Stat 135 Lab 1 - Jin Kweon

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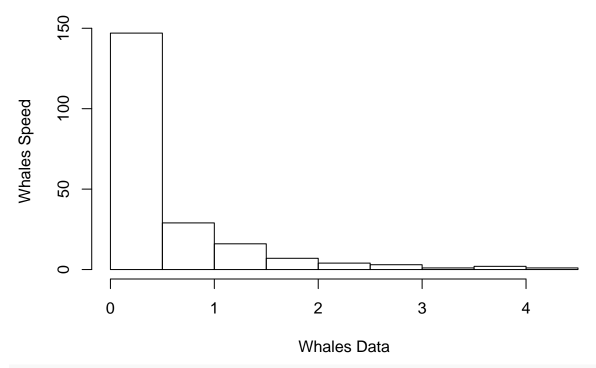
Data import

```
whales <- read.csv("/Users/yjkweon24/Desktop/Cal/2017Spring/Stat 135/Data/whales.txt", header = FALSE, colnames(whales) <- "Data"
options(digits=9, stringsAsFactors = FALSE)
whales$Data <- as.numeric(whales$Data)
newdata <- as.numeric(whales[[1]])</pre>
```

\mathbf{A}

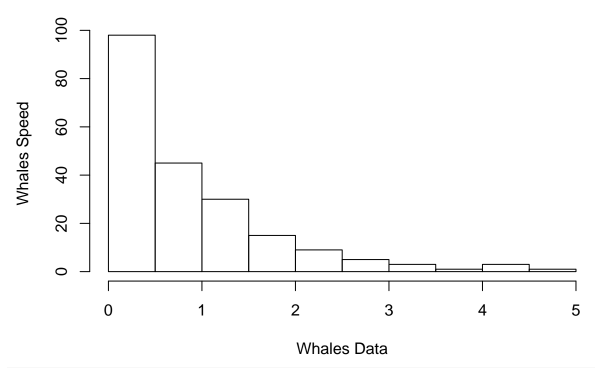
```
hist(newdata, main = "Whales Swimming", xlab = "Whales Data", ylab = "Whales Speed")
```

Whales Swimming



hist(rgamma(newdata, 0.9), main = "Whales Swimming with Gamma Fit", xlab = "Whales Data", ylab = "Whale

Whales Swimming with Gamma Fit



So, I can say the histogram looks like it fits into the gamma distribution.

The left portion is much bigger compared to the rightmost portions. The frequency gets smaller rapidl

To prove my statement, I used rgamma function and get the data, and plot them as histogram.

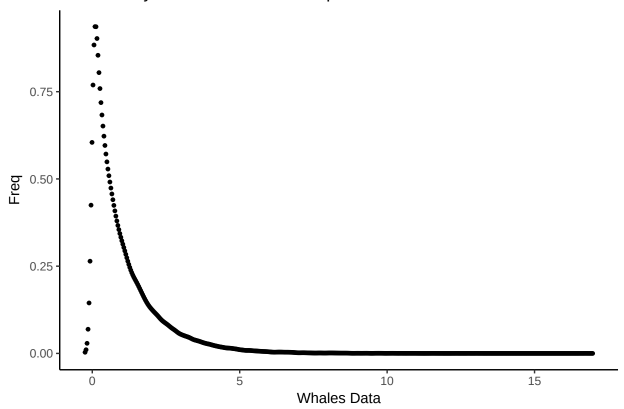
 \mathbf{B}

```
# Method of Moments Way
# when I say x ~ gamma(alpha, lambda), and x_bar = mean of sample space (210), expectation(x^2) = get2,
# alph(a) = (x_bar)^2 / [get2 - (x_bar)^2] and
# lambd(a) = (x_bar) / [get2 - (x_bar)^2]
x_bar = mean(newdata)
get2 = (x_bar)^2 + var(newdata)
alph = (x_bar)^2 / (get2 - (x_bar)^2) # Get Alpha.
lambd = (x_bar) / (get2 - (x_bar)^2) # Get Lambda

x <- rgamma(100000, shape=alph, scale=lambd) # Randomly get 100000 data from the alpha and lambda I got
densit <- density(x)
datas <- data.frame(x = densit$x, y = densit$y)

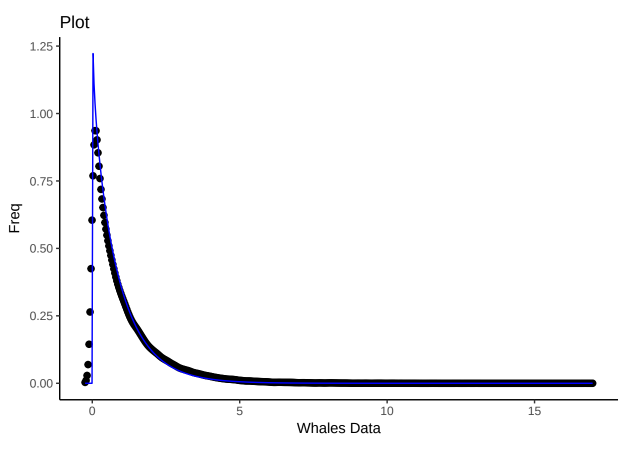
# Plot density from Rgamma points with MoM
ggplot(data = datas, aes(x = x, y = y)) +
geom_point(size = 1) +
theme_classic() + xlab("Whales Data") + ylab("Freq") + ggtitle("Plot Density from random Gamma points</pre>
```

