Stat5361-Homework 3

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Question1

Derive the updating rules in the given algorithm based on the construction of an EM algorithm.

$$\begin{split} Q[\Psi|\Psi(k)] &= \sum_{i=1}^{n} \sum_{j=1}^{m} p_{ij}^{(k+1)} [log(\pi_i) + log(\phi(y_i - x_i^T \beta_j; 0, \sigma^2))] \\ &\qquad \qquad \sum_{j=1}^{m} \pi_j = 1 \\ L(\pi_1, ..., \pi_m, \lambda) &= Q[\Psi|\Psi(k)] - \lambda(\sum_{j=1}^{m} \pi_j - 1) = 0 \end{split}$$

Calculate the first derivative of $L(\pi_1,...,\pi_m,\lambda)$ respect to π_j and λ , for j=1,...,m:

$$L(\pi_{1},...,\pi_{m},\lambda)_{\pi_{j}}^{'}=0;L(\pi_{1},...,\pi_{m},\lambda)_{\lambda}^{'}=0$$

$$\sum_{i=1}^{n} p_{ij}^{(k+1)}(\frac{1}{\pi_j}) - \lambda = 0$$

Slove for π_j

$$\pi_j = \frac{\sum_{i=1}^n p_{ij}^{(k+1)}}{\lambda}$$

$$\sum_{j=1}^{m} \pi_j = \frac{\sum_{j=1}^{m} \sum_{i=1}^{n} p_{ij}^{(k+1)}}{\lambda} = \frac{\sum_{i=1}^{n} \sum_{j=1}^{m} p_{ij}^{(k+1)}}{\lambda} = \frac{n}{\lambda} = 1; \lambda = n$$

$$\pi_j = \frac{\sum_{i=1}^n p_{ij}^{(k+1)}}{n}$$

$$Q[\Psi|\Psi(k)] = \sum_{i=1}^{n} \sum_{j=1}^{m} p_{ij}^{(k+1)} [log(\pi_j) + log(\frac{1}{\sqrt{2\pi}\sigma} exp(-\frac{(y_i - x_i^T \beta_j)^2}{2\sigma^2}))]$$

$$= \sum_{i=1}^{n} \sum_{j=1}^{m} p_{ij}^{(k+1)} [log(\pi_j) + log(\frac{1}{\sqrt{2\pi}\sigma}) + (-\frac{(y_i - x_i^T \beta_j)^2}{2\sigma^2})]$$

$$=\sum_{i=1}^{n}\sum_{j=1}^{m}p_{ij}^{(k+1)}log(\pi_{j})-\frac{1}{2}\sum_{i=1}^{n}\sum_{j=1}^{m}p_{ij}^{(k+1)}log(2\pi\sigma^{2})-\frac{1}{2}\sum_{i=1}^{n}\sum_{j=1}^{m}p_{ij}^{(k+1)}\frac{(y_{i}-x_{i}^{T}\beta_{j})^{2}}{\sigma^{2}}=I_{1}-\frac{1}{2}I_{2}-\frac{1}{2}I_{3}$$

Where $I_1 = \sum_{i=1}^n \sum_{j=1}^m p_{ij}^{(k+1)} log(\pi_j); I_2 = \sum_{i=1}^n \sum_{j=1}^m p_{ij}^{(k+1)} log(2\pi\sigma^2); \text{ and } I_3 = \sum_{i=1}^n \sum_{j=1}^m p_{ij}^{(k+1)} \frac{(y_i - x_i^T \beta_j)^2}{\sigma^2}$

$$I_3 = \sum_{i=1}^{n} \sum_{j=1}^{m} p_{ij}^{(k+1)} \left(\frac{(y_i - x_i^T \beta_j)^2}{2\sigma^2} \right)$$

Which is a qudratic form. It is the sum of m quadratic forms each involving a single β_j for every certain j.

$$\sum_{i=1}^{n} p_{ij}^{(k+1)} (\frac{(y_i - x_i^T \beta_j)^2}{2\sigma^2}) = \sum_{i=1}^{n} p_{ij}^{(k+1)} [x_i^T (\frac{y_i}{x_i^T} - \beta_j)^2]$$

