$t0)/length(boot_samples_loess$t)
cat("p-value for linear regression = ", pvalue_lm, "\n")
cat("p-value for loess regression = ", pvalue_loess, "\n")
# plot 2 hist using 1*2 layout
par(mfrow = c(1,2))
# Plot the histogram of bootstrap sample statistics(lm)
hist(boot_samples_lm$t, main = "Bootstrap S Distribution(lm)", xlab = "S", breaks=100)
# Plot the histogram of bootstrap sample statistics(loess)
hist(boot_samples_loess$t, main = "Bootstrap S Distribution(loess)", xlab = "S", breaks=100)
test_hypothesis <- function(data, B = 2000) {</pre>
  # bootstrap samples
  bootstrap_samples_lm <- boot(data = data, statistic = fun_lm_s, R = B,
                  formula=Draft_No ~ Day_of_year,parallel = "multicore")
  bootstrap_samples_loess <- boot(data = data, statistic = fun_loess_s, R = B,
                  formula=Draft_No ~ Day_of_year,parallel = "multicore")
  # p-values
  pvalue_lm = sum(boot_samples_lm$t >= boot_samples_lm$t0)/length(boot_samples_lm$t)
  pvalue_loess = sum(boot_samples_loess$t >= boot_samples_loess$t0)/length(boot_samples_loess$t)
  mean_boot_samples_lm <- mean(boot_samples_lm$t)</pre>
  mean_boot_samples_loess <- mean(boot_samples_loess$t)</pre>
```

```
result <- list(
   s_lm_value = mean_boot_samples_lm,
   p_lm_value = pvalue_lm,
   s loess value = mean boot samples loess,
   p_loess_value = pvalue_loess
 return(result)
}
#(a) generate a dataset of the same dimensions as the original data.
gen_df_1_5 <- function (k){</pre>
 df_1_5 <- data.frame(Day_of_year = 1:k, Draft_No = sample(min(data$Draft_No):max(data$Draft_No),1))
 df_1_5 <- rbind(data[k+1,],df_1_5)
 return(df_1_5)
alpha \leftarrow 0.05
k <- 9
data_random <- gen_df_1_5(k)</pre>
result <- test_hypothesis(data = data_random)</pre>
cat("P-value(lm):", result$p_lm_value, "\n")
cat("P-value(loess):", result$p_loess_value, "\n")
\#max_k = dim(data)[1]
max_k = 50
for(k in 10:max_k){
 if (k \% 3 != 0) {
   next
 data_random <- gen_df_1_5(k)</pre>
 res <- test hypothesis(data = data random)</pre>
 # we only test p loess value here.
 if(result$p_loess_value <= alpha ){</pre>
   cat("Reject HO Hypothesis","\n")
 }else{
   cat("Fail to Reject HO Hypothesis","\n")
 }
}
### Init Code
rm(list = ls())
library(ggplot2)
library(boot)
data <- read.csv("prices1.csv", sep = ";")</pre>
```

```
g_2_1 <- ggplot(data, aes(x = SqFt, y = Price)) + geom_point()</pre>
# fit a linear regression model
fit2_1_lm <- lm(Price ~ SqFt, data = data)
summary(fit2_1_lm)
minimal_x <- min(data$SqFt)</pre>
maximum x <- max(data$SqFt)</pre>
x_2_1_{m \leftarrow seq(minimal_x, maximum_x, 0.1)}
y_2_1_lm <- predict(fit2_1_lm, newdata = data.frame(SqFt = x_2_1_lm))</pre>
data_2_1 \leftarrow data.frame(SqFt = x_2_1_lm, Price = y_2_1_lm)
g_21 + geom_line(data = data_2_1, aes(x = x_2_1_lm, y = y_2_1_lm), color = "red")
rss_fun <- function(par,SqFt,Price) {</pre>
  a1 <- par[1]
  a2 <- par[2]
  b <- par[3]
  c <- par[4]
  predicted <- ifelse(SqFt > c, b + a1 * SqFt + a2 * (SqFt - c), b + a1 * SqFt)
 rss <- sum((Price - predicted)^2)
  return(rss)
}
c_func <- function(c, SqFt, Price) {</pre>
  # Init
  init_pars \leftarrow c(2, 0.1, 50,c)
  # Optimize using optim()
  result <- optim(par = init_pars, rss_fun, SqFt = SqFt, Price = Price)</pre>
  opt_pars <- result$par</pre>
  rss <- result$value
  return(list(c = opt_pars, params = opt_pars, rss = rss))
}
c <- 150
results <- c_func(c, SqFt = data$SqFt, Price = data$Price)</pre>
results
stat1 <- function(data, vn){</pre>
    data = as.data.frame(data[vn,])
    res <- c_func(c, SqFt = data$SqFt, Price = data$Price)</pre>
    return(res$param[4])
}
set.seed(12345)
```

```
res1 = boot(data, stat1, R=1000)
res1
print(boot.ci(res1))
# plot
plot(res1)
# Jackknife Function
jackknife <- function(data, fun) {</pre>
  n <- length(data)</pre>
  indices <- 1:n
  estimates <- numeric(n)</pre>
  for (i in 1:n) {
    jackknife_sample <- data[-i]</pre>
    estimates[i] <- fun(jackknife_sample)</pre>
  bias_correction <- (n - 1) * mean(estimates) - fun(data)</pre>
  jackknife_var <- ((n - 1) / n) * sum((estimates - mean(estimates) - bias_correction)^2)</pre>
 return(list(estimates = estimates, jackknife_var = jackknife_var))
}
# not working here,c_values
jackknife_results <- jackknife(c_values, function(c_val) {</pre>
  c_func(c_val, SqFt, Price)$params[1]
})
# Print Jackknife Results
cat("Jackknife Estimates of c:", jackknife_results$estimates, "\n")
cat("Jackknife Variance of c:", jackknife_results$jackknife_var, "\n")
```