Plot Density from random Gamma points with MoM



```
#Fit Parameters - check Fit
fit.params <- fitdistr(x, "gamma", lower = c(0, 1))

# Plot with density points - Fitparams into the MOM density
ggplot(data = datas, aes(x = x,y = y)) +
  geom_point(size = 2) +
  geom_line(aes(x=datas$x, y=dgamma(datas$x, fit.params$estimate["shape"], fit.params$estimate["rate"])
  theme_classic() + xlab("Whales Data") + ylab("Freq") + ggtitle("Plot")</pre>
```



So, basically, I conclude that alpha and lambda from MLE are nearly twice as the ones from MoM. alpha, lambda(MLE) = 2*alpha, lambda(Mom) ### C

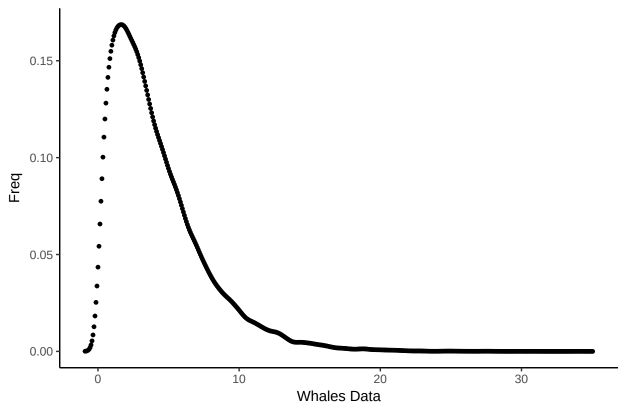
```
# Maximum Likehoold Way
# As the text book mentions "MLE are not given in closed form, obtaining their exact sampling distribut
\# So I went over "https://pdfs.semanticscholar.org/306d/d2f94d4e2a4e460ba26d07aba05c6f0b587a.pdf" and "
\# lambda = alpha / x_bar
# 1. alpha \sim ((3-s + sqrt((s-3)^2 + 24*s)) / 12*s) where s = log(x_bar) - mean(log(newdata)) -> within
# 2. alpha \sim (0.5 \ / \ s) \rightarrow (https://pdfs.semanticscholar.org/306d/d2f94d4e2a4e460ba26d07aba05c6f0b587a.p)
# 3. alpha -> from the textbook
# 4. alpha -> from mle() function -> lambda: 2.63269 and alpha: 1.59541
# Textbook way
loglikehood <- function(x){</pre>
  answer \leftarrow 210*log(x) - 210*log(x_bar) + sum(log(x))-n*digamma(x)
  return(answer)
\#z \leftarrow nleqslv(x, loglikehood)[[1]]
# MLE function using package of stats4
MLEf <- function(alph_mle1, lambd_mle1) {</pre>
     R = dgamma(whales$Data, alph_mle1, lambd_mle1)
     -sum(log(R))
}
```

```
mle1 = mle(MLEf, start = list(alph_mle1 = 1, lambd_mle1=1), method = "L-BFGS-B", lower = c(-Inf, 0), up
alph_mle1 = mle1@coef[[1]]
lambd_mle1 = mle1@coef[[2]]

x2 <- rgamma(50000, shape=alph_mle1, scale=lambd_mle1)
densit2 <- density(x2)
datas2 <- data.frame(x_mle1 = densit2$x, y_mle1 = densit2$y)

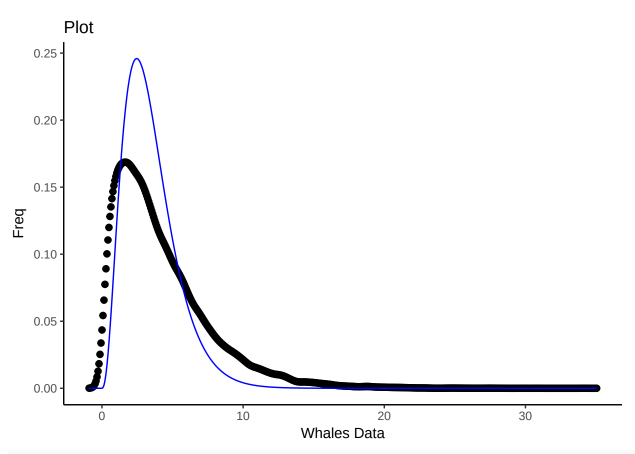
# Plot density from Rgamma points with MLE2
ggplot(data = datas2, aes(x = x_mle1, y = y_mle1)) +
    geom_point(size = 1) +
    theme_classic() + xlab("Whales Data") + ylab("Freq") + ggtitle("Plot Density from random Gamma points</pre>
```