From the property of sample mean, β_j must be the mean of a weighted sample of $\frac{y_1}{x_1}, ..., \frac{y_n}{x_n^T}$. And for each element, it has the weight: $p_{ij} \cdot x_i \cdot x_i^T$. It can be represented in the following form:

$$\sum_{i=1}^{n} p_{ij}^{(k+1)} x_i x_i^T (\frac{y_i}{x_i^T} - \beta_j)^2$$

Solve for β_j :

$$\beta_j = \frac{\sum_{i=1}^n p_{ij}^{(k+1)} x_i x_i^T (\frac{y_i}{x_i^T})}{\sum_{i=1}^n p_{ij}^{(k+1)} x_i x_i^T} = \frac{\sum_{i=1}^n p_{ij}^{(k+1)} x_i y_i}{\sum_{i=1}^n p_{ij}^{(k+1)} x_i x_i^T}$$

Only I_2 and I_3 contain σ

$$I_2 + I_3 = \sum_{i=1}^{n} \sum_{j=1}^{m} p_{ij}^{(k+1)} ln(\sigma) + \sum_{i=1}^{n} \sum_{j=1}^{m} p_{ij}^{(k+1)} \frac{(y_i - x_i^T \beta_j)^2}{2\sigma^2}$$

Minimizing $I_2 + I_3$ to maximaze $Q[\Psi|\Psi(k)]$. Now we knew that β_j is the weighted mean of $\frac{y_i}{x_i^T}$, therefore, σ must be the sample variance of all data for every j = 1, ..., m.

$$\sigma^{2(k+1)} = \frac{\sum_{i=1}^{n} \sum_{j=1}^{m} p_{ij}^{(k+1)} (y_i - x_i^T \beta_j^{(k+1)})^2}{\sum_{i=1}^{n} \sum_{j=1}^{m} p_{ij}^{(k+1)}}$$
$$= \frac{\sum_{i=1}^{n} \sum_{j=1}^{m} p_{ij}^{(k+1)} (y_i - x_i^T \beta_j^{(k+1)})^2}{n}$$

Question2

(a)

$$g(x) \propto (2x^{\theta-1} + x^{\theta-\frac{1}{2}}e^{-x})$$

$$C\int_{0}^{\infty} (2x^{\theta-1} + x^{\theta-\frac{1}{2}})e^{-x}dx = 1$$

$$2C\gamma_{(\theta)} + C\gamma_{(\theta+\frac{1}{2})} = 1$$

$$C = \frac{1}{2\gamma_{(\theta)} + C\gamma_{(\theta+\frac{1}{2})}}$$

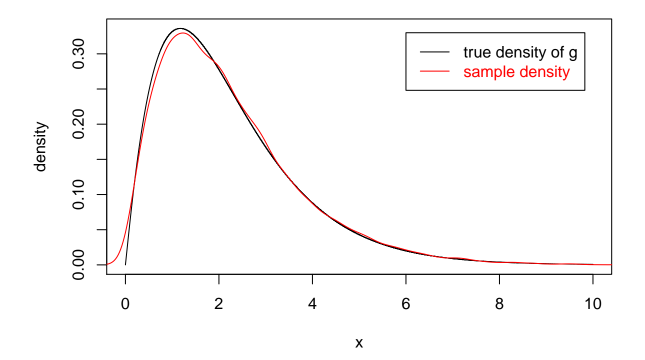
$$\therefore g(x) = \frac{2\gamma_{(\theta)}}{2\gamma_{(\theta)} + \gamma_{(\theta+\frac{1}{2})}} \frac{1}{\gamma_{(\theta)}} x^{\theta-1} e^{-x} + \frac{\gamma_{(\theta+\frac{1}{2})}}{2\gamma_{(\theta)} + \gamma_{(\theta+\frac{1}{2})}} * \frac{1}{\gamma_{(\theta+\frac{1}{2})}} x^{\theta-\frac{1}{2}} e^{-x}$$

It is the mixture of $\gamma(\theta, 1)$ and $\gamma(\theta + \frac{1}{2}, 1)$, where the weights are $\frac{2\gamma_{(\theta)}}{2\gamma_{(\theta)} + \gamma_{(\theta + \frac{1}{2})}}$ and $\frac{\gamma_{(\theta + \frac{1}{2})}}{2\gamma_{(\theta)} + \gamma_{(\theta + \frac{1}{2})}}$

(b)

