ETC 2420/5242 Lab 4 2016

Di Cook SOLUTION

Question 1

- 1. Simulate samples of size n = 30, 100, 500 from these distributions
 - a. Lognormal(2, 0.5)
 - b. Gamma(2, 4)

```
library(ggplot2)
library(gridExtra)

df_30 <- data.frame(x1=rlnorm(30, 2, 0.5), x2=rgamma(30, 2, 4))

ggplot(df_30, aes(x=x1)) + geom_histogram()

ggplot(df_30, aes(x=x2)) + geom_histogram()

df_100 <- data.frame(x1=rlnorm(100, 2, 0.5), x2=rgamma(100, 2, 4))

ggplot(df_100, aes(x=x1)) + geom_histogram()

ggplot(df_100, aes(x=x2)) + geom_histogram()

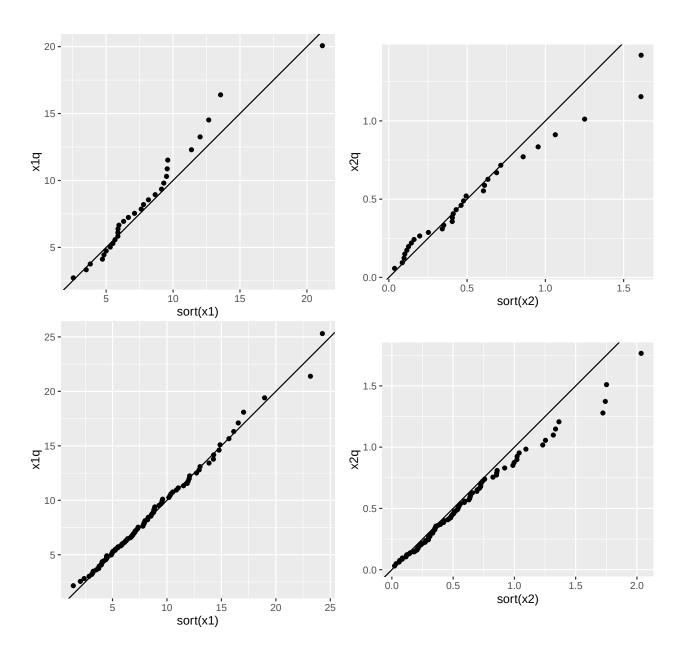
df_500 <- data.frame(x1=rlnorm(500, 2, 0.5), x2=rgamma(500, 2, 4))

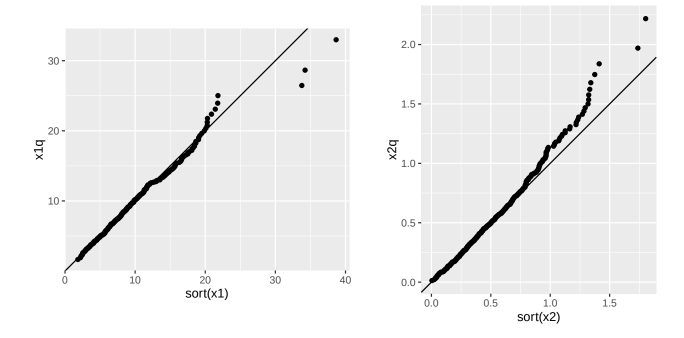
ggplot(df_500, aes(x=x1)) + geom_histogram()

ggplot(df_500, aes(x=x2)) + geom_histogram()</pre>
```

2. Make a QQ-plot of each these samples.

```
n <- 30
df_{30}x1q = qlnorm(c(1 - 0.5^(1/n), (2:(n-1) - 0.3175) /
                        (n + 0.365), 0.5^{(1/n)}, 2, 0.5)
df_{30}x2q = qgamma(c(1 - 0.5^(1/n), (2:(n-1) - 0.3175)) /
                        (n + 0.365), 0.5^{(1/n)}, 2, 4)
p1 \leftarrow ggplot(df_30, aes(x=sort(x1), y=x1q)) +
  geom_abline(intercept=0, slope=1) +
  geom_point() + coord_equal()
p2 \leftarrow ggplot(df_30, aes(x=sort(x2), y=x2q)) +
  geom_abline(intercept=0, slope=1) +
  geom_point() + coord_equal()
grid.arrange(p1, p2, ncol=2)
n <- 100
df_{100}x1q = qlnorm(c(1 - 0.5^(1/n), (2:(n-1) - 0.3175)) /
                        (n + 0.365), 0.5^{(1/n)}, 2, 0.5)
df_{100}x2q = qgamma(c(1 - 0.5^(1/n), (2:(n-1) - 0.3175) /
                        (n + 0.365), 0.5^{(1/n)}, 2, 4)
p1 <- ggplot(df_100, aes(x=sort(x1), y=x1q)) +
  geom_abline(intercept=0, slope=1) +
  geom_point() + coord_equal()
p2 \leftarrow ggplot(df_100, aes(x=sort(x2), y=x2q)) +
  geom_abline(intercept=0, slope=1) +
  geom_point() + coord_equal()
grid.arrange(p1, p2, ncol=2)
n <- 500
```





