

STA510 Assignment 02 Solution

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2024-10-27

Problem 01 (a) || R

```
# Set seed for reproducibility
set.seed(123)

# Number of simulations
n <- 10000

# Define the limits of the integral
a <- -2 - sqrt(3)
b <- -1 / sqrt(3)

# Define the function f(x)
f <- function(x) {
  exp(-x) / (1 + x^2)^2
}

# Generate n uniformly distributed random variables in the interval [a, b]
x <- runif(n, min = a, max = b)

# Calculate the crude Monte Carlo estimate
I_CMC <- (b - a) * mean(f(x))

# Print the result
cat("Estimated value of I using Crude Monte Carlo:", I_CMC)
```

Estimated value of I using Crude Monte Carlo: 1.166561

Problem 01 (b) || R

```
# Define the function f(x)
f <- function(x) {
  exp(-x) / (1 + x^2)^2
}

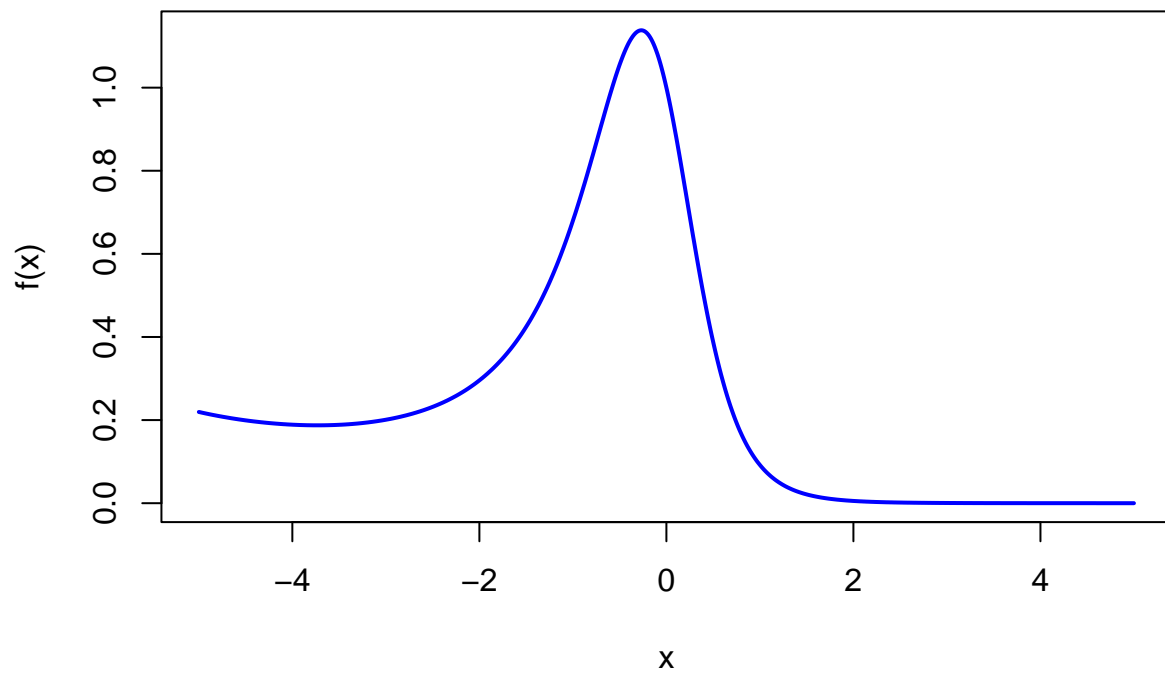
# Define the interval for plotting
x_values <- seq(-5, 5, length.out = 1000)

# Evaluate the function at each x value
f_values <- f(x_values)

# Plot the function f(x) over the interval [-5, 5]
```

```
plot(x_values, f_values, type = "l", col = "blue", lwd = 2,  
     xlab = "x", ylab = "f(x)",  
     main = expression(paste("Plot of ", f(x), " = e^(-x) / (1 + x^2)^2")),  
     xlim = c(-5, 5))
```

Plot of $f(x) = e^{-x} / (1 + x^2)^2$



PROBLEM 01(b):

Estimating I using Antithetic Random Variables —

The integral $I = \int_a^b f(x) dx$.

Where $f(x) = \frac{e^{-x}}{(1+x^2)^2}$ and $[a,b]$ is the interval $[-2-\sqrt{3}, \frac{1}{\sqrt{3}}]$

Let X be a uniform Random Variable on $[a,b]$. We generate $n/2$ independent samples of X . Let's denote these sample as $x_1, x_2, \dots, x_{n/2}$. For each x_i , create an antithetic variable y_i defined as $y_i = a+b - x_i$. By construction, y_i is also uniformly distributed on $[a,b]$ but is negatively correlated with x_i .

Now, for each pair (x_i, y_i) , we have to calculate $f(x_i)$ and $f(y_i)$.

The antithetic estimator \hat{I}_{AT} is the average of $f(x_i)$ and $f(y_i)$.

$$\hat{I}_{AT} = \frac{b-a}{n} \sum_{i=1}^{n/2} [f(x_i) + f(y_i)]$$

Why this approach reasonable —

The use of antithetic variable reduces the variance of the estimator because the values $f(x_i)$ and $f(y_i)$ tend to balance out each other's deviations from the mean, leading to a more stable estimate of I .

Problem 01 (c) || R

```
# Number of simulations
n <- 10000 # Total number of simulations
n_half <- n / 2 # Half of the simulations for generating antithetic pairs

# Define the limits of the integral
a <- -2 - sqrt(3)
b <- -1 / sqrt(3)

# Define the function f(x)
f <- function(x) {
  exp(-x) / (1 + x^2)^2
}

# Generate n/2 uniformly distributed random variables in the interval [a, b]
X <- runif(n_half, min = a, max = b)

# Create antithetic variables
Y <- a + b - X

# Calculate the Monte Carlo estimate using antithetic random variables
I_AT <- (b - a) * mean(f(X) + f(Y)) / 2

# Print the result
cat("Estimated value of I using Antithetic Monte Carlo:", I_AT)

## Estimated value of I using Antithetic Monte Carlo: 1.17512
```

