# Applying the common random number technique as a Markov chain convergence diagnostic

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The following code generated the graphs and calculations found in "Applying the common random number technique as a Markov chain convergence diagnostic."

#### Bayesian Regression Gibbs Sampler

Numerical example

```
# Data is obtained from the general linear models textbook by Dobson
#Table 6.3 Carbohydrate, age, relative weight and protein for twenty male insulin dependent diabetics;
df <- c(33, 33, 100, 14,
40, 47, 92, 15,
37, 49, 135, 18,
27, 35, 144, 12,
30, 46, 140, 15,
43, 52, 101, 15,
34, 62, 95, 14,
48, 23, 101, 17,
30, 32, 98, 15,
38, 42, 105, 14,
50, 31, 108, 17,
51, 61, 85, 19,
30, 63, 130, 19,
36, 40, 127, 20,
41, 50, 109, 15,
42, 64, 107, 16,
46, 56, 117, 18,
24, 61, 100, 13,
35, 48, 118, 18,
37, 28, 102, 14)
df <- matrix(df, nrow = 20, byrow=TRUE)</pre>
Y <- df[,1]
X \leftarrow cbind(1,df[,-1])
n <- length(Y)
p \leftarrow dim(X)[2]
df <- data.frame(df)</pre>
colnames(df) <- c("carbs", "age", "weight", "protein")</pre>
```

```
# setting priors: beta ~ N(b_0,E_0) sigma^2 ~ Inv-Chi^2(v_0, c_0)
b_0 <- rep(0,4)
E_0 <- diag(4)
v_0 <- 1
c_0 <- 6
```

Consistent with equation (16) we define

$$g(\beta, \sigma^2) = \frac{1}{(\sigma^2)^{(n+\nu_0)/2+1}} \exp\left(-\frac{1}{2\sigma^2}(y - X\beta)^T(y - X\beta) - \frac{1}{2}(\beta - \beta_0)^T \Sigma_{\beta}^{-1}(\beta - \beta_0) - \frac{\nu_0 c_0^2}{\sigma^2}\right)$$

We further define

## [1] 0.971513

$$f(\beta, \sigma^2) = \log(g(\beta, \sigma^2))$$

```
# g(x,y) = g(beta,sigma^2) as defined in equation 6
g <- function(x){
b <- c(x[1], x[2], x[3], x[4])
o <- x[5]
z = exp(-((n+v_0)/2+1)*log(o) - 1/(2*o)*t(Y-(X %*% b)) %*% (Y-(X %*% b)) - 0.5*t(b-b_0) %*% inv(E_0) return(z)}

f <- function(x){
b <- c(x[1], x[2], x[3], x[4])
o <- x[5]
z = -((n+v_0)/2+1)*log(o) - 1/(2*o)*t(Y-(X %*% b)) %*% (Y-(X %*% b)) - 0.5*t(b-b_0) %*% inv(E_0) %*% return(-z)
}</pre>
```

We want to find a lower bound (L) on  $\int_{\mathbb{R}^4 \times \mathbb{R}} g(\beta, \sigma^2) d(\beta, \sigma^2)$ . To do so, we apply the following,