Plot Density from random Gamma points with MLE1



```
#Fit Parameters - check Fit
fit.params2 <- fitdistr(x2, "gamma", lower = c(0, 1))

# Plot with density points - Fitparams into the MLE2 density
ggplot(data = datas2, aes(x = x_mle1,y = y_mle1)) +
    geom_point(size = 2) +
    geom_line(aes(x=datas2$x, y=dgamma(datas2$x, fit.params2$estimate["shape"], fit.params2$estimate["rat
    theme_classic() + xlab("Whales Data") + ylab("Freq") + ggtitle("Plot")</pre>
```

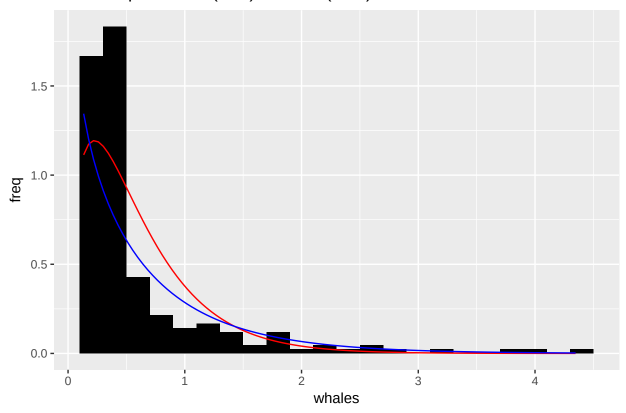


```
# The second way (Extra) - This might be wrong, although I got closed to alph_mle1 and lambd_mle1.
s = log(x_bar) - mean(log(newdata))
alph_mle2 = 0.5 / s
lambd_mle2 = alph_mle2 / mean(newdata)
```

 \mathbf{D}

```
hist1 <- ggplot(whales, aes(x = Data)) +
  geom_histogram(binwidth = 0.2, fill = "black", aes(y = ..density..), position = "identity", size = 1)
  xlab("whales") +ylab("freq") + ggtitle("Whales Speed MLE (Red) vs MoM (Blue)")+
  stat_function(fun = dgamma, args= c(shape=alph_mle1, scale=1/lambd_mle1), color = "red")+
  stat_function(fun = dgamma, args= c(shape=alph, scale=1/lambd), color = "blue")
hist1</pre>
```

Whales Speed MLE (Red) vs MoM (Blue)