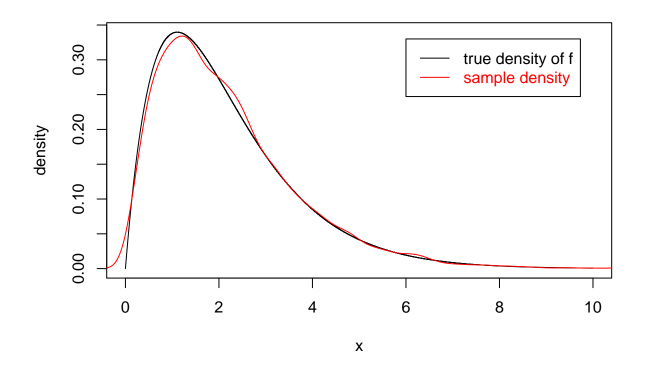
```
Repeat the following step 1 and step 2 for 10,000 times:
Step 1: u \sim unif(0,1)
Step 2: find x, where Cg(x) = u
sample_q \leftarrow c(sample_q, x)
#you can change the value of theta, we set it equals to 2 in order to calculate conveniently
theta \leftarrow 2
C <- 1 / (2 * gamma(theta) + gamma(theta + 1/2))
#pdf of g(x)
func_g <- function(x){</pre>
  return(C * (2 * x ** (theta - 1) + x ** (theta - 1/2)) * exp(-x))
#calculate the inverse of g-unit(1)
inverse_function <- function(x){</pre>
  #u <- runif(1)
  cdf_g <- integrate(func_g, lower = 0, upper = x)$value</pre>
  inverse_value <- cdf_g - u
 return(inverse value)
  #return()
}
#generate 10000 samples
sample_g <- c()</pre>
for(i in 1:10000){
  u <- runif(1)
  inverse_method <- uniroot(inverse_function,interval = c(0,20))$root
  sample_g <- c(sample_g,inverse_method)</pre>
sampleg_pdf <- density(sample_g)</pre>
#calculate the true value of q_pdf
x \leftarrow seq(from = 0, to = 10, by = 0.0001)
g_value <- sapply(x,func_g)</pre>
##-----Q2-b-----
```

```
#plot the true density and the sample density in one figure
plot(g_value~x,type = "l", xlab = "x", ylab = "density")
lines(sampleg_pdf, col = "red")
#add legend the fist two parameters denote the location, lty =1 denotes the type of legend is line
legend(6,0.33,c("true density of g", "sample density"), col = c("black","red"),text.col = c("black","red")
```



```
(c) \sqrt{4+x} < 2 + \sqrt{x} \text{ for all } x > 0 \text{ because } (2+\sqrt{x})^2 - (\sqrt{4+x})^2 = 4 \cdot \sqrt{x} > 0. Therefore, \frac{f(x)}{g(x)} \le 1, we choose \alpha = \frac{1}{C}. Step 1: u \sim unif(0,1) Step 2: find x^*, where Cg(x) = u Calculate f(x^*) and g(x^*) u^* \sim unif(0,1) if u^* < \frac{f(x^*)}{g(x^*)} sample \leftarrow c(sample,x^*) Repeat Step 1 and Step 2 until the length of sample is 10,000 #you can change the value of theta, we set it equals to 2 in order to calculate conveniently theta <-2 C <-1 / (2 * gamma(theta) + gamma(theta + 1/2)) #pdf of g(x)
```

```
func_g <- function(x){</pre>
 return(C * (2 * x ** (theta - 1) + x ** (theta - 1/2)) * exp(-x))
}
#calculate the inverse of g-unit(1)
inverse_function <- function(x){</pre>
  #u <- runif(1)
  cdf_g <- integrate(func_g, lower = 0, upper = x)$value</pre>
  inverse_value <- cdf_g - u
 return(inverse_value)
  #return()
}
##-----Q2-c-----
#pdf of f
func_f <- function(x){</pre>
 return(((4 + x) ** 0.5) * (x ** (theta - 1)) * exp(-x))
}
const <- 1 / integrate(func_f,lower = 0, upper = Inf)$value</pre>
func_f_true <- function(x){</pre>
 value <- func_f(x) * const</pre>
 return(value)
}
#calculate the true value of f_pdf
x \leftarrow seq(from = 0, to = 10, by = 0.0001)
f_value <- sapply(x,func_f_true)</pre>
f_sample_rejection <-c()</pre>
while(length(f_sample_rejection)<10000){
  u <- runif(1)
  sample_value <- uniroot(inverse_function,interval = c(0,20))$root
  sample_value_g <- func_g(sample_value) / C</pre>
  sample_value_f <- func_f(sample_value)</pre>
  ratio <- sample_value_f / sample_value_g
  rejection_value <- runif(1)</pre>
  if(ratio >= rejection_value){
    f_sample_rejection <- c(f_sample_rejection, sample_value)</pre>
  }
}
sampleg_pdf <- density(f_sample_rejection)</pre>
##-----Q2-c-----
#plot the true density and the sample density in one figure
plot(f_value~x,type = "l", xlab = "x", ylab = "density")
lines(sampleg_pdf, col = "red")
#add legend the fist two parameters denote the location, lty =1 denotes the type of legend is line
legend(6,0.33,c("true density of f", "sample density"), col = c("black", "red"), text.col = c("black", "re
```