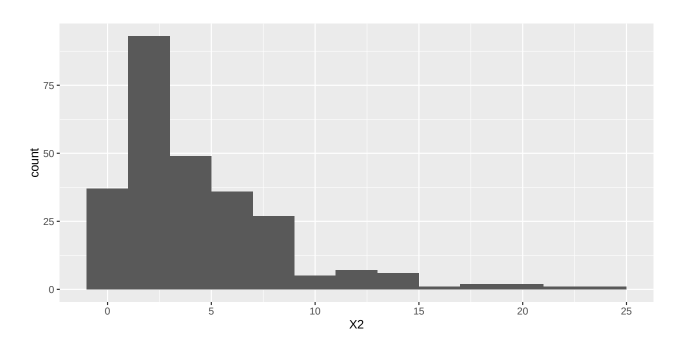
Question 2

Using this code, generate a sample of size n=267 from a Gamma(1.2,0.25) distribution.

```
set.seed(123)
X2 <- rgamma(n=267, 1.2, 0.25)</pre>
```

a. Plot the sample, using a histogram, describe the shape of the distribution.

```
ggplot(data.frame(X2), aes(x=X2)) + geom_histogram(binwidth=2)
```

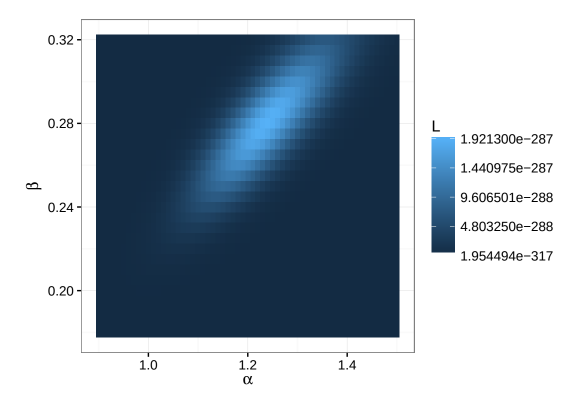


- b. What parameters of the gamma distribution were used to simulate the sample? (α, β) 1.2, 0.25
- c. If we are to use maximum likelihood distribution what values would we expect to get as the parameter estimates? Close to 1.2, 0.25
- d. Write a function to compute the likelihood function.

```
nmle <- function(x, a, b) {
  f <- prod(dgamma(x, a, b))
  return(f)
}</pre>
```

e. Plot the likelihood function for a range of values of α, β that shows the maximum likelihood estimates for each parameter.

```
a <- seq(0.9, 1.5, 0.01)
b <- seq(0.18, 0.32, 0.005)
g <- expand.grid(x=a, y=b)
g$f <- 0
for (i in 1:nrow(g)) {
    g$f[i] <- nmle(X2, g$x[i], g$y[i])
}
ggplot(g, aes(x=x, y=y, fill=f)) + geom_tile() + xlab(expression(alpha)) + ylab(expression(beta)) + them
    scale_fill_continuous("L") +
    theme(aspect.ratio=1)</pre>
```



f. Look up the function fitdistr from the MASS library. Explain what this does. Use it to find the MLE estimates for α, β . How do these compare with the values you read off your plot?

```
library(MASS)
fitdistr(X2, "gamma")

# shape rate

# 1.23994762 0.28051981

# (0.09624333) (0.02667898)
```

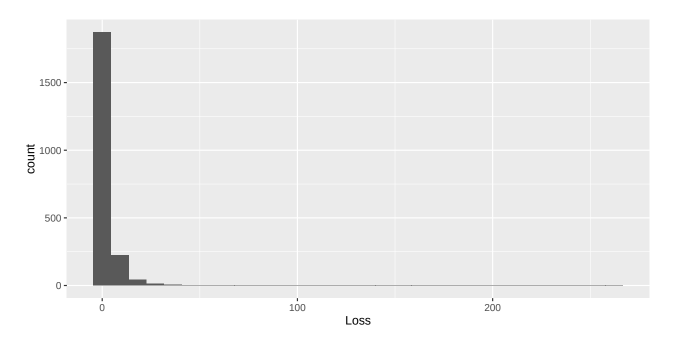
Question 3

Take a look at the data set danishuni from the CASdatasets library.

```
library(CASdatasets)
data(danishuni)
```

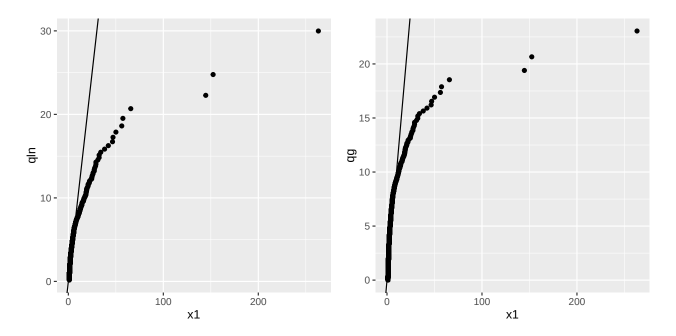
a. Make a histogram of the Loss. Describe the shape.

```
ggplot(danishuni, aes(x=Loss)) + geom_histogram()
```



b. Fit both a gamma and lognormal distribution to the sample, i.e. find the MLEs.

c. Produce a QQ-plot for each of the distributions.



d. Which is the better fit to the sample? Neither!! Maybe we should try a pareto?

TURN IN

- Your .Rmd file
- Your Word (or pdf) file that results from knitting the Rmd.
- Make sure your group members are listed as authors, one person per group will turn in the report
- DUE: Wednesday after the lab, by 7am, loaded into moodle

Resources

- PSU lecture notes on MLE
- CASdatasets