Problem 01 (d) || Theory

PROBLEM 01 (d):

$$f(x) = \frac{e^{-x}}{(1+x^2)^2} \quad ; \quad g(x) = \frac{c}{1+x^2} \text{ for } -2-\sqrt{3} \leq x \leq -1/\sqrt{3}$$

$$g(x) = 0, \text{ otherwise}$$

for $g(x)$ to be a pdf, the integral over the interval must equal 1,

$$\int_{-2-\sqrt{3}}^{-1/\sqrt{3}} g(x) dx = 1$$

$$\Rightarrow \int_{-2-\sqrt{3}}^{-1/\sqrt{3}} \frac{c}{1+x^2} dx = 1$$

$$\Rightarrow c \int_{-2-\sqrt{3}}^{-1/\sqrt{3}} \frac{1}{1+x^2} dx = 1 \quad \left[\int \frac{1}{1+x^2} dx = \arctan(x) \right]$$

$$\Rightarrow c \left[\arctan\left(\frac{-1}{\sqrt{3}}\right) - \arctan(-2-\sqrt{3}) \right] = 1$$

$$\Rightarrow c \left(\frac{-\pi}{6} - \frac{-5\pi}{12} \right) = 1$$

$$\Rightarrow c \left(\frac{5\pi}{12} - \frac{\pi}{6} \right) = 1$$

$$\Rightarrow c \left(\frac{5\pi - 2\pi}{12} \right) = 1$$

$$\Rightarrow c \cdot \frac{3\pi}{12} = 1$$

$$\Rightarrow c = \frac{4}{\pi} \approx 1.2732$$

Problem 01 (e) || R

```
# Define the function f(x)
f <- function(x) {
  exp(-x) / (1 + x^2)^2
}

# Calculate the constant c for g(x)
# Define the limits of the interval
lower_limit <- -2 - sqrt(3)
upper_limit <- -1 / sqrt(3)

# Calculate the integral of 1 / (1 + x^2) over the given interval
integral_value <- atan(upper_limit) - atan(lower_limit)

# Calculate the constant c such that g(x) integrates to 1
c <- 1 / integral_value

# Define the function g(x)
g <- function(x) {
  if (x >= -2 - sqrt(3) && x <= -1 / sqrt(3)) {
    return(c / (1 + x^2))
  } else {
    return(0)
  }
}

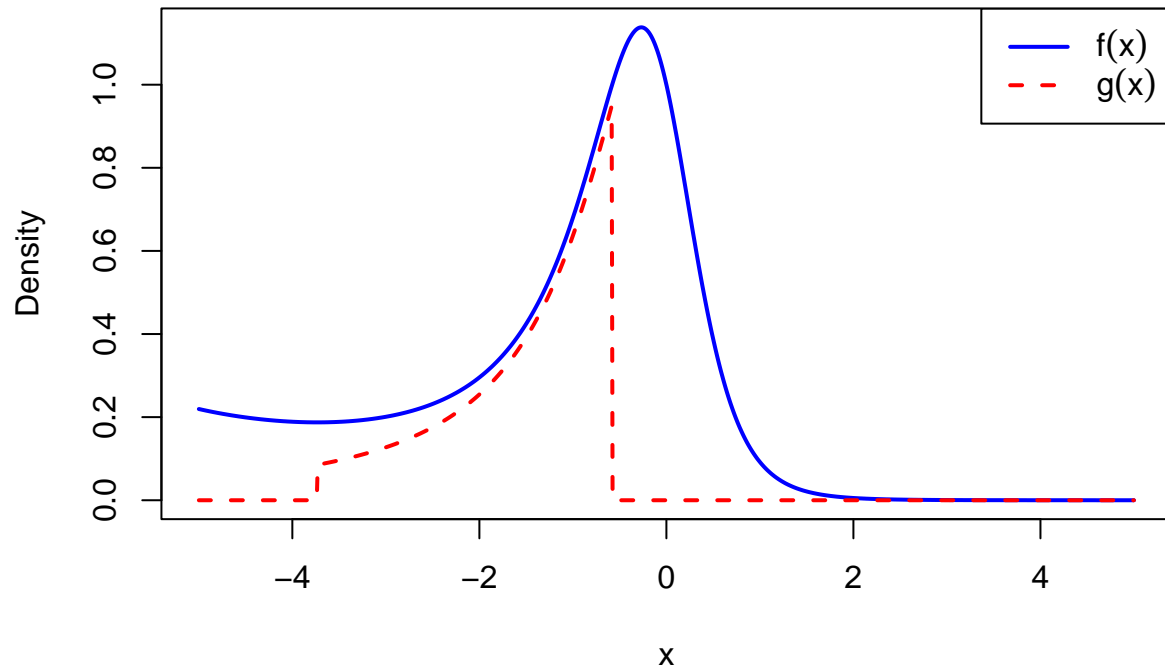
# Create a vectorized version of g(x) for plotting
g_vectorized <- Vectorize(g)

# Define the interval for plotting
x_values <- seq(-5, 5, length.out = 1000)

# Evaluate the functions at each x value
f_values <- f(x_values)
g_values <- g_vectorized(x_values)

# Plot f(x) and g(x) on the same graph
plot(x_values, f_values, type = "l", col = "blue", lwd = 2,
     xlab = "x", ylab = "Density",
     main = expression(paste("Plot of ", f(x), " and ", g(x))),
     xlim = c(-5, 5))
lines(x_values, g_values, col = "red", lwd = 2, lty = 2)
legend("topright", legend = c(expression(f(x)), expression(g(x))),
     col = c("blue", "red"), lty = c(1, 2), lwd = 2)
```

Plot of $f(x)$ and $g(x)$



```
# Print the result
```

```
cat("The value of c for g(x) is:", c)
```

```
## The value of c for g(x) is: 1.27324
```

```
# Assess g(x) as an importance function-----  
# The function g(x) approximates f(x) in the interval where g(x) is non-zero. If g(x)  
# captures the general behavior of f(x) in this interval, it can serve as a good  
# importance function. Ideally, g(x) should closely follow the shape of f(x) to be  
# effective.
```

PROBLEM 01(f):

$$g(x) = \frac{c}{1+x^2}, \text{ where } -2-\sqrt{3} \leq x \leq -1/\sqrt{3}$$

The CDF is defined as

$$\begin{aligned} h(x) &= \int_{-2-\sqrt{3}}^x g(t) dt \\ &= \int_{-2-\sqrt{3}}^x \frac{c}{1+t^2} dt \quad \left[\int \frac{1}{1+t^2} dt = \arctan(t) \right] \\ &= c [\arctan(x) - \arctan(-2-\sqrt{3})] \end{aligned}$$

To ensure $h(x)$ ranges from 0 to 1 over the interval $[-2-\sqrt{3}, -1/\sqrt{3}]$, we normalize it,

$$h(x) = \frac{\arctan(x) - \arctan(-2-\sqrt{3})}{\arctan(-1/\sqrt{3}) - \arctan(-2-\sqrt{3})}$$

