# Final Project Part 1

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# **Metropolis-Hastings**

## 1. Algorithm and main function

#### Algorithm

- 1. Choose a starting value  $\theta^{(0)}$
- 2. At iteration t, compute the jumping distribution  $J_t(\theta^*|\theta^{(t-1)})$
- 3. Computing an acceptance ratio

$$r = \frac{p(\theta')/J(\theta')}{p(\theta)/J(\theta)}$$

Here we split into two part: Posterior =  $\frac{L(\phi')}{L(\phi)}$ , Proposal =  $\frac{L'(\phi)}{L'(\phi')}$ r = Posterior \* Proposal

- 4. Accept  $\theta^*$  as  $\theta^{(t)}$  with probability min(r,1). If  $\theta^*$  is not accepted, then  $\theta^{(t)} = \theta^{(t-1)}$
- 5. Repeat step 2-4 Iteration times to get samples, and then apply burn-in or thinning

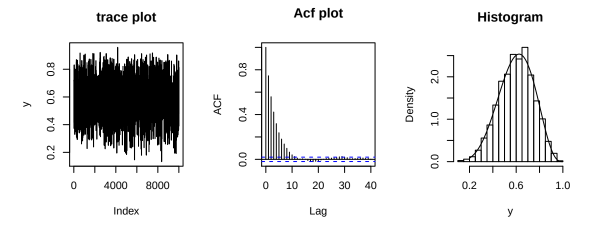
#### Main function

```
Beta_MetropolisHastings = function(true_alpha, true_beta, c, iter){
  set.seed(1)
  # start value random from a uniform distribution
  start value = runif(1); chain = rep(0, iter)
  # the proposal function given in the problem
  proposalfunction = function(c, phi_old){return(rbeta(1, c * phi_old, c * (1 - phi_old)))}
  for (i in 1 : iter){
    # generate a bata sample
   beta = proposalfunction(c, start_value)
    # compute the posterior and propsal value from generated beta
   post = dbeta(beta, true_alpha, true_beta) / dbeta(start_value, true_alpha, true_beta)
   proposal = dbeta(start_value, c * beta, c * (1 - beta)) / dbeta(beta, c * start_value,
                                                                     c * (1 - start_value))
    # if it's in the accept field, add beta into chain, otherwise,
    # use the same value with last element
   if(runif(1) < min(1, post * proposal)){</pre>
      start_value = beta
   }
    chain[i] = start_value
  return(chain)
```

## 2. Evaluate the performance of the sampler

```
Variables: \alpha = 6, \beta = 4, c = 1, iteration = 10000
```

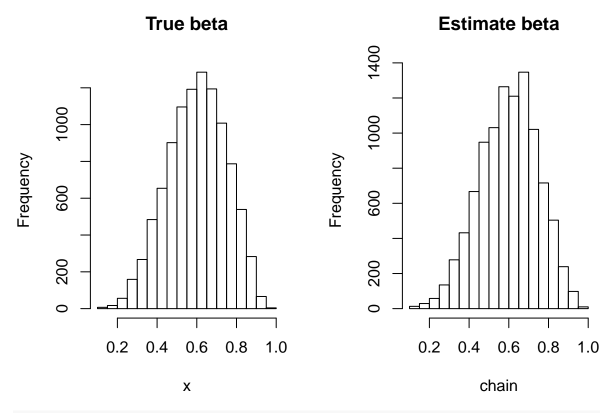
```
set.seed(1)
true_alpha = 6; true_beta = 4; c = 1; iter = 10000
chain = Beta_MetropolisHastings(true_alpha = 6, true_beta = 4, c = 1, iter = 10000)
par(mfrow = c(1, 3)) #1 row, 3 columns
plot(chain, type = "l", main = "trace plot", ylab = "y")
acf(chain, main = "Acf plot")
hist(chain, freq = FALSE, main = "Histogram", xlab = "y")
grid = seq(0.2,1,length = 300)
beta_density = dbeta(grid, true_alpha, true_beta)
lines(grid,beta_density)
```



### Comparision

In the histogram plot, I change the histogram from frequency to density and also add the true beta (6,4) density line in the graph. It seems that the generated samples follow a beta distribution. The following graph shows the true beta distribution with emprical density.

```
x = rbeta(iter, 6, 4)
par(mfrow = c(1, 2))
hist(x, main = "True beta")
hist(chain, main = "Estimate beta")
```



