Computational Statistics - Lab 04

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# Question 1: Computations with Metropolis-Hastings

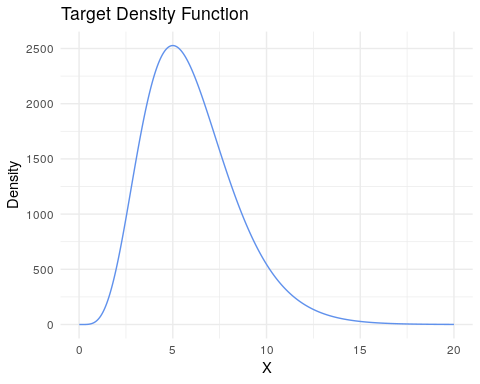
Consider the following probability density function:

You can see that the distribution is known up to some constant of proportionality. If you are interested (**NOT** part of the Lab) this constant can be found by applying integration by parts multiple times and equals 120.

1. Use Metropolis–Hastings algorithm to generate samples from this distribution by using proposal distribution as log–normal , take some starting point. Plot the chain you obtained as a time series plot. What can you guess about the convergence of the chain? If there is a burn–in period, what can be the size of this period?

**Answer:** First we will define the original function with the constant set to get a brief overview how the function actually looks like. We also define the given function without the constant, as we’re going to us it later as the target distribution. We use the function without the constant to show that Metropilis-Hastings works without knowing the constant. As random samplings can actually give an outside of the scope of the function, we will set all values of to a relatively small value.

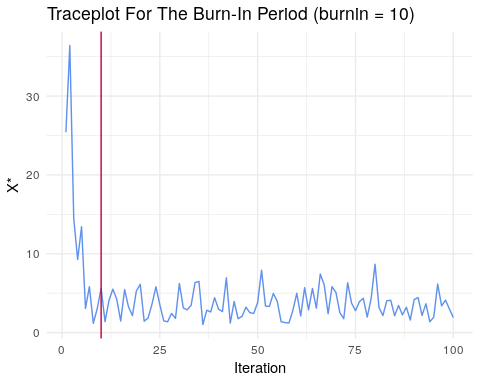
# Target function with original scaling  
f = function(x) {  
 return(120 \* x^5 \* exp(-x))  
}  
  
# Target function  
df = function(x) {  
 if (sum(c(x) <= 0) > 0) x = 0.0001  
 return(x^5 \* exp(-x))  
}



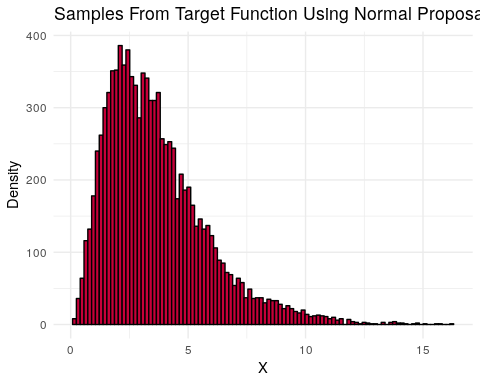
The following code chunk defines the Metropolis-Hastings algorithm. The correction factor c is used to handle the asymmetry in the proposal function. The code is commented.

#' Metropolis Hasting Algorithm  
#'  
#' @param n Number of samples from the target distribution.  
#' @param x\_0 Initial state from Pi.   
#' @param b Burn-in steps to remove from samples.  
#' @param proposal Selects the proposal function.  
#' @param keep\_burnin Decides if to keep the samples during the burn-in period.  
#'  
#' @return Returns a list containing samples.  
#' @export  
#'  
#' @examples  
metropolis\_hastings = function(n = 1, x\_0 = 1, b = 50, proposal = "lnorm",  
 keep\_burnin = FALSE) {  
   
 # Vectors to store the samples in  
 samples = c()  
   
 # Samples from proposel from the random walk (MC)  
 xt = x\_0  
 xt\_1 = x\_0  
   
 while (length(samples) < n) {  
   
 # Generate proposal state  
 if (proposal == "lnorm") {  
 x\_star = rlnorm(n = 1, meanlog = log(xt), sdlog = 1)  
   
 # Calculate correction factor C  
 c = dlnorm(xt\_1, meanlog = xt, sdlog = 1) /  
 dlnorm(x\_star, meanlog = xt, sdlog = 1)  
 }  
 else if (proposal == "chisquared") {  
 x\_star = rchisq(n = 1, df = floor(xt))  
   
 # Calculate correction factor C  
 c = dchisq(x = xt\_1, df = floor(xt)) /  
 dchisq(x\_star, df = floor(xt))  
 }  
 else {  
 stop("Invalid proposal.")  
 }  
   
 # Calculate acceptance probability alpha  
 if (df(xt\_1) <= 0) {  
 alpha = 0  
 }  
 else {  
 alpha = min(1, df(x\_star)/df(xt\_1) \* c)  
 }  
   