Now, let u represent a value of the CDF such that $u \in [0, 1]$. To find the inverse CDF $h^{-1}(u)$, we set:

$$u = \frac{\arctan(x) - \arctan(-2-\sqrt{3})}{\arctan(-1/\sqrt{3}) - \arctan(-2-\sqrt{3})}$$

$$\Rightarrow u \cdot [\arctan(-1/\sqrt{3}) - \arctan(-2-\sqrt{3})] = \arctan(x) - \arctan(-2-\sqrt{3})$$

$$\Rightarrow \arctan(x) = u \cdot [\arctan(-1/\sqrt{3}) - \arctan(-2-\sqrt{3})] + \arctan(-2-\sqrt{3})$$

$$\Rightarrow x = \tan [u \cdot (\arctan(-1/\sqrt{3}) - \arctan(-2-\sqrt{3})) + \arctan(-2-\sqrt{3})]$$

So, the inverse CDF $h^{-1}(x)$ is

$$h^{-1}(x) = \tan [x \cdot (\arctan(-1/\sqrt{3}) - \arctan(-2-\sqrt{3})) + \arctan(-2-\sqrt{3})]$$

Problem 01 (g) || R

```
# Number of simulations
n <- 10000

# Define the limits of the integral
lower_limit <- -2 - sqrt(3)
upper_limit <- -1 / sqrt(3)

# Define the constant c (from previous calculations)
c <- 1.27324

# Define the function f(x)
f <- function(x) {
  exp(-x) / (1 + x^2)^2
}

# Define the density function g(x)
g <- function(x) {
  if (x >= lower_limit && x <= upper_limit) {
    return(c / (1 + x^2))
  } else {
    return(0)
  }
}

# Create a vectorized version of g(x) for calculation
g_vectorized <- Vectorize(g)

# Define the inverse CDF function G^-1(u) for sampling from g(x)
G_inverse <- function(u) {
  tan(u * (atan(upper_limit) - atan(lower_limit)) + atan(lower_limit))
}

# Generate random samples from g(x) using inverse transform sampling
u_samples <- runif(n)
x_samples <- G_inverse(u_samples)

# Calculate the importance sampling estimator
weights <- f(x_samples) / g_vectorized(x_samples)
I_IM <- mean(weights)

# Print the result
cat("Estimated value of I using Importance Sampling:", I_IM)
```

Estimated value of I using Importance Sampling: 1.17835

Problem 01 (h) || R

```
# Number of replications
num_replications <- 1000
n <- 10000 # Number of samples for each estimate

# Define the limits of the integral
```

```

lower_limit <- -2 - sqrt(3)
upper_limit <- -1 / sqrt(3)

# Define the constant c (from previous calculations)
c <- 1.27324

# Define the function f(x)
f <- function(x) {
  exp(-x) / (1 + x^2)^2
}

# Define the density function g(x)
g <- function(x) {
  if (x >= lower_limit && x <= upper_limit) {
    return(c / (1 + x^2))
  } else {
    return(0)
  }
}

# Vectorize the density function g(x) for calculation
g_vectorized <- Vectorize(g)

# Define the inverse CDF function G^-1(u) for sampling from g(x)
G_inverse <- function(u) {
  tan(u * (atan(upper_limit) - atan(lower_limit)) + atan(lower_limit))
}

# Function to calculate the Crude Monte Carlo estimate
estimate_CMC <- function() {
  x <- runif(n, min = lower_limit, max = upper_limit)
  (upper_limit - lower_limit) * mean(f(x))
}

# Function to calculate the Antithetic estimate
estimate_AT <- function() {
  x <- runif(n / 2, min = lower_limit, max = upper_limit)
  y <- lower_limit + upper_limit - x
  (upper_limit - lower_limit) * mean((f(x) + f(y)) / 2)
}

# Function to calculate the Importance Sampling estimate
estimate_IM <- function() {
  u_samples <- runif(n)
  x_samples <- G_inverse(u_samples)
  weights <- f(x_samples) / g_vectorized(x_samples)
  mean(weights)
}

# Generate 1,000 replications of each estimator
estimates_CMC <- replicate(num_replications, estimate_CMC())
estimates_AT <- replicate(num_replications, estimate_AT())
estimates_IM <- replicate(num_replications, estimate_IM())

```

```

# Calculate the mean and standard deviation for each set of estimates
mean_CMC <- mean(estimates_CMC)
sd_CMC <- sd(estimates_CMC)

mean_AT <- mean(estimates_AT)
sd_AT <- sd(estimates_AT)

mean_IM <- mean(estimates_IM)
sd_IM <- sd(estimates_IM)

# Print the results
cat("Results of 1,000 Replications:\n")

## Results of 1,000 Replications:
cat("Crude Monte Carlo Estimator ||
    Mean:", mean_CMC, " Standard Deviation:", sd_CMC, "\n")

## Crude Monte Carlo Estimator ||
##      Mean: 1.176997 Standard Deviation: 0.006840798

cat("Antithetic Estimator ||
    Mean:", mean_AT, " Standard Deviation:", sd_AT, "\n")

## Antithetic Estimator ||
##      Mean: 1.177121 Standard Deviation: 0.00415498

cat("Importance Sampling Estimator ||
    Mean:", mean_IM, " Standard Deviation:", sd_IM, "\n")

## Importance Sampling Estimator ||
##      Mean: 1.17714 Standard Deviation: 0.002221089

# Compare and Comment
#
# Mean Estimates:
# All three estimators provide mean estimates that are very close to each other,
# suggesting that all methods are accurately estimating I. This consistency across
# methods validates the implementation and the effectiveness of each technique.
#
# Standard Deviation (Variability):
# 01. Crude Monte Carlo has the highest standard deviation (0.0068), which is
# expected since it does not incorporate any variance reduction techniques.
# 02. Antithetic Sampling reduces the standard deviation to 0.00415 by using
# negatively correlated random variables, thus lowering variability and
# increasing precision.
# 03. Importance Sampling achieves the lowest standard deviation (0.00222) due to
# the choice of  $g(x)$  that approximates  $f(x)$ . This targeted sampling approach
# significantly reduces the estimator's variability.
#
#
# The reduction in standard deviation indicates that both Antithetic Sampling
# and Importance Sampling are more efficient than the Crude Monte Carlo method.
# Among these, Importance Sampling is the most efficient due to its smallest
# standard deviation.