Question 3

(a)

For

0 < x < 1

,

$$f(x) = \frac{x^{\theta - 1}}{1 + x^2} + \sqrt{2 + x^2} \cdot (1 - x)^{\beta - 1} \le x^{\theta - 1} + \sqrt{3} \cdot (1 - x)^{\beta - 1}$$

$$g_1^*(x) = x^{\theta - 1} \int_0^1 g_1^*(x) dx = \frac{x^{\theta}}{\theta} \Big|_0^1 = \frac{1}{\theta} = \frac{1}{C_1} \Rightarrow g_1(x) = \theta \cdot x^{\theta - 1}$$

$$g_2^*(x) = \sqrt{3} \cdot (1 - x)^{\beta - 1} \int_0^1 g_2^*(x) dx = -\sqrt{3} \cdot \frac{(1 - x)^{\beta}}{\beta} \Big|_0^1 = \frac{\sqrt{3}}{\beta} = \frac{1}{C_2} \Rightarrow g_2(x) = \beta \cdot (1 - x)^{\beta - 1}$$

$$g(x) = \frac{1}{2} g_1(x) + \frac{1}{2} g_2(x)$$

$$f(x) < x^{\theta - 1} + \sqrt{3} \cdot (1 - x)^{\beta - 1}$$

$$\alpha g(x) = \alpha \cdot \frac{1}{2} \cdot \theta \cdot x^{\theta - 1} + \alpha \cdot \frac{1}{2} \cdot \beta \cdot (1 - x)^{\beta - 1}$$

$$1 \leq \frac{1}{2} \cdot \alpha \cdot \theta$$

$$\frac{2}{\theta} \leq \alpha$$

$$1 \leq \frac{1}{2} \cdot \alpha \cdot \beta$$

$$\frac{2}{\beta} \leq \alpha$$

$$\therefore \alpha = max(\frac{2}{\theta}, \frac{2}{\beta})$$
Step 1: u -weight $\sim unif(0, 1)$
Step 2: if u -weight $\leq unif(0, 1)$, find x^* , where $g_i(x) = u$ elase, $u \sim unif(0, 1)$, find x^* , where $g_i(x) = u$ else, $u \sim unif(0, 1)$, find x^* , where $g_i(x) = u$ Step 3: calculate $f(x^*)$ and $g(x^*)$

$$u^* \sim unif(0, 1)$$
if $u^* \leq \frac{f(x^*)}{g(x^*)}$
sample $\leftarrow c(sample, x^*)$
Repeat Step 1,2 and 3 until the length of sample is 10,000
#you can change the value of theta and beta theta factor $\leftarrow 0$
$U^* \subset (C^* \cap C^*) = 0$
$U^* \subset (C^* \cap C^*$

Step 2: if u-weight $\leq w_1$

 $sample \leftarrow c(sample, x^*)$

theta_factor <- 9 beta_factor <- 6</pre>

#pdf of g(x)

return(g1)

return(g2)

}

func_g1 <- function(x){</pre>

func_g2 <- function(x){</pre>

g1 <- func_g1(x)</pre> $g2 \leftarrow func_g2(x)$ return(W1*g1+W2*g2)

