Lab3 report

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Q1 Gibbs sampler for a normal model

(a Implement (code!) a Gibbs sampler that simulates from the joint posterior

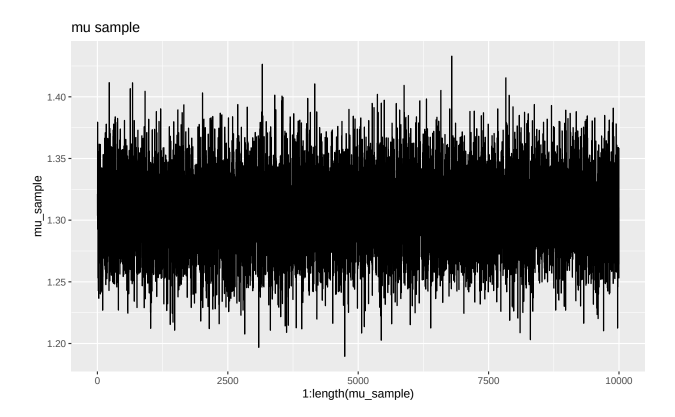
```
#####Q1
library(ggplot2)
library(mvtnorm)
#library(extraDistr)
precipitation <- readRDS("Precipitation.rds")</pre>
log_prec <- log(precipitation)</pre>
mu_post <- function(mu_0,tau_0_sqr,sigma_square_current,y,n){</pre>
  tau_n_sqr = 1 / (1/tau_0_sqr + n/sigma_square_current)
  mu_n <- tau_n_sqr * (mu_0/tau_0_sqr + sum(y)/sigma_square_current)</pre>
  mupost <- rnorm(1, mu_n, sqrt(tau_n_sqr))</pre>
  return(mupost)
}
sigma_square_post <- function(mu_current, v_0, sigma_square_0, y,n, use_myinvchi=TRUE) {</pre>
  v_n \leftarrow v_0 + n
  elem1 <- v_0*sigma_square_0
  elem2 <- sum((y - mu_current)^2)</pre>
  elem3 \leftarrow n+v_0
  elem_comb <- (elem1+elem2)/elem3</pre>
  if (use_myinvchi){
    sigma_square_post <- my_inv_chi(v_n,elem_comb)</pre>
  }
  else {
    sigma_square_post <- rinvchisq(1, v_n, elem_comb) #Requires package extraDistr</pre>
  return(sigma_square_post)
my_inv_chi<- function(df,tau_sqr) {</pre>
 X <- rchisq(1,df)</pre>
```

```
inv_chi <- (df*tau_sqr)/X</pre>
  return(inv_chi)
}
#init
mu 0 <- 0
tau_0_sqr <- 1
sigma_square_0 <- 1
v_0 \leftarrow 1 #degree of freedom for chi square
gibbs_sampler <- function(nstep, data, mu_0, tau_0_sqr, v_0, sigma_square_0) {
  # Init parameters
  mu_current <- 0
  sigma_square_current <- 1</pre>
  mu_samples <- rep(0,nstep)</pre>
  sigma_square_samples <- rep(0,nstep)</pre>
  for (i in 1:nstep) {
    mu_current <- mu_post(mu_0, tau_0_sqr, sigma_square_current, y=data, length(data))</pre>
    #print(mu_current)
    sigma_square_current <- sigma_square_post(mu_current, v_0, sigma_square_0, y=data,</pre>
                                             length(data),use_myinvchi=TRUE)
    #print(sigma square current)
    mu_samples[i] <- mu_current</pre>
    sigma_square_samples[i] <- sigma_square_current</pre>
  output_df <- data.frame(mu_sample = mu_samples, sigma_sample = sigma_square_samples)</pre>
  return(output_df)
}
sample_gibbs <- gibbs_sampler(nstep=10000, data=log_prec, mu_0, tau_0_sqr, v_0, sigma_square_0)</pre>
In lecture slide IF = 1 + 2\sum_{k=1}^{\infty} \rho_k where \rho_k is autocorrelation at lag k
my_acf <- acf(sample_gibbs$mu_sample,plot = F) # getting the autocorrelation</pre>
if_mu <- 1 + 2 * sum(my_acf$acf[-1])
my_acf <- acf(sample_gibbs$sigma_sample,plot = F)</pre>
if_sigma \leftarrow 1 + 2 * sum(my_acf_acf[-1])
```

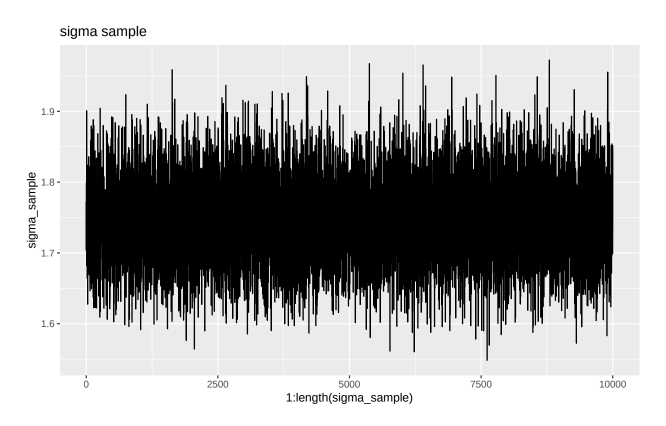
The the Inefficiency Factors (IF) of $\mu = 0.9982076$ and IF of $\sigma = 0.9522093$. The value of IF is around 1 indicates that the chain has converged to the stationary distribution.