```

Problem 01 (i) || R

```
# Standard deviation of the Crude Monte Carlo estimator (from previous results)
sigma_CMC <- 0.006840798

# Set the desired margin of error and confidence level
margin_of_error <- 0.001
z_value <- qnorm(0.995) # z-value for 99% confidence level (0.5% on each tail)

# Calculate the required number of simulations
n_required <- (z_value * sigma_CMC / margin_of_error)^2

# Print the result
cat("Number of simulations required for 99% confidence and a margin of 0.001:",
    ceiling(n_required), "\n")
```

```
## Number of simulations required for 99% confidence and a margin of 0.001: 311
```

Problem 02 (a) || R

```
# Define the intensity (lambda) and the time interval
lambda <- 3
time_interval <- 30

# Generate interarrival times from an exponential distribution
interarrival_times <- rexp(1000, rate = lambda)

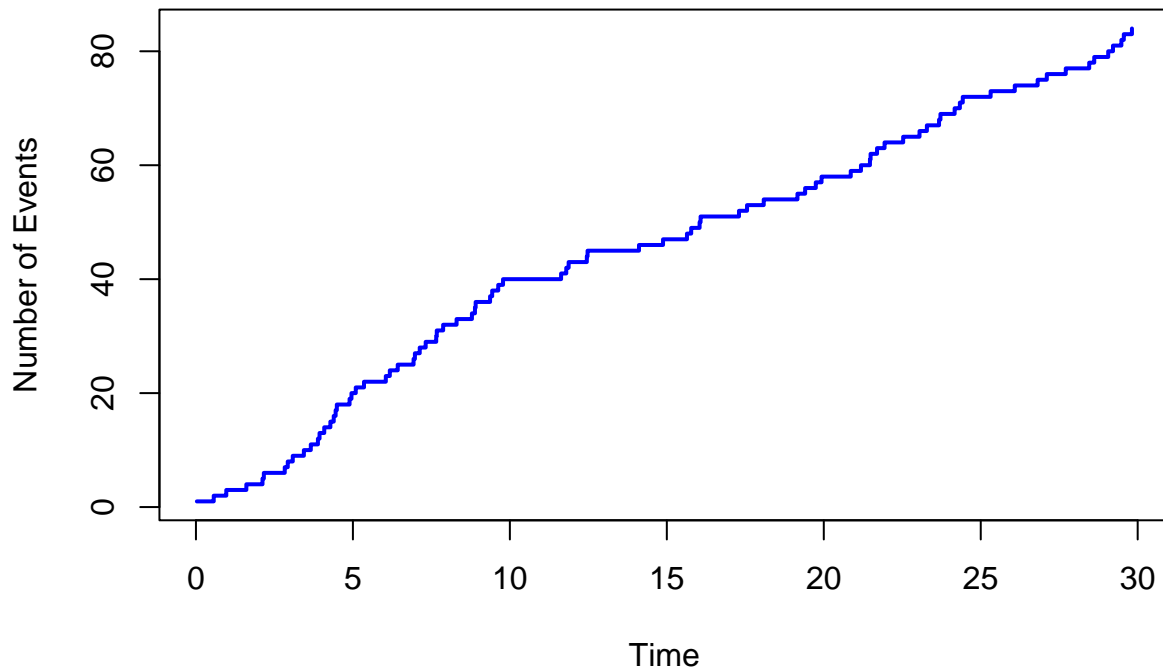
# Calculate the cumulative sum of interarrival times to get event times
event_times <- cumsum(interarrival_times)

# Filter event times to be within the interval [0, 30]
event_times <- event_times[event_times <= time_interval]

# Count the number of events over time to plot N(t)
N_t <- 1:length(event_times)

# Plot the Poisson process N(t)
plot(event_times, N_t, type = "s", col = "blue", lwd = 2,
      xlab = "Time", ylab = "Number of Events",
      main = expression(paste("Poisson Process N(t) with ", lambda, " = 3")))
```

Poisson Process $N(t)$ with $\lambda = 3$



```
# Print Result
cat("Number of events in [0, 30]:", length(event_times), "\n")
```

```
## Number of events in [0, 30]: 84
```

Problem 02 (b) || Theory

PROBLEM 02 (b):

for Poisson process,

Expected Value, $E = \lambda t$

Variance, $\text{Var} = \lambda t$

So, Standard Deviation, $\sigma = \sqrt{\lambda t}$

Here, $\lambda = 3$,

for $N(5)$, $t = 5$, So, $E = \lambda t = 15$

$$\sigma = \sqrt{\lambda t} = \sqrt{15} = 3.87$$

for $N(20)$, $t = 20$, So, $E = \lambda t = 60$

$$\sigma = \sqrt{\lambda t} = \sqrt{60} = 7.75$$

Problem 02 (c) || R

```
# Define the intensity (lambda) and the time interval
lambda <- 3
time_interval <- 30
num_realizations <- 40

# Create a plot with a fixed range
plot(0, 0, type = "n", xlim = c(0, time_interval), ylim = c(0, 100),
     xlab = "Time", ylab = "Number of Events",
     main = paste(num_realizations, "Realizations of Poisson Process with",
                  expression(lambda), "= 3"))

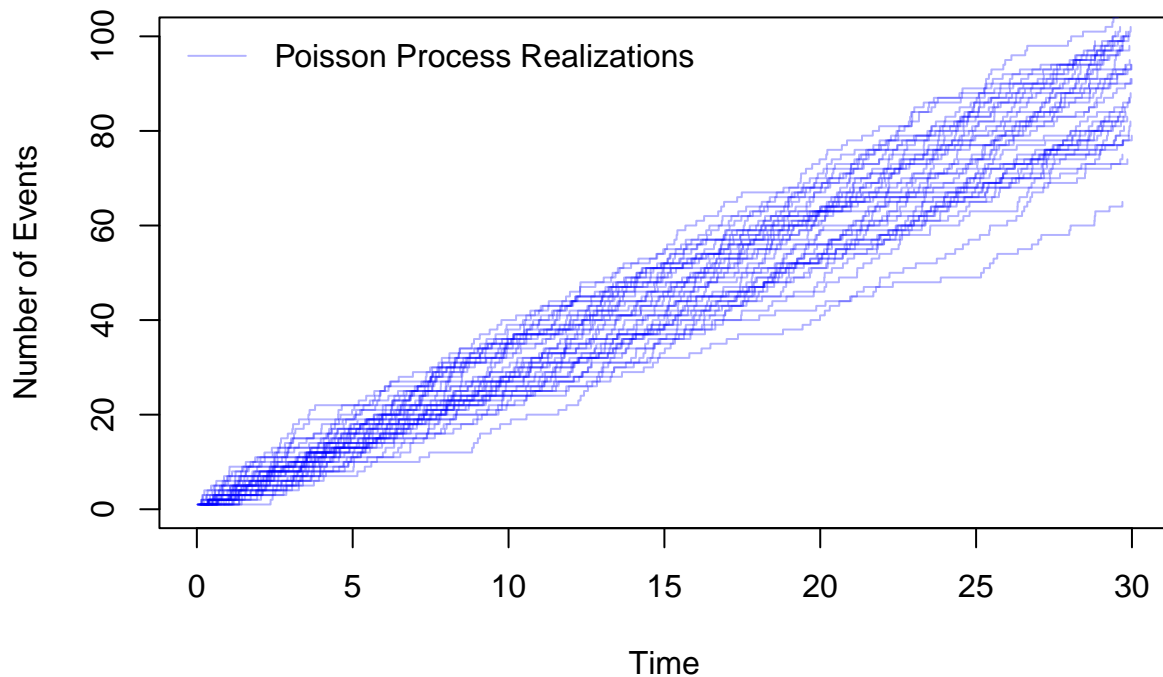
# Generate and plot each realization
for (i in 1:num_realizations) {
  # Generate interarrival times for one realization
  interarrival_times <- rexp(1000, rate = lambda)
  event_times <- cumsum(interarrival_times)

  # Filter event times to be within the interval [0, 30]
  event_times <- event_times[event_times <= time_interval]
  N_t <- 1:length(event_times)

  # Plot the realization as a step function
  lines(event_times, N_t, type = "s", col = rgb(0, 0, 1, alpha = 0.3))
}

# Adding a legend
legend("topleft", legend = "Poisson Process Realizations",
      col = rgb(0, 0, 1, alpha = 0.3), lwd = 1, bty = "n")
```