 # Generate u from uniform  
 u = runif(n = 1, min = 0, max = 1)  
   
 # Decide if to accept or reject the proposal  
 if (u <= alpha) {  
 # Accept  
 xt\_1 = xt  
 xt = x\_star  
 samples = c(samples, x\_star)  
 }  
 else {  
 # Reject  
 xt = xt\_1  
 }  
 }  
   
 # Return samples  
 if (keep\_burnin) return(samples)  
 return(samples[b+1:length(samples)])  
}

For the lognormal distribution as the proposal we selected a starting point which is far from the converging value to illistrate the burn-in period. We can see that we reach convergence rather quickly after around 8 iterations. We increased the limit to 10 to be sure to get into the area of the converging value.

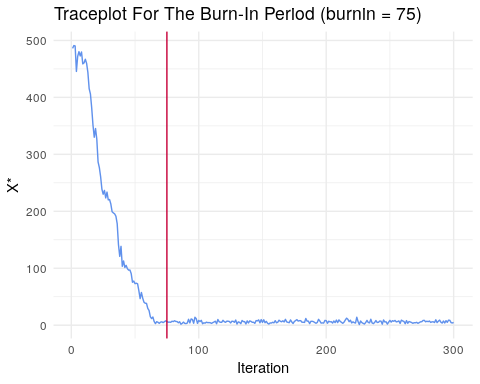


The following plot shows 10.000 samples with as a starting point and burn-in iterations set to 100.

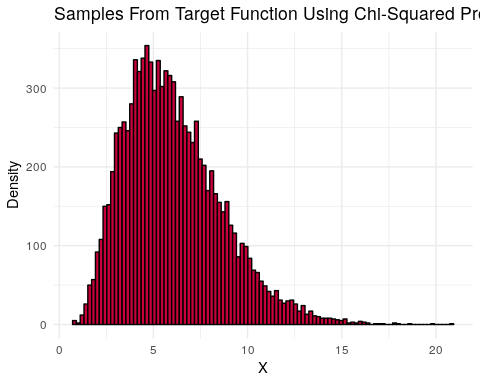


1. Perform Step 1 by using the chi–square distribution as a proposal distribution, where is the floor function, meaning the integer part of for positive , i.e.

**Answer:** The trace plot shows the burn-in period for the chi-square distribution. We see that it takes quite longer to achieve convergence. We set the burn-in iterations to 75 to be sure to converge.



Again you can see 10.000 samples with as a starting point and burn-in iterations set to 75.



1. Compare the results of Steps 1 and 2 and make conclusions.

**Answer:** The samples seem to actually be taken from the real distribution and in this context ist doesn’t matter which proposal distribution we take. Comparing the burn-in times we see that it seems to take longer for the chi-squared proposal to converge. We have to keep in mind that this mostly depends on the initial state of the MCMC. It’s not so easy to take an initial value which is far of from the converging value as we get values around 0 as the functions have a really low density which makes it quite hard to actually calculate c and . As long as we take an initial state not too far off from a low density, we shoudl be fine, as the algorithm automatically avoids really low density regions.

We assume as long as the proposal distribution mostly covers the target distribution and the initial state is selected quite reasonable (which is not always the case in the “real world”) the Metropolis-Hastings works quite well.

1. Generate 10 MCMC sequences using the generator from Step 2 and starting points 1, 2, …, or 10. Use the Gelman–Rubin method to analyze convergence of these sequences.

**Answer:** The following output is based three sets of multiple sequences (with starting points ). The first set of sequences uses 5, the second 10 and the last uses 50 iterations. We can see that the Upper C.I. (at 95% confidence) converges towards 1 which indicates that the set has converged. This works by comparing the variance of the sequences within a set. As more iterations are performed, the MCMC depends less on the starting point until it finally holds no more information about that. This is when the Gelman-Ruby factor is near 1.

**TODO:** Implement also the formulas from the slides.

## Potential scale reduction factors:  
##   
## Point est. Upper C.I.  
## [1,] 1.07 1.85

## Potential scale reduction factors:  
##   
## Point est. Upper C.I.  
## [1,] 1.04 1.19

## Potential scale reduction factors:  
##   
## Point est. Upper C.I.  
## [1,] 1.02 1.06

1. Estimate

using the samples from Steps 1 and 2.

f\_integrate = function(x) {  
 return(x \* df(x))  
}  
  
valid\_samples\_lnorm = results\_lnorm[!is.na(results\_lnorm)]  
  
b = max(valid\_samples\_lnorm)  
a = min(valid\_samples\_lnorm)  
diff = b - a  
  
sum(f\_integrate(valid\_samples\_lnorm) \* diff / df(valid\_samples\_lnorm)) /  
 length(valid\_samples\_lnorm)

## [1] 58.75335

valid\_samples\_chisquared = results\_chisquared[!is.na(results\_chisquared)]  
  
b = max(valid\_samples\_chisquared)  
a = min(valid\_samples\_chisquared)  
diff = b - a  
  
sum(f\_integrate(valid\_samples\_chisquared) \* diff / df(valid\_samples\_chisquared)) /  
 length(valid\_samples\_chisquared)

## [1] 121.3373

**TODO:** The lnorm returns only half of the integral. Check out why (on purpose or if there is a reason).

1. The distribution generated is in fact a gamma distribution. Look in the literature and define the actual value of the integral. Compare it with the one you obtained.