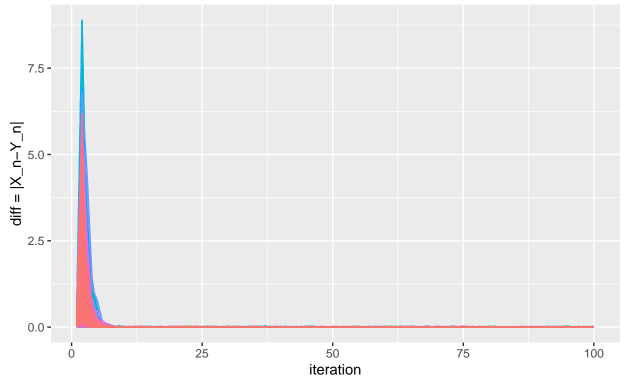
```
\int_{\mathbb{R}^4\times\mathbb{R}}g(\beta,\sigma^2)d(\beta,\sigma^2)\geq \int_Cg(\beta,\sigma^2)d(\beta,\sigma^2)\geq e^{\int_Cf(\beta,\sigma^2)d(\beta,\sigma^2)}
x \leftarrow c(0.1, 0.1, 0.1, 0.1, 0.1)
x \leftarrow optim(x, f)par
#c(0.1,0.1,0.1,0.1,0.1)
#c(0.2,0.2,0.2,0.2,0.2)
int_f \leftarrow adaptIntegrate(f, lowerLimit = x-c(0.1,0.1,0.1,0.1,0.1), upperLimit = x+c(0.1,0.1,0.1,0.1,0.1)
int_f
## $integral
## [1] 0.02890064
##
## $error
## [1] 1.018664e-13
## $functionEvaluations
## [1] 93
## $returnCode
## [1] 0
L <- exp(-int_f$integral)</pre>
```

```
alpha <- (n+v_0)/2
beta <- v_0*c_0/2
K <- 1/L*gamma(alpha)/beta^alpha</pre>
## [1] 11.40551
tvbound <- (n+v_0)/(v_0*c_0)
tvbound
## [1] 3.5
The following is function that generates \sigma_{n+1}^2 given \sigma_n^2, equation 7.
b_hat <- inv(t(X) %*% X) %*% t(X) %*% Y
invE_0 \leftarrow inv(E_0)
nextIt <- function(o, Z, G){</pre>
  eigenV \leftarrow eigen(t(X) %*% X/o + inv(E_0))
  Q <- eigenV$vectors
  L <- diag(eigenV$values)</pre>
  Vinv12 <- Q %*% inv(sqrt(L)) %*% inv(Q)</pre>
  Vinv \leftarrow Q \%*\% inv(L) \%*\% inv(Q)
  b_tilde <- Vinv %*% (t(X) %*% X %*% b_hat/o + inv(E_0) %*% b_0)
  W <- X %*% b_tilde - Y + X %*% Vinv12 %*% Z
  o1 <- (v_0*c_0/2 +t(W) %*% W/(2*G))
  return(o1)
}
Now we apply the common random number technique to generate an estimate of E[X_k - Y_k], N = 100
I = 1000 \text{ and } X_0, Y_0 \sim \Gamma^{-1}(\alpha', \beta') = \Gamma^{-1}(10.5, 2)
I = 1000
J = 100
diff <- matrix(0, I, J)</pre>
for(i in 1:I){
  it <- matrix(0, ncol=2, nrow=J)</pre>
  it[1,] <- 1/rgamma(2, shape = alpha, rate = beta)
  for(j in 2:J){
    Z \leftarrow rnorm(p, 0, 1)
    G <- rgamma(1, shape = alpha, rate =1)
    it[j,1] <- nextIt(it[j-1,1], Z, G)
    it[j,2] \leftarrow nextIt(it[j-1,2], Z, G)
  }
  diff[i,] <- abs(it[,1]-it[,2])
}
diff_df <- data.frame(t(diff), iter_no = 1:J)</pre>
diff_df <- diff_df %>%
  pivot_longer(cols = starts_with("X"), names_to = "sim_no", values_to = "val")
diff_df %>%
  ggplot(aes(x = iter_no, y = val)) +
  geom_line(aes(color = sim_no)) + theme(legend.position = "none") +
  labs(title = "1000 simulations of |X_n-Y_n|", subtitle = "Using common random number technique") +
```

xlab("iteration") + ylab("diff = |X\_n-Y\_n|")