40 Realizations of Poisson Process with $\lambda = 3$



```
# Comment on Pattern  
# The realizations exhibit variability around the expected number of events calculated  
# in Problem 2b.  
# At t=5, the expected number of events is 15, with a standard deviation of 3.87.  
# At t=20, the expected number of events is 60, with a standard deviation of 7.75.  
#  
#  
# In the plot, we can see that the number of events at any time t tends to fluctuate  
# around these expected values. The step functions illustrate the random nature of event  
# occurrences in a Poisson process. The density of events increases over time due to the  
# constant intensity lambda, reflecting the increasing number of events as time progresses.
```

Problem 02 (d) || R

```
# Define the intensity (lambda) and the number of simulations  
lambda <- 3  
num_simulations <- 10000 # Number of simulations  
  
### (i) Probability of Observing at Least 80 Events in [0, 30]  
  
# Define the time interval for part (i)  
time_interval_30 <- 30  
  
# Simulate the number of events in each realization for the interval [0, 30]  
event_counts_30 <- replicate(num_simulations, {  
  # Generate interarrival times and calculate cumulative sum for each realization
```



```

interarrival_times <- rexp(1000, rate = lambda)
event_times <- cumsum(interarrival_times)

# Count the number of events within the interval [0, 30]
sum(event_times <= time_interval_30)
})

# Calculate the probability of observing at least 80 events in [0, 30]
probability_at_least_80 <- mean(event_counts_30 >= 80)

# Print the result
cat("Estimated probability of observing at least 80 events in [0, 30]:",
    probability_at_least_80, "\n")

```

Estimated probability of observing at least 80 events in [0, 30]: 0.8759

(ii) Probability of Observing Less Than 30 Events in [0, 10]

```

# Define the time interval for part (ii)
time_interval_10 <- 10

# Simulate the number of events in each realization for the interval [0, 10]
event_counts_10 <- replicate(num_simulations, {
  # Generate interarrival times and calculate cumulative sum for each realization
  interarrival_times <- rexp(1000, rate = lambda)
  event_times <- cumsum(interarrival_times)

  # Count the number of events within the interval [0, 10]
  sum(event_times <= time_interval_10)
})

# Calculate the probability of observing less than 30 events in [0, 10]
probability_less_than_30 <- mean(event_counts_10 < 30)

# Print the result
cat("Estimated probability of observing less than 30 events in [0, 10]:",
    probability_less_than_30, "\n")

```

Estimated probability of observing less than 30 events in [0, 10]: 0.479

Problem 02 (e) || R

```

# Define the parameters for the truncated normal distribution
a <- 8
b <- 17.5
mu <- 12.5
sigma <- 1

# Define the truncated normal pdf function
f_truncated <- function(t, mu, sigma, a, b) {
  # Calculate the normal pdf and CDF values
  numerator <- dnorm((t - mu) / sigma) / sigma
  denominator <- pnorm((b - mu) / sigma) - pnorm((a - mu) / sigma)
}

```

```

# Compute the pdf of the truncated normal distribution
if (t >= a && t <= b) {
  return(numerator / denominator)
} else {
  return(0)
}
}

# Define the intensity function lambda(t)
lambda <- function(t) {
  100 * sapply(t, f_truncated, mu = mu, sigma = sigma, a = a, b = b)
}

# Define the interval for the process
t_min <- 7
t_max <- 18

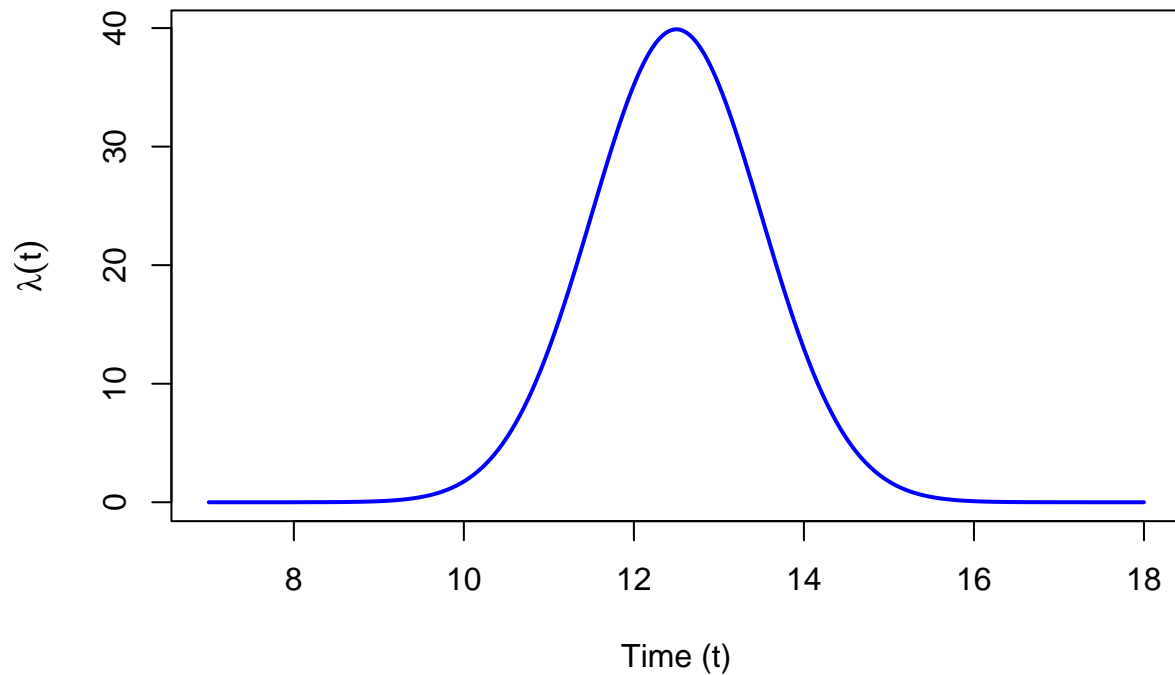
# Define the interval for plotting lambda(t)
t_values <- seq(t_min, t_max, length.out = 1000)

# Evaluate the intensity function lambda(t) over the interval
lambda_values <- lambda(t_values)

# Plot the intensity function lambda(t)
plot(t_values, lambda_values, type = "l", col = "blue", lwd = 2,
     xlab = "Time (t)", ylab = expression(lambda(t)),
     main = expression(paste("Intensity Function ", lambda(t), " over [7, 18]")))