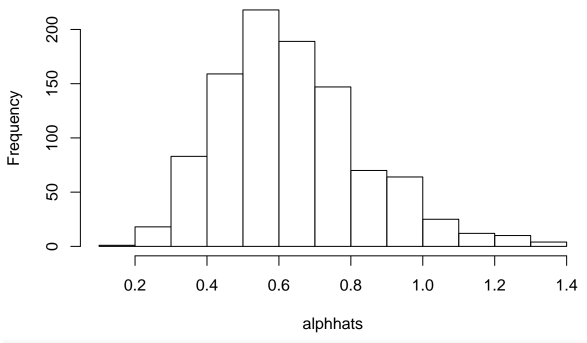


 \mathbf{E}

```
lambdhats <- numeric(0)</pre>
alphhats <- numeric(0)</pre>
for (i in 1:1000){
  do <- rgamma(210, shape = alph, scale = lambd)</pre>
  xbar <- mean(do)</pre>
  s <- var(do)
  lambdhat <- (xbar / s^2)</pre>
  alphhat <- (xbar^2 / s^2)
  lambdhats <- c(lambdhats, lambdhat)
  alphhats <- c(alphhats, alphhat)</pre>
mean(lambdhats)
## [1] 0.621232166
sd(lambdhats)
## [1] 0.225996432
mean(alphhats)
## [1] 0.637111713
sd(alphhats)
## [1] 0.201657391
```

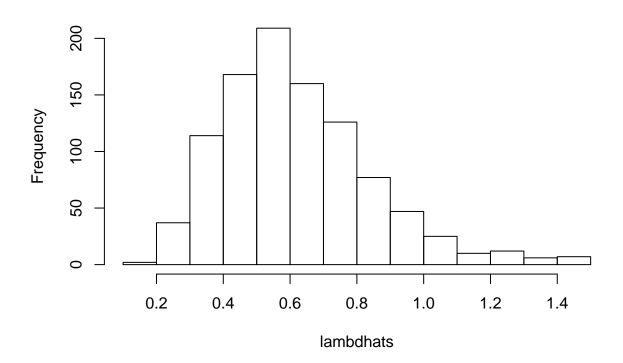
hist(alphhats)

Histogram of alphhats



hist(lambdhats)

Histogram of lambdhats



```
\mathbf{F}
```

```
lambdhats2 <- numeric(0)</pre>
alphhats2 <- numeric(0)</pre>
for (i in 1:1000){
  do2 <- rgamma(210, shape = alph_mle1, scale = lambd_mle1)</pre>
  xbar <- mean(do2)</pre>
  s <- var(do2)
  MLEf <- function(alph_mle1, lambd_mle1) {</pre>
     R = dgamma(do2, alph_mle1, lambd_mle1)
    -sum(log(R))
  }
  mle2 = mle(MLEf, start = list(alph_mle1 = 1, lambd_mle1=1), method = "L-BFGS-B", lower = c(-Inf, 0),
  lambdhat2 <- mle2@coef[[2]]</pre>
  alphhat2 <- mle2@coef[[1]]</pre>
  lambdhats2 <- c(lambdhats2, lambdhat2)</pre>
  alphhats2 <- c(alphhats2, alphhat2)</pre>
}
mean(lambdhats2)
sd(lambdhats2)
mean(alphhats2)
sd(alphhats2)
hist(alphhats2)
hist(lambdhats2)
# Looks normal
\mathbf{G}
# Three methods for confidence interval for MLE: exact method, approximations based on the large sample
#Since Bootstrap uses a lot of data, by the central theorem, we say the distribution is likely to be a
# alpha
Lower <-function(t){</pre>
  return(mean(alphhats2) - (sd(alphhats2)/sqrt(1000))*t)
Upper <- function(t2){</pre>
  \verb"return(mean(alphhats2) + (sd(alphhats2)/sqrt(1000))*t2)"
# For 90%
Lower()
Upper()
# For 95%
```

```
Lower()
Upper()
# For 99%
Lower()
Upper()
# Lambda
Boundary <- function(chi){</pre>
  return((1000*sd(lambdhats2)^2) / chi)
}
# For 90%
Boundary() #Lower
Boundary() # Upper
# For 95%
Boundary() #Lower
Boundary() # Upper
# For 99%
Boundary() #Lower
Boundary() # Upper
```