## 1000 simulations of $|X_n-Y_n|$

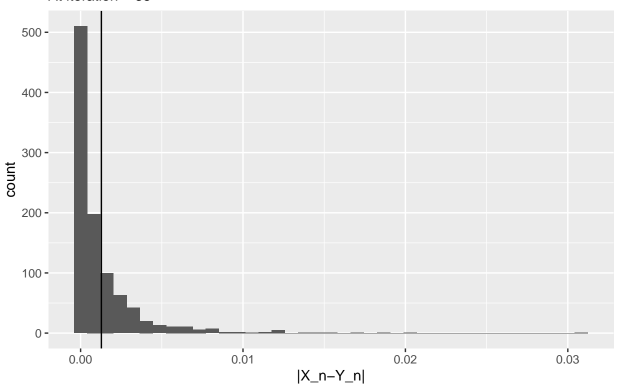
Using common random number technique



```
expdiff <- diff_df %>%
  filter(iter_no==35) %>%
  summarise(expdiff = mean(val))
expdiff
## # A tibble: 1 x 1
##
     expdiff
       <dbl>
##
## 1 0.00128
diff_df %>%
  filter(iter_no==35) %>%
  ggplot(aes(x=val)) +
  geom_histogram(bins = "39") + labs(title = "Histogram of |X_n-Y_n|", subtitle = "At iteration = 35")
  xlab("|X_n-Y_n|") +
  geom_vline(aes(xintercept = expdiff$expdiff), colour="black")
```

### Histogram of |X\_n-Y\_n|

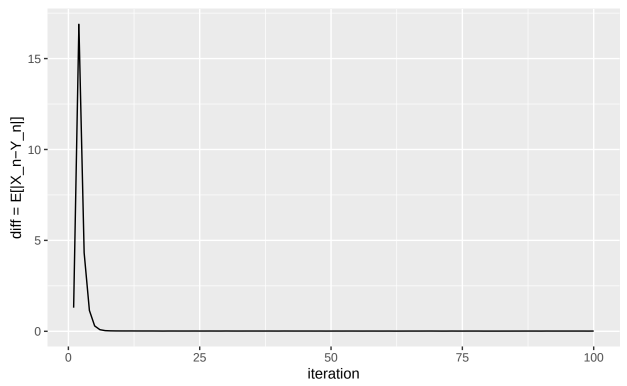




#### #geom\_vline(xintercept = expdiff)

```
diff_df %>%
  group_by(iter_no) %>%
  summarise(mean_val = mean(val)*K) %>%
  ggplot(aes(x = iter_no, y = mean_val)) +
  geom_line() + theme(legend.position = "none") +
  labs(title = "Sample average value of |X_n-Y_n|", subtitle = "Based on 1000 simulations") +
  xlab("iteration") + ylab("diff = E[|X_n-Y_n|]")
```

# Sample average value of |X\_n-Y\_n| Based on 1000 simulations



```
diff_df %>%
  group_by(iter_no) %>%
  summarise(mean_val = mean(val)*K) %>%
  filter(iter_no==35)
## # A tibble: 1 x 2
##
     iter_no mean_val
##
       <int>
                <dbl>
## 1
          35
               0.0146
diff_df %>%
  group_by(iter_no) %>%
  summarise(mean_val = mean(val)*K*tvbound) %>%
 filter(iter_no==35)
## # A tibble: 1 x 2
     iter_no mean_val
                <dbl>
##
       <int>
## 1
          35
               0.0511
```

#### Autoregressive process

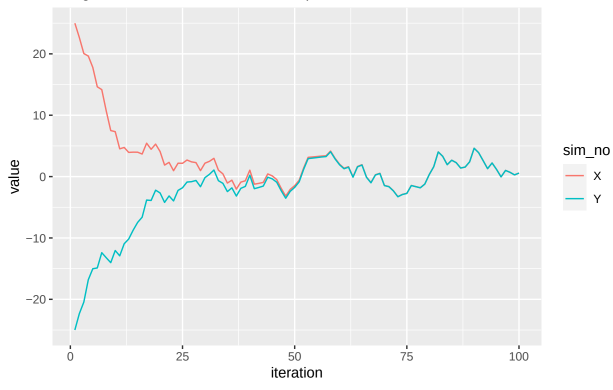
Define the autorregressive process to be

$$X_n = 0.9X_{n-1} + Z_n, Z_n \sim N(0, 1)$$

```
X <- 1:100
Y <- X
Z \leftarrow rnorm(100, 0, 1)
X[1] < -25
Y[1] <- -25
for(i in 2:100){
  X[i] \leftarrow 0.9*X[i-1]+Z[i]
  Y[i] \leftarrow 0.9*Y[i-1]+Z[i]
}
x_forwards <- X</pre>
df <- data.frame(X,Y, iter=1:100)</pre>
df %>%
  pivot_longer(X:Y, names_to = "sim_no", values_to = "val") %>%
  ggplot(aes(x = iter, y = val)) +
  geom_line(aes(color = sim_no)) + labs(title = "Simulations of X_n and Y_n", subtitle = "Using common :
  xlab("iteration") + ylab("value")
```