```

Intensity Function $\lambda(t)$ over [7, 18]



Problem 02 (f) || R

```
# Define the parameters for the truncated normal distribution
a <- 8
b <- 17.5
mu <- 12.5
sigma <- 1

# Define the truncated normal pdf function (same as Problem 2e)
f_truncated <- function(t, mu, sigma, a, b) {
  numerator <- dnorm((t - mu) / sigma) / sigma
  denominator <- pnorm((b - mu) / sigma) - pnorm((a - mu) / sigma)

  if (t >= a && t <= b) {
    return(numerator / denominator)
  } else {
    return(0)
  }
}

# Define the intensity function lambda(t) (same as Problem 2e)
lambda <- function(t) {
  100 * sapply(t, f_truncated, mu = mu, sigma = sigma, a = a, b = b)
}
```

```

# Define the interval for the process
t_min <- 7
t_max <- 18

# Find the maximum value of lambda(t) over the interval [7, 18]
t_values <- seq(t_min, t_max, length.out = 1000)
lambda_max <- max(lambda(t_values))

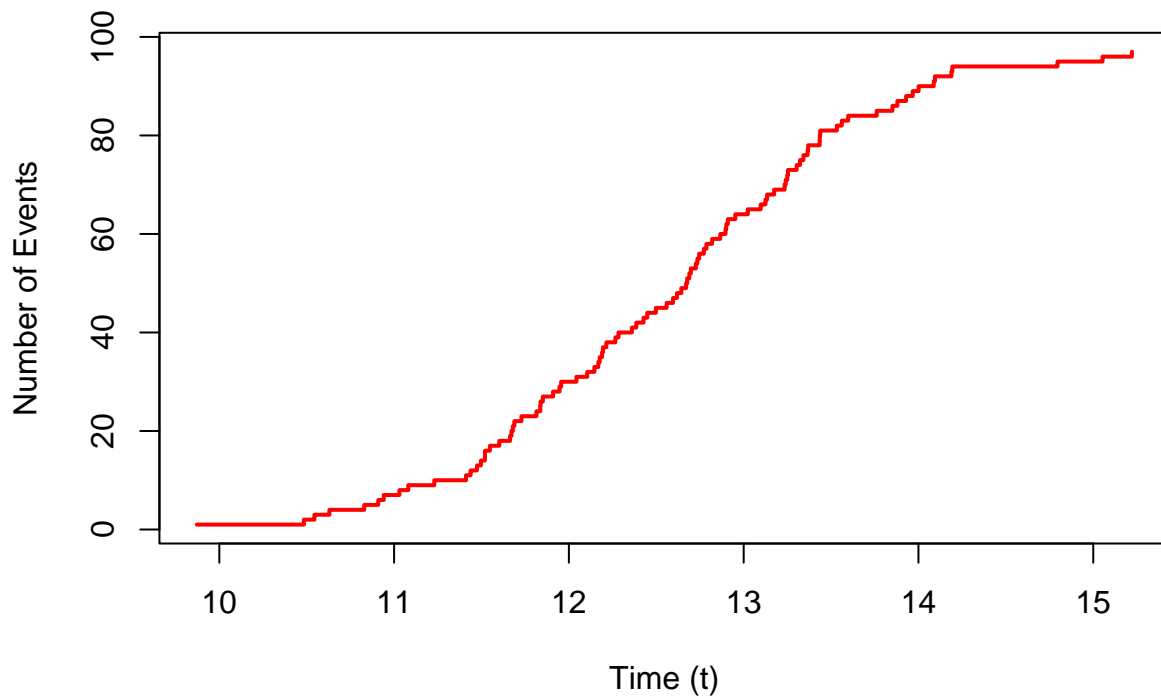
# Simulate a homogeneous Poisson process with intensity lambda_max
candidate_times <- cumsum(rexp(10000, rate = lambda_max))
candidate_times <-
  candidate_times[candidate_times <= t_max & candidate_times >= t_min]

# Apply the thinning algorithm to select events based on lambda(t) / lambda_max
accepted_times <-
  candidate_times[runif(length(candidate_times)) <=
    lambda(candidate_times) / lambda_max]

# Plot the non-homogeneous Poisson process
plot(accepted_times, 1:length(accepted_times), type = "s", col = "red", lwd = 2,
     xlab = "Time (t)", ylab = "Number of Events",
     main =
       expression(paste("Non-Homogeneous Poisson Process with ", lambda(t))))