elase,

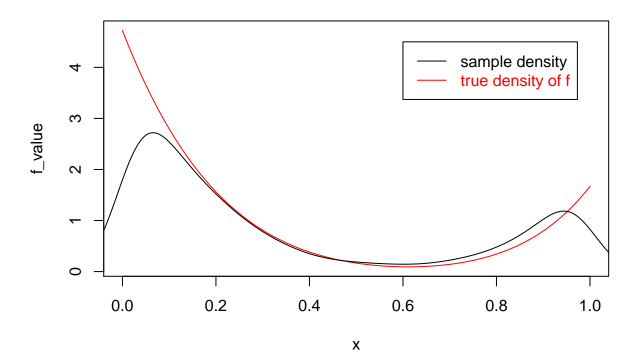
 $u^* \sim unif(0,1)$ if $u^* \le \frac{f(x^*)}{g(x^*)}$

```
\# x \leftarrow seq(from = 0, to = 1, by = 0.001)
\# cdf_g \leftarrow c()
# for(i in 1:length(x)){
\# cdf_g \leftarrow c(cdf_g, integrate(func_g, lower = 0, upper = x[i])$value)
# #g_value <- sapply(x, func_g)
\# plot(cdf_g~x, type = "l", xlab = "x", ylab = "density")
#calculate the inverse of g-unit(1)
inverse_function <- function(x){</pre>
  #u <- runif(1)
  cdf_g <- integrate(func_g, lower = 0, upper = x)$value</pre>
  inverse_value <- cdf_g - u
  return(inverse_value)
}
##-----Q3-a-----
#pdf of f
func_f <- function(x){</pre>
  f1 \leftarrow (x \hat theta_factor-1)) / (1+x^2)
  f2 \leftarrow ((2+x^2)^0.5) * ((1-x)^(beta_factor - 1))
  return(f1+f2)
func_f_true <- function(x){</pre>
  value <- func_f(x) * const</pre>
  return(value)
}
#solve for the constant for f_pdf
integrate(func_f, lower = 0, upper = 1)$value
## [1] 0.2993621
const <- 1/integrate(func_f, lower = 0, upper = 1)$value</pre>
#true value of f_pdf
x \leftarrow seq(from = 0, to = 1, by = 0.001)
f_value <- sapply(x,func_f_true)</pre>
f_sample_rejection <-c()</pre>
while(length(f_sample_rejection)<10000){
  u_weight <- runif(1)</pre>
  if(u_weight < 0.5){
    u <- runif(1)
    func_g <- func_g1</pre>
    sample_value <- uniroot(inverse_function,interval = c(0,1))$root</pre>
  }
  else{
    u <- runif(1)
    func_g <- func_g2</pre>
    sample_value <- uniroot(inverse_function,interval = c(0,1))$root</pre>
  sample_value_g <- func_g_mix(sample_value)</pre>
  sample_value_f <- func_f(sample_value)</pre>
  ratio <- sample_value_f / (alpha_factor * sample_value_g)
  rejection_value <- runif(1)
  if(ratio >= rejection_value){
```

```
f_sample_rejection <- c(f_sample_rejection, sample_value)
}

sample_pdf <- density(f_sample_rejection)

#plot the true density and the sample density in one figure
plot(f_value~x,type = "l",col = "red")
lines(sample_pdf,type = "l", xlab = "x", ylab = "density", main = "using rejection method to get sample
legend(0.6,4.5,c("sample density", "true density of f"), col = c("black","red"),text.col = c("black","r</pre>
```



str(f_sample_rejection)
max(f_sample_rejection)

(b)

$$f(x_1) = \frac{x^{\theta-1}}{1+x^2} \le x^{\theta-1} = \alpha_1 \cdot g_1(x)$$

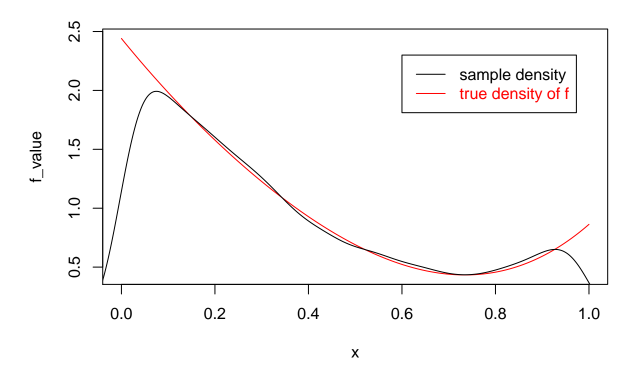
$$f(x_2) = \sqrt{2 + x^2} \cdot (1 - x)^{\beta - 1} \le \sqrt{3} \cdot (1 - x)^{\beta - 1} = \alpha_2 \cdot g_2(x)$$

