HUDM6026 Homework_02

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Question 01 SCR 3.3

MY SOLUTION:

The inverse transformation of the Pareto(a,b)'s cdf function is as followed.

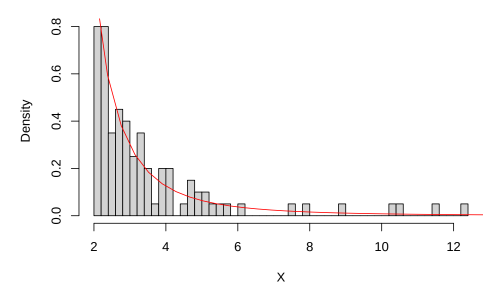
$$F^{-1}(u) = \frac{b}{(1-u)^{\frac{1}{a}}}$$

This inverse function runs well. Before comparing the simulated density and the original density, I derivate the CDF to get the pdf function of Pareto(a,b), that is:

$$f(x) = \frac{ab^a}{x^{a+1}}$$

Figure 1. Comparing the simulated data with Pareto(a,b)

$$f(x) = ab^a/x^{(a+1)}$$



Question 02 SCR 3.9

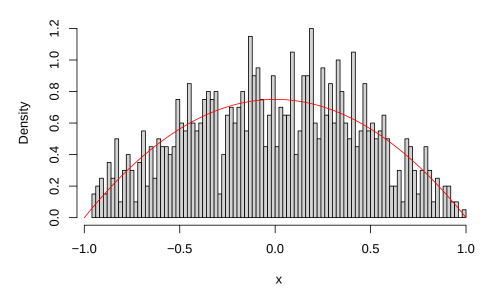
MY SOLUTION:

This question has already given the clues to generate random variable for the rescaled Epanechnikov kernel

```
> # write a function based on text's information
> gen_var <- function(n){ # n is the sample size</pre>
    U_1 <- runif(n, -1, 1)
    U_2 <- runif(n, -1, 1)
    U_3 <- runif(n, -1, 1)
    U_output <- c()</pre>
    for (i in c(1:n)) {
      if (abs(U_3[i]) > abs(U_2[i]) &
          abs(U_3[i]) > abs(U_1[i]))
        {U_output[i] <- U_2[i]}
+
        {U_output[i] <- U_3[i]}
    }
+
    return(U_output)
>
> # generate 1000 data
> U_output <- gen_var(1000)
> hist(U_output, prob = T,
       breaks = 100,
       xlab = "x",
      main = expression(f(x) == (3/4)*(1-x^2)))
> x_{vec} < - seq(-1,1,0.001)
> lines(x_vec, f_x, col="red")
```

Figure 2. Rescaled Epanechnikov kernel Distribution

$$f(x) = (3/4)(1-x^2)$$



Question 03 SCR 3.11

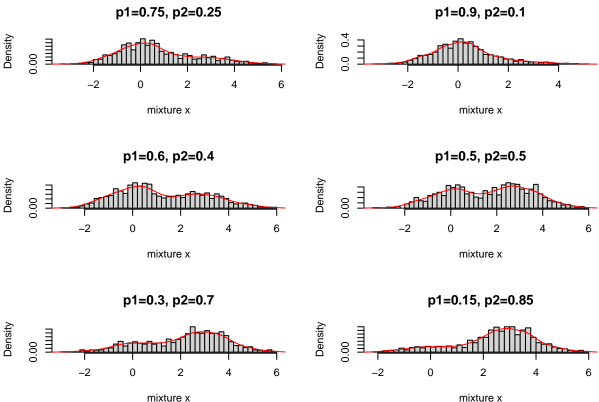
MY SOLUTION:

How to better understand the mixing weights (i.e., the mixing probabilities)? The mixing weights is about **how much** each individual distribution contributes to the mixture distribution (Stephanie Glen. StatisticsHowTo.com). Therefore, when constructing the mixture function, one should not directly use the probability of each parent distribution as a coefficient!!

```
> set.seed(1000)
> n <- 1000
> # generate two vectors from normal distribution
> x1 <- rnorm(n,0,1)
> x2 < rnorm(n,3,1)
> # use a for-loop to draw graphs at different mixing weights
> par(mfrow=c(3,2))
> for (p1 in c(0.75, 0.90, 0.60, 0.5, 0.30, 0.15)){}
    # define the mixing prob
    p2 <- 1 - p1
    # use n data from uniform distribution to construct
    # the proportion of each parent distribution.
    u <- runif(n)
    k <- as.integer(u > p2)
    x \leftarrow k * x1 + (1-k) * x2
    hist(x, prob = T,
```

```
+ breaks = 50,
+ xlab = "mixture x",
+ main = sprintf("p1=%s, p2=%s", p1, p2))
+ lines(density(x), col= "red")
+ }
> mtext("Figure 3. Mixture Distributions With Different Mixing Weights",
+ side = 3,
+ line = -1,
+ outer = T)
```

Figure 3. Mixture Distributions With Different Mixing Weights



From the graphs, one can find that when the mixing weights are .5 and .5, the mixture distribution is apparently a bimodal distribution. At this circumstance the two samples contribute equally to the final mixture. But this bimodal distribution might not be symmetrical because the two parent distribution's shapes are different in variance. With increasing in P1, the bimodal distribution will be more positively skewed.

Question 04 SCR 3.14

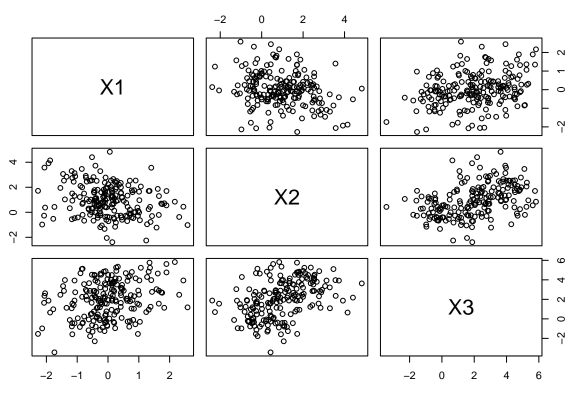
MY SOLUTION:

For solving this question, I refer to Prof.Keller's in-class demo codes and remove unnecessary if-else expressions.

```
> # create a function to gen data matrix from a multivariate normal distribution.
> mvn_gen <- function(n, mu, sigma){
+ # to determine the dimension</pre>
```

```
d <- length(mu)
   # generate a n*d matrix from a standard normal distribution
   Z <- matrix(rnorm(n*d), nrow = n, ncol = d)</pre>
   # use Cholesky function to factorize the Cov-var matrix
   Q <- chol(sigma)
   # to transform the mu from dataframe to matrix
    mu <- matrix (mu, nrow = d, ncol = 1)</pre>
+ # J is column vector of ones
+ J <- matrix(mu, nrow = n, ncol = 1)
   # define the output X matrix
+ X <- Z %*% Q + J %*% t(mu)
   return(data.frame(X))
+ }
>
> # input the given cov-var matrix
> sig \leftarrow matrix(c(1.0, -0.5, 0.5,
                  -0.5, 1.0, -0.5,
+
                  0.5, -0.5, 1.0), 3, 3)
> set.seed(100)
> X \leftarrow mvn_gen(n=200, mu = c(0,1,2), sigma = sig)
> pairs(X)
> mtext("Figure 4. Generate Data From Multivariate Normal Distribution",
       side = 3,
        line = -1,
        outer = T)
> head(X)
          X1
                      X2
1 -0.50219235 0.275493640 -1.3082869
2\quad 0.13153117\ 0.625320206\ 2.0529727
3 -0.07891709 2.777854646 4.8891723
4 0.88678481 0.001194942 -0.5540068
5 0.11697127 1.823304027 0.9285956
6 0.31863009 0.956058119 5.3480697
```





From the cov-var matrix, one can easily get the correlation coefficient between the x_1 and x_2 is -0.5. The middle-right graph demonstrated that these two variables are negatively correlated. By visually check, the joint mean point is at around (0,1), which agrees with the given condition. The bottom-right graph also meets the given condition. However, the correlation between the x_2 and x_3 is obscure by visual check.