```

Non-Homogeneous Poisson Process with $\lambda(t)$



Problem 02 (g) || R

```
# Define the parameters for the truncated normal distribution
a <- 8
b <- 17.5
mu <- 12.5
sigma <- 1

# Define the truncated normal pdf function (same as Problem 2e)
f_truncated <- function(t, mu, sigma, a, b) {
  numerator <- dnorm((t - mu) / sigma) / sigma
  denominator <- pnorm((b - mu) / sigma) - pnorm((a - mu) / sigma)

  if (t >= a && t <= b) {
    return(numerator / denominator)
  } else {
    return(0)
  }
}

# Define the intensity function lambda(t) (same as Problem 2e)
lambda <- function(t) {
  100 * sapply(t, f_truncated, mu = mu, sigma = sigma, a = a, b = b)
}

# Define the interval for the process
t_min <- 7
t_max <- 18

# Find the maximum value of lambda(t) over the interval [7, 18]
t_values <- seq(t_min, t_max, length.out = 1000)
lambda_max <- max(lambda(t_values))

# Function to generate a single replication of non-homogeneous Poisson process
generate_process <- function() {
  # Generate candidate event times using homogeneous Poisson process with lambda_max
  candidate_times <- cumsum(rexp(1000, rate = lambda_max))
  candidate_times <-
    candidate_times[candidate_times <= t_max & candidate_times >= t_min]

  # Apply the thinning algorithm to select events based on lambda(t) / lambda_max
  accepted_times <- candidate_times[runif(length(candidate_times)) <=
    lambda(candidate_times) / lambda_max]

  return(accepted_times)
}

# Number of replications
num_replications <- 10000

# Simulate 10,000 replications and count arrivals in the specified intervals
arrivals_before_10 <- numeric(num_replications)
arrivals_11_to_13 <- numeric(num_replications)

for (i in 1:num_replications) {
```

```

# Generate a single replication of the process
event_times <- generate_process()

# Count the number of arrivals before 10:00
arrivals_before_10[i] <- sum(event_times < 10)

# Count the number of arrivals between 11:00 and 13:00
arrivals_11_to_13[i] <- sum(event_times >= 11 & event_times <= 13)
}

# Calculate the average number of arrivals for each interval
average_before_10 <- mean(arrivals_before_10)
average_11_to_13 <- mean(arrivals_11_to_13)

# Print the results
cat("Average number of arrivals before 10:00:", average_before_10, "\n")

## Average number of arrivals before 10:00: 0.615
cat("Average number of arrivals between 11:00 and 13:00:", average_11_to_13, "\n")

## Average number of arrivals between 11:00 and 13:00: 62.4591

```

Problem 03 (a) || R

```
# Number of Monte Carlo samples
n <- 5000

# Define the limits of the integration
x_lower <- 1
x_upper <- 4
y_lower <- -2
y_upper <- 2
z_lower <- 0
z_upper <- 1
w_lower <- 0
w_upper <- 10

# Sampling uniformly for x, y, z, and exponentially for w
x_samples <- runif(n, min = x_lower, max = x_upper)
y_samples <- runif(n, min = y_lower, max = y_upper)
z_samples <- runif(n, min = z_lower, max = z_upper)
w_samples <- rexp(n, rate = 4) # Exponential distribution with rate = 4

# Define the integrand function
integrand <- function(x, y, z, w) {
  (1 / (1 + x^2 + y^2 + z^2)) * exp(-w / 4)
}

# Calculate the value of the integrand at the sampled points
integrand_values <- integrand(x_samples, y_samples, z_samples, w_samples)

# Calculate the volume of the integration region
volume <-
  (x_upper - x_lower) * (y_upper - y_lower) * (z_upper - z_lower) * w_upper

# Estimate the integral using Monte Carlo
integral_estimate <- volume * mean(integrand_values)

# Print the result
cat("Estimated value of the nested integral using Monte Carlo integration:",
    integral_estimate, "\n")
```

```
## Estimated value of the nested integral using Monte Carlo integration: 15.23773
```

PROBLEM 03(b):

We know that, a χ^2 distribution with k -degrees of freedom can be generated as the sum of the squares of k independent standard normal random variables. That is, if Z_1, Z_2, \dots, Z_k are independent and follow a standard normal distribution $N(0,1)$, then

$$\chi_k^2 = Z_1^2 + Z_2^2 + Z_3^2 + Z_4^2 + Z_5^2 + Z_6^2$$

Let, U_1 and U_2 be two independent random variables, that are uniformly distributed on the interval $(0,1)$.

Using Box-Muller transformation,
for Uniform Random Variables U_1 and U_2 ,

$$Z_1 = \sqrt{-2 \ln(U_1)} \cos(2\pi U_2) ; \quad Z_2 = \sqrt{-2 \ln(U_1)} \sin(2\pi U_2)$$

for Uniform Random Variables U_3 and U_4 ,

$$Z_3 = \sqrt{-2 \ln(U_3)} \cdot \cos(2\pi U_4) ; \quad Z_4 = \sqrt{-2 \ln(U_3)} \cdot \sin(2\pi U_4)$$

for Uniform Random Variables U_5 and U_6

$$Z_5 = \sqrt{-2 \ln(U_5)} \cdot \cos(2\pi U_6) ; \quad Z_6 = \sqrt{-2 \ln(U_5)} \cdot \sin(2\pi U_6)$$

Now,

$$\begin{aligned} \chi_6^2 &= Z_1^2 + Z_2^2 + Z_3^2 + Z_4^2 + Z_5^2 + Z_6^2 \\ &= [-2 \ln(U_1)] \{ \cos^2(2\pi U_2) + \sin^2(2\pi U_2) \} + \\ &\quad [-2 \ln(U_3)] \{ \cos^2(2\pi U_4) + \sin^2(2\pi U_4) \} + \\ &\quad [-2 \ln(U_5)] \{ \cos^2(2\pi U_6) + \sin^2(2\pi U_6) \} \\ &= -2 \ln(U_1) - 2 \ln(U_3) - 2 \ln(U_5) \\ &= -2 (\ln U_1 + \ln U_3 + \ln U_5) \end{aligned}$$

Problem 04 (a) || R

```
# Install and load the "shapes" package if not already installed
if (!require(shapes)) {
  install.packages("shapes")
  library(shapes)
}
```

```
## Loading required package: shapes
```

```
## Warning: package 'shapes' was built under R version 4.3.3
```

```
# Load the female data
```

```
load("C:/Users/USER/Documents/panf.dat.rda")
```

```
female_data <- get("panf.dat")
```

```
# Load the male data
```

```
load("C:/Users/USER/Documents/panm.dat.rda")
```

```
male_data <- get("panm.dat")
```

```
# Plot the data for female and male chimpanzees using plotshapes
```

```
par(mfrow = c(1, 2))
```

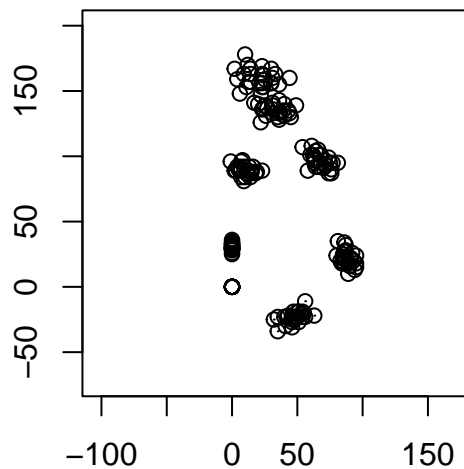
```
plotshapes(female_data)
```

```
title("Female Chimpanzee Skull")
```