#### Simulations of X\_n and Y\_n

Using common random number technique



```
n <- 100
X <- 1:n
Y <- 1:n
#Z <- rnorm(n, 0,1)

X[1] <- 25
Y[1] <- 25</pre>
```

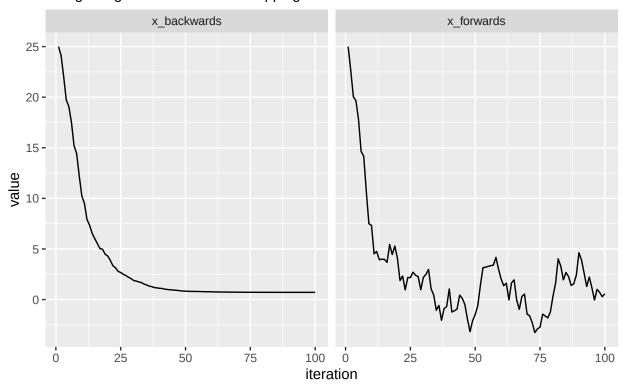
```
for(i in 2:n){
  for(j in 2:i){
    Y[j] <- 0.9*Y[j-1]+Z[i+1-j]
  }
  X[i] <- Y[i]
}

x_backwards <- X

df <- data.frame(x_forwards,x_backwards, iter=1:100)
  df %>%
  pivot_longer(x_forwards:x_backwards, names_to = "sim_no", values_to = "val") %>%
  ggplot(aes(x = iter, y = val)) +
  geom_line() + facet_wrap(vars(sim_no)) + labs(title = "Backwards process vs forwards process", subtit xlab("iteration") + ylab("value")
```

### Backwards process vs forwards process

Using using the same random mappings



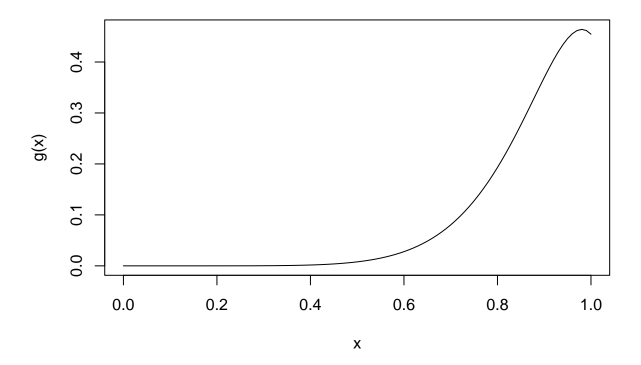
#### Metropolis Algorithm

The unnormalised target density is defined as follows,

$$g(x) = x^3 \sin(x^4) \cos(x^5) I_{x \in [0,1]}$$

```
g <- function(y){
  if ((0<=y) && (y<=1)){
    h <- y^3*sin(y^4)*cos(y^5)
}</pre>
```

```
else {
    h <- 0
}
return(h)
}
g_norestrict <- function(y){
    h <- y^3*sin(y^4)*cos(y^5)
    return(h)
}
curve(g_norestrict, from=0, to=1, xlab="x", ylab="g(x)")</pre>
```