Then,
$$\alpha_1 = \frac{1}{\theta}$$
, and $\alpha_2 = \frac{\sqrt{3}}{\beta}$

```
Thus, when runif > \frac{\overline{\theta}}{1 + \sqrt{3}}, sample from g_2(x)
Step 1: u-weight \sim unif(0,1)
Step 2: if u-weight > weighted-boundary
u \sim unif(0,1), find x^*, where g_2(x^*) = u
calculate ratio = \frac{f_2(x^*)}{\alpha g_2(x^*)}
else.
u \sim unif(0,1), find x^*, where g_1(x^*) = u
calculate ratio = \frac{f_1(x^*)}{\alpha g_1(x^*)}
Step 3: u^* \sim unif(0,1)
if u^* < ratio
sample \leftarrow c(sample, x^*)
Repeat Step 1, 2 and 3 until the length of sample is 10,000
#you can change the value of theta and beta
theta_factor <- 6
beta_factor <- 3</pre>
alpha_1 <- 1 / theta_factor</pre>
alpha_2 \leftarrow (3^0.5) / beta_factor
weight_boundary <-alpha_1 / (alpha_1 + alpha_2)</pre>
#pdf of g(x)
func_g1 <- function(x){</pre>
  g1 <- theta_factor * (x ^ (theta_factor-1))</pre>
  return(g1)
func_g2 <- function(x){</pre>
  g2 <- (beta_factor) * ((1-x)^(beta_factor-1))</pre>
  return(g2)
\# x \leftarrow seq(from = 0, to = 1, by = 0.001)
# cdf_g <- c()
\# for(i in 1:length(x)) \{
  cdf_g \leftarrow c(cdf_g, integrate(func_g, lower = 0, upper = x[i])$value)
# }
# #g_value <- sapply(x, func_g)
# plot(cdf_g~x,type = "l", xlab = "x", ylab = "density")
#calculate the inverse of g-unit(1)
inverse_function <- function(x){</pre>
  cdf_g <- integrate(func_g, lower = 0, upper = x)$value</pre>
  inverse_value <- cdf_g - u
  return(inverse_value)
}
##-----Q3-b-----
#pdf of f
func_f1 <- function(x){</pre>
 f1 <- (x ^ (theta_factor-1)) / (1+x^2)
```

```
return(f1)
}
func_f2 <- function(x){</pre>
  f2 \leftarrow ((2+x^2)^0.5) * ((1-x)^(beta_factor - 1))
  return(f2)
func f <- function(x){</pre>
  f1 <- func f1(x)
  f2 <- func_f2(x)
  return(f1+f2)
integrate(func_f, lower = 0, upper = 1)$value
## [1] 0.5793803
const <- 1/integrate(func_f, lower = 0, upper = 1)$value</pre>
func_f_true <- function(x){</pre>
  value <- func_f(x) * const</pre>
  return(value)
#solve for the constant for f_pdf
#true value of f_pdf
x \leftarrow seq(from = 0, to = 1, by = 0.001)
f_value <- sapply(x,func_f_true)</pre>
f_sample_rejection <-c()</pre>
while(length(f sample rejection)<10000){
  random_weight <- runif(1)</pre>
  if(random_weight > weight_boundary){
    u <- runif(1)
    func_g <- func_g2</pre>
    sample_value <- uniroot(inverse_function,interval = c(0,1))$root</pre>
    sample_value_g <- func_g2(sample_value)</pre>
    sample_value_f <- func_f2(sample_value)</pre>
    ratio <- sample_value_f / (alpha_2 * sample_value_g)</pre>
  }
  else{
    u <- runif(1)
    func_g <- func_g1</pre>
    sample_value <- uniroot(inverse_function,interval = c(0,1))$root</pre>
    sample_value_g <- func_g1(sample_value)</pre>
    sample_value_f <- func_f1(sample_value)</pre>
    ratio <- sample_value_f / (alpha_1 * sample_value_g)</pre>
  rejection_value <- runif(1)</pre>
  if(ratio >= rejection_value){
    f_sample_rejection <- c(f_sample_rejection, sample_value)</pre>
}
sample_pdf <- density(f_sample_rejection)</pre>
```

```
#plot the true density and the sample density in one figure
plot(f_value~x,type = "l",col = "red")
lines(sample_pdf,type = "l", xlab = "x", ylab = "density", main = "using rejection method to get sample
legend(0.6,2.3,c("sample density", "true density of f"), col = c("black","red"),text.col = c("black","r
```



```
# str(f_sample_rejection)
# max(f_sample_rejection)
```