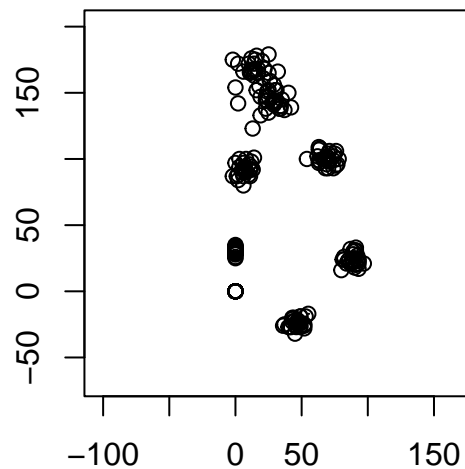
```
plotshapes(male_data)
```

```
title("Male Chimpanzee Skull")
```

Female Chimpanzee Skull



Male Chimpanzee Skull



Problem 04 (b) || R

```
# Calculate the centroid size for female chimpanzees
female_centroid_sizes <- apply(female_data, 3, centroid.size)
mean_female_centroid <- mean(female_centroid_sizes)

# Calculate the centroid size for male chimpanzees
male_centroid_sizes <- apply(male_data, 3, centroid.size)
mean_male_centroid <- mean(male_centroid_sizes)

# Print the mean centroid sizes
cat("Mean Centroid Size for Female Chimpanzees:", mean_female_centroid,
    "and Male Chimpanzees:", mean_male_centroid)

## Mean Centroid Size for Female Chimpanzees: 196.0349 and Male Chimpanzees: 203.1697

# Perform bootstrap sampling with B = 5,000
B <- 5000

# Bootstrap mean centroid size for female chimpanzees
bootstrap_female <- replicate(B, {
  resample_indices <- sample(1:length(female_centroid_sizes), replace = TRUE)
  mean(female_centroid_sizes[resample_indices])
})

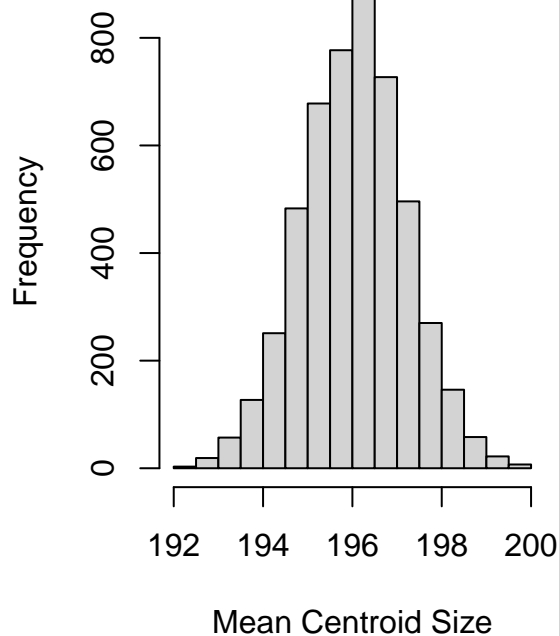
# Bootstrap mean centroid size for male chimpanzees
bootstrap_male <- replicate(B, {
  resample_indices <- sample(1:length(male_centroid_sizes), replace = TRUE)
  mean(male_centroid_sizes[resample_indices])
})

# Print the bootstrap estimates for the means
cat("Bootstrap Mean for Female Chimpanzees:", mean(bootstrap_female),
    "and Male Chimpanzees:", mean(bootstrap_male))

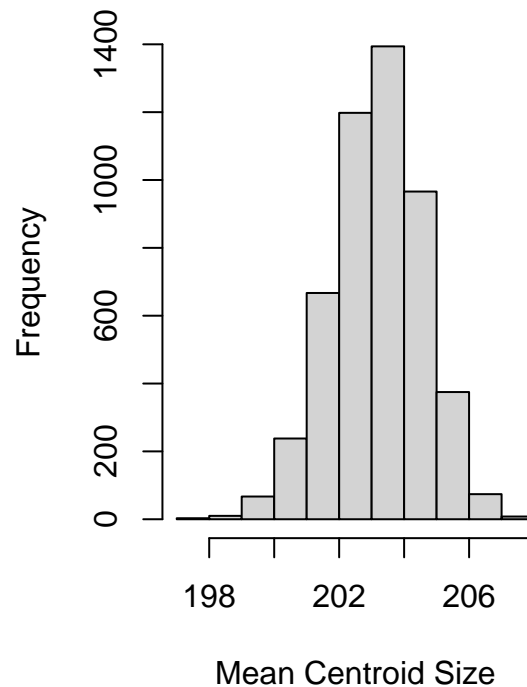
## Bootstrap Mean for Female Chimpanzees: 196.0345 and Male Chimpanzees: 203.1747

# Plot the bootstrap distributions
par(mfrow = c(1, 2))
hist(bootstrap_female,
     main = "Bootstrap Distribution - Female", xlab = "Mean Centroid Size")
hist(bootstrap_male,
     main = "Bootstrap Distribution - Male", xlab = "Mean Centroid Size")
```

Bootstrap Distribution – Female



Bootstrap Distribution – Male



Problem 04 (c) || R

```
# Calculate the 95% percentile bootstrap confidence intervals
ci_female <- quantile(bootstrap_female, probs = c(0.025, 0.975))
ci_male <- quantile(bootstrap_male, probs = c(0.025, 0.975))

# Print the confidence intervals
cat("95% Percentile Bootstrap Confidence Interval for Female Mean Centroid Size:", ci_female)

## 95% Percentile Bootstrap Confidence Interval for Female Mean Centroid Size: 193.7075 198.3498
cat("95% Percentile Bootstrap Confidence Interval for Male Mean Centroid Size:", ci_male)

## 95% Percentile Bootstrap Confidence Interval for Male Mean Centroid Size: 200.2601 205.8049

# Interpretation of the Results
if (ci_female[2] < ci_male[1] || ci_male[2] < ci_female[1]) {
  cat("The confidence intervals for female and male mean centroid sizes do not overlap,
      suggesting a significant difference in skull size between female and male chimpanzees.")
} else {
  cat("The confidence intervals for female and male mean centroid sizes overlap, suggesting
      no clear significant difference in skull size between female and male chimpanzees.")
}

## The confidence intervals for female and male mean centroid sizes do not overlap,
##      suggesting a significant difference in skull size between female and male chimpanzees.
```