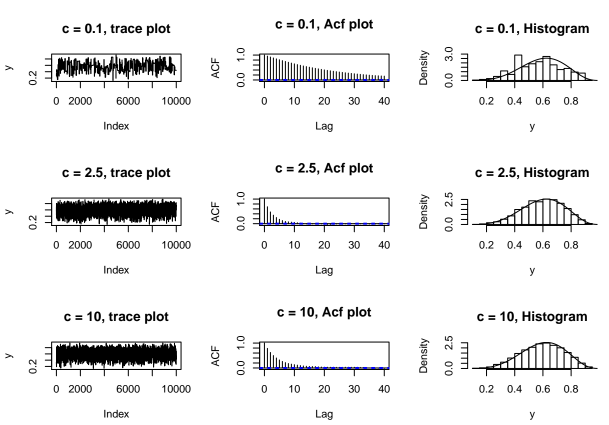
```
ks.test(chain[-(1:5000)], x)
```

```
##
## Two-sample Kolmogorov-Smirnov test
##
## data: chain[-(1:5000)] and x
## D = 0.0186, p-value = 0.199
## alternative hypothesis: two-sided
```

Burin the first 5000 samples, and compare the random data from beta(6,4). The Kolmogorov-Smirnov test show that the p-value is greater than 0.05, which means that we can NOT reject the null hypothesis. Then we can conclude that data may follow beta distribution.

### 3. Re-run the sample with more parameter

```
c1 = 0.1; c2 = 2.5; c3 = 10
chain1 = Beta_MetropolisHastings(true_alpha = 6, true_beta = 4, c = c1, iter = 10000)
chain2 = Beta_MetropolisHastings(true_alpha = 6, true_beta = 4, c = c2, iter = 10000)
chain3 = Beta MetropolisHastings(true alpha = 6, true beta = 4, c = c3, iter = 10000)
grid = seq(0.2,1,length = 300)
beta density = dbeta(grid, true alpha, true beta)
par(mfrow = c(3, 3))
plot(chain1, type = "l",main = "c = 0.1, trace plot", ylab = "y")
acf(chain1, main = "c = 0.1, Acf plot")
hist(chain1, freq = FALSE, main = "c = 0.1, Histogram", xlab = "y")
lines(grid, beta_density)
plot(chain2, type = "l",main = "c = 2.5, trace plot", ylab = "y")
acf(chain2, main = "c = 2.5, Acf plot")
hist(chain2, freq = FALSE, main = "c = 2.5, Histogram", xlab = "y")
lines(grid, beta_density)
plot(chain3, type = "l",main = "c = 10, trace plot", ylab = "y")
acf(chain3, main = "c = 10, Acf plot")
hist(chain3, freq = FALSE, main = "c = 10, Histogram", xlab = "y")
lines(grid, beta_density)
```



#### Analysis 1

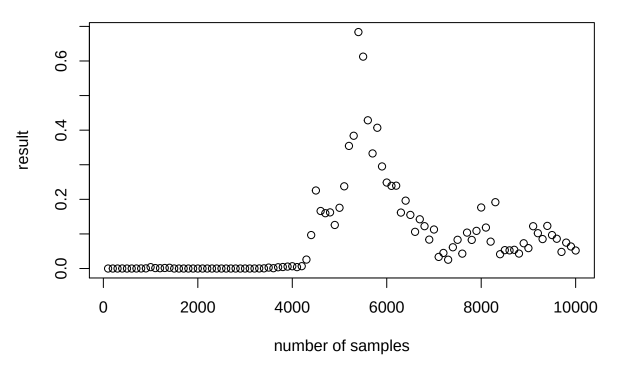
From the trace, Acf plots and histograms, we find that when c=2.5, it has the best fit for beta distribution and least autocorrelation. For smaller c or larger c, it has high autocorrelation, and the samples should be generated independently. Therefore, c=2.5 would be the most effective at drawing from the target distribution.

#### Analysis 2

Consider different number of draws with Kolmogorov-Smirnov test:

```
x = rbeta(iter, 6, 4); result = c()
par(mfrow = c(1, 1))
sample = Beta_MetropolisHastings(true_alpha = 6, true_beta = 4, c = 2.5, iter = 10000)
for(i in 1:100){
    result = c(result, ks.test(sample[1:(i*100)],x)$p.value)
}
plot((1:100)*100,result, main = "p-value", xlab = "number of samples")
```

# p-value



For this certain samples, we could see that when the sample number is greater than 4000, the p-value will be significant not zero and greater than 0.05, and we can NOT reject null hypothesis. The generated samples could be from a beta distribution when the number of draws is greater than 4000.