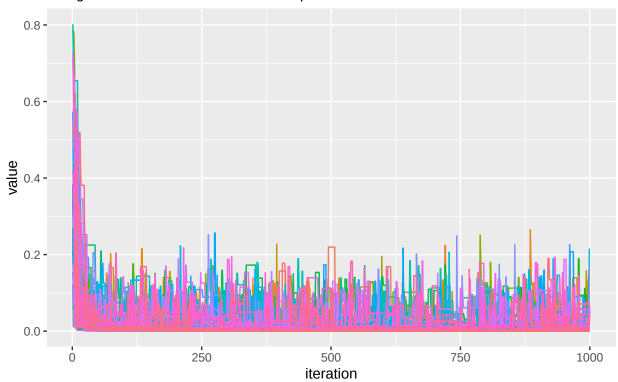


```
n <- 10000
x <- rep(0, n)
y <- x
z <- x
x[1] = 0.7
y[1] = 0.2

acceptreject <- function(u, a, x, x_prop){
  if(u < a) {
    x_next <- x_prop
  }
  else {
    x_next <- x
}</pre>
```

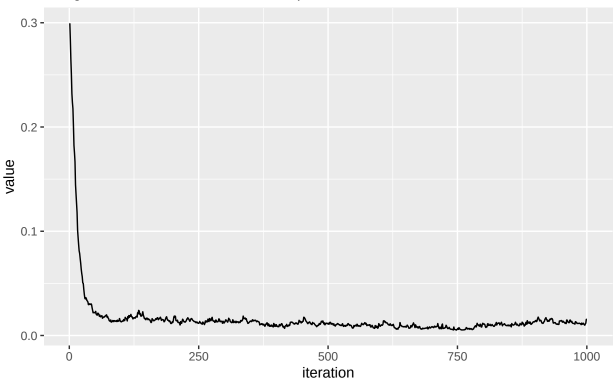
```
getsim_crn <- function(n, i){</pre>
  x <- 1:n
  y <- 1:n
  x[1] \leftarrow runif(1,0,1)
  y[1] \leftarrow runif(1,0,1)
  for (j in 2:n){
    z <- rnorm(1)
    u <- runif(1)
    x_{prop} = x[j-1] + 0.1*z
    y_{prop} = y[j-1] + 0.1*z
   A_x = g(x_prop)/g(x[j-1])
    A_y = g(y_prop)/g(y[j-1])
    x[j] = acceptreject(u, A_x, x[j-1], x_prop)
    y[j] = acceptreject(u, A_y, y[j-1], y_prop)
  df <- data.frame(sim_no = as.character(i), iter = 1:n, x, y, val = abs(x-y))</pre>
  return(df)
}
I <- 100
N <- 1000
df <- getsim_crn(N, 1)</pre>
for(i in 2:I){
  df <- rbind(df, getsim_crn(N, i))</pre>
}
df %>%
  ggplot(aes(x = iter, y = val)) +
  geom_line(aes(color = sim_no)) + labs(title = "Simulations of |X_n - Y_n|", subtitle = "Using common :
 xlab("iteration") + ylab("value") + theme(legend.position = "none")
```

# Simulations of |X\_n - Y\_n| Using common random number technique



```
df %>%
  group_by(iter) %>%
  summarise(mval = mean(val))%>%
  ggplot(aes(x = iter, y = mval)) +
  geom_line() + labs(title = "Mean |X_n-Y_n|", subtitle = "Using common random number technique") +
  xlab("iteration") + ylab("value")
```

Mean |X\_n-Y\_n|
Using common random number technique



#### Example of the set A

```
eq = function(x){cos(2*x*pi)}
eq2 = function(x){cos(x*pi)}
ggplot(data.frame(x=c(0, 2)), aes(x=x)) +
    stat_function(fun=eq, xlim=c(0,0.5), linewidth=1) +
    stat_function(fun=eq2, xlim=c(0,0.5), linewidth=1) +
    stat_function(fun=eq, xlim=c(0.5,1.5), linewidth=0.5) +
    stat_function(fun=eq2, xlim=c(0.5,1.5), linewidth=0.5) +
    stat_function(fun=eq2, xlim=c(1.5,2), linewidth=1) +
    stat_function(fun=eq2, xlim=c(1.5,2), linewidth=1) + theme_classic() +
    labs(x="theta", y="value of f")
```