**Answer:** As we know it’s a gamma distribution where the integral over D is 1 and we also know the constant C given from above, we know that the integral has to add upp to .

**TODO:** Check out and write down the real gamma distribution mentioned here.

# Question 2: Gibbs Sampling

concentration of a certain chemical was measured in a water sample, and the result was stored in the data chemical.RData having the following variables:

* day of the measurement
* measured concentration of the chemical.

The instrument used to measure the concentration had certain accuracy; this is why the measurements can be treated as noisy. Your purpose is to restore the expected concentration values.

1. Import the data to R and plot the dependence of Y on X. What kind of model is reasonable to use here?
2. A researcher has decided to use the following (random–walk) Bayesian model (n=numberof observations, are unknown parameters):

where the prior is

Present the formulae showing the likelihood and the prior . **Hint:** a chain rule can be used here

1. Use Bayes’ Theorem to get the posterior up to a constant proportionality, and then find out the distributions of , where is a vector containing all values except of .

Hint A: consider for separate formulae for and then a formula for all remaining .

Hint B:

Hint C:

1. Use the distributions derived in Step 3 to implement a Gibbs sampler that uses as a starting point. Run the Gibbs sampler to obtain 1000 values of and then compute the expected value of versus X and Y versus X in the same graph. Does it seem that you have managed to remove the noise? Does it seem that the expected value of can catch the true underlying dependence between Y and X?
2. Make a trace plot for and comment on the burn–in period and convergence.

# Loading RData  
#data2 = get(load("data.RData"))  
#head(data2)

# Source Code

knitr::opts\_chunk$set(echo = TRUE, cache = FALSE, include = TRUE, eval = TRUE)  
library(knitr)  
library(readxl)  
library(ggplot2)  
library(gridExtra)  
library(coda)  
set.seed(12345)  
  
# Target function with original scaling  
f = function(x) {  
 return(120 \* x^5 \* exp(-x))  
}  
  
# Target function  
df = function(x) {  
 if (sum(c(x) <= 0) > 0) x = 0.0001  
 return(x^5 \* exp(-x))  
}  
  
  
sequence = seq(from = 0.01, to = 20, by = 0.01)  
  
real\_f = f(sequence)  
plotdf = data.frame(sequence, real\_f)  
  
ggplot(plotdf) +  
 geom\_line(aes(x = sequence, y = real\_f), color = "#6091ec") +  
 labs(title = "Target Density Function", y = "Density",  
 x = "X", color = "Legend") +  
 theme\_minimal()  
  
  
#' Metropolis Hasting Algorithm  
#'  
#' @param n Number of samples from the target distribution.  
#' @param x\_0 Initial state from Pi.   
#' @param b Burn-in steps to remove from samples.  
#' @param proposal Selects the proposal function.  
#' @param keep\_burnin Decides if to keep the samples during the burn-in period.  
#'  
#' @return Returns a list containing samples.  
#' @export  
#'  
#' @examples  
metropolis\_hastings = function(n = 1, x\_0 = 1, b = 50, proposal = "lnorm",  
 keep\_burnin = FALSE) {  
   
 # Vectors to store the samples in  
 samples = c()  
   
 # Samples from proposel from the random walk (MC)  
 xt = x\_0  
 xt\_1 = x\_0  
   
 while (length(samples) < n) {  
   
 # Generate proposal state  
 if (proposal == "lnorm") {  
 x\_star = rlnorm(n = 1, meanlog = log(xt), sdlog = 1)  
   
 # Calculate correction factor C  
 c = dlnorm(xt\_1, meanlog = xt, sdlog = 1) /  
 dlnorm(x\_star, meanlog = xt, sdlog = 1)  
 }  
 else if (proposal == "chisquared") {  
 x\_star = rchisq(n = 1, df = floor(xt))  
   
 # Calculate correction factor C  
 c = dchisq(x = xt\_1, df = floor(xt)) /  
 dchisq(x\_star, df = floor(xt))  
 }  
 else {  
 stop("Invalid proposal.")  
 }  
   
 # Calculate acceptance probability alpha  
 if (df(xt\_1) <= 0) {  
 alpha = 0  
 }  
 else {  
 alpha = min(1, df(x\_star)/df(xt\_1) \* c)  
 }  
   
 # Generate u from uniform  
 u = runif(n = 1, min = 0, max = 1)  
   
 # Decide if to accept or reject the proposal  
 if (u <= alpha) {  
 # Accept  
 xt\_1 = xt  
 xt = x\_star  
 samples = c(samples, x\_star)  
 }  
 else {  
 # Reject  
 xt = xt\_1  
 }  
 }  
   
 # Return samples  
 if (keep\_burnin) return(samples)  
 return(samples[b+1:length(samples)])  
}  
  
  
burnin = 10  
  
results = metropolis\_hastings(100, x\_0 = 40, b = burnin, proposal = "lnorm",  
 keep\_burnin = TRUE)  
  
plotdf = data.frame(index = 1:length(results), values = results)  
  
ggplot(plotdf) +  
 geom\_line(aes(x = index, y = values), color = "#6091ec") +  
 labs(title = "Traceplot For The Burn-In Period (burnin = 10)", y = "X\*",  
 x = "Iteration", color = "Legend") +  
 geom\_vline(xintercept = burnin, color = "#C70039") +   
 theme\_minimal()  
  
  
results\_lnorm = metropolis\_hastings(10000, x\_0 = 5, b = 100, proposal = "lnorm")  
  
plotdf = data.frame(results\_lnorm)  
  
ggplot(plotdf) +  
 geom\_histogram(aes(x = results\_lnorm),  
 color = "#000000", fill = "#C70039", bins = length(results\_lnorm)/100) +  
 labs(title = "Samples From Target Function Using Normal Proposal", y = "Density",  
 x = "X", color = "Legend") +  
 theme\_minimal()  
  
  
burnin = 75  
  
results = metropolis\_hastings(300, x\_0 = 500, b = burnin, proposal = "chisquared", keep\_burnin = TRUE)  
  
plotdf = data.frame(index = 1:length(results), values = results)  
  
ggplot(plotdf) +  
 geom\_line(aes(x = index, y = values), color = "#6091ec") +  
 labs(title = "Traceplot For The Burn-In Period (burnin = 75)", y = "X\*",  
 x = "Iteration", color = "Legend") +  
 geom\_vline(xintercept = burnin, color = "#C70039") +   
 theme\_minimal()  
  
  
results\_chisquared = metropolis\_hastings(10000, x\_0 = 5, b = 100, proposal = "chisquared")  
  
plotdf = data.frame(results\_chisquared)  
  
ggplot(plotdf) +  
 geom\_histogram(aes(x = results\_chisquared),  
 color = "#000000", fill = "#C70039", bins = length(results\_chisquared)/100) +  
 labs(title = "Samples From Target Function Using Chi-Squared Proposal", y = "Density",  
 x = "X", color = "Legend") +  
 theme\_minimal()  
  
  
results5 = list()  
results10 = list()  
results50 = list()  
k = 10  
  
for (i in 1:k) {  
 results5[[i]] = mcmc(metropolis\_hastings(5, x\_0 = i, b = 0,  
 proposal = "chisquared", keep\_burnin = TRUE))  
}  
  
for (i in 1:k) {  
 results10[[i]] = mcmc(metropolis\_hastings(10, x\_0 = i, b = 0,  
 proposal = "chisquared", keep\_burnin = TRUE))  
}  
  
for (i in 1:k) {  
 results50[[i]] = mcmc(metropolis\_hastings(50, x\_0 = i, b = 0,  
 proposal = "chisquared", keep\_burnin = TRUE))  
}  
  
mcmc\_list5 = mcmc.list(results5)  
gelman.diag(mcmc\_list5)  
  
mcmc\_list10 = mcmc.list(results10)  
gelman.diag(mcmc\_list10)  
  
mcmc\_list50 = mcmc.list(results50)  
gelman.diag(mcmc\_list50)  
  
  
f\_integrate = function(x) {  
 return(x \* df(x))  
}  
  
valid\_samples\_lnorm = results\_lnorm[!is.na(results\_lnorm)]  
  
b = max(valid\_samples\_lnorm)  
a = min(valid\_samples\_lnorm)  
diff = b - a  
  
sum(f\_integrate(valid\_samples\_lnorm) \* diff / df(valid\_samples\_lnorm)) /  
 length(valid\_samples\_lnorm)  
  
  
  
valid\_samples\_chisquared = results\_chisquared[!is.na(results\_chisquared)]  
  
b = max(valid\_samples\_chisquared)  
a = min(valid\_samples\_chisquared)  
diff = b - a  
  
sum(f\_integrate(valid\_samples\_chisquared) \* diff / df(valid\_samples\_chisquared)) /  
 length(valid\_samples\_chisquared)  
  
  
# Loading RData  
#data2 = get(load("data.RData"))  
#head(data2)