Below, we present two trace plot that also confirm the samples' convergence.

```
ggplot(data=sample_gibbs, aes(x = 1:length(mu_sample), y = mu_sample)) +
  geom_line()+labs(title = "mu_sample")
```



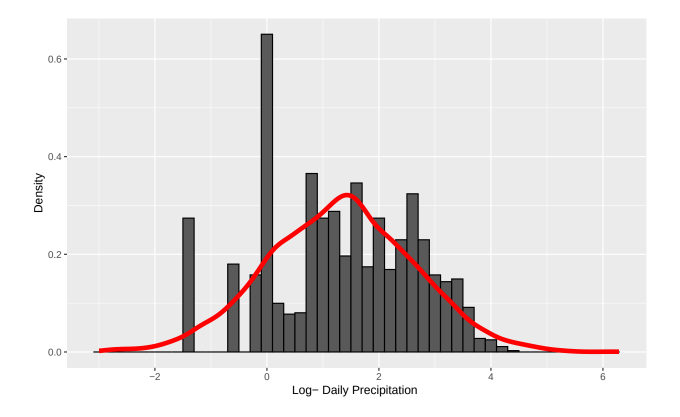
```
ggplot(data=sample_gibbs, aes(x = 1:length(sigma_sample), y = sigma_sample)) +
geom_line() +labs(title = "sigma_sample")
```



(b)

Plot the following in one figure:

- 1) a histogram or kernel density estimate of the daily precipitation
- 2) The resulting posterior predictive density using the simulated posterior draws from How well does the posterior predictive density agree with this data?



Above is the plot of the histogram of the daily precipitation, and the resulting posterior predictive density (red line). We can say that the posterior prediction does not totally agree with the existing data, especially towards to left-hand side of the mode.

Metropolis Random Walk for Poisson regression

#####Q2#####

##

##

AIC: 3610.3

a) Obtain the maximum likelihood estimator of β in the Poisson regression model for the eBay data [Hint: glm.R, don't forget that glm() adds its own intercept so don't input the covariate Const] Which covariates are significant?

```
ebay_data <- read.table("eBayNumberOfBidderData.dat", header = TRUE)</pre>
model1 <- glm(formula = nBids ~ .,data = ebay_data[,-2],family = 'poisson')</pre>
summary(model1)
##
## Call:
## glm(formula = nBids ~ ., family = "poisson", data = ebay_data[,
       -2])
##
##
## Deviance Residuals:
##
       Min
                 10
                      Median
                                   30
                                           Max
## -3.5800 -0.7222 -0.0441
                               0.5269
                                        2.4605
##
## Coefficients:
               Estimate Std. Error z value Pr(>|z|)
##
                           0.03077 34.848 < 2e-16 ***
## (Intercept) 1.07244
## PowerSeller -0.02054
                           0.03678
                                    -0.558
                                             0.5765
## VerifyID
               -0.39452
                           0.09243
                                    -4.268 1.97e-05 ***
## Sealed
                           0.05056
                                     8.778 < 2e-16 ***
                0.44384
## Minblem
               -0.05220
                           0.06020 -0.867
                                             0.3859
## MajBlem
               -0.22087
                           0.09144 - 2.416
                                             0.0157 *
## LargNeg
                0.07067
                           0.05633
                                     1.255
                                             0.2096
## LogBook
               -0.12068
                           0.02896 -4.166 3.09e-05 ***
## MinBidShare -1.89410
                           0.07124 -26.588 < 2e-16 ***
## Signif. codes: 0 '*** 0.001 '** 0.01 '* 0.05 '.' 0.1 ' 1
##
## (Dispersion parameter for poisson family taken to be 1)
```

Null deviance: 2151.28 on 999 degrees of freedom

Residual deviance: 867.47 on 991 degrees of freedom

Number of Fisher Scoring iterations: 5

The estimates of beta are 1.0724421, -0.0205408, -0.3945165, 0.4438426, -0.0521983, -0.2208712, 0.0706725, -0.1206776, -1.8940966

The significant covariates are 'VerfiyID', 'Sealed', 'LogBook' and 'MinBidShare' and 'Intercept'! MajBlem' is also significant but to a less degree compared to the ones mentioned before.

b) Let's do a Bayesian analysis of the Poisson regression. Let the prior be $\beta \sim \mathcal{N}[0, 100 \cdot (X^TX)^{-1}]$, where X is the n × p covariate matrix. This is a commonly used prior, which is called Zellner's g-prior. Assume first that the posterior density is approximately multivariate normal:

$$\beta \mid y \sim \mathcal{N}\left(\tilde{\beta}, J_{\mathbf{y}}^{-1}(\tilde{\beta})\right)$$

where $\tilde{\beta}$ is the posterior mode and $J_{\mathbf{y}}(\tilde{\beta})$ is the negative Hessian at the posterior mode. $\tilde{\beta}$ and $J_{\mathbf{y}}(\tilde{\beta})$ can be obtained by numerical optimization (optim.R) exactly like you already did for the logistic regression in Lab 2 (but with the log posterior function replaced by the corresponding one for the Poisson model, which you have to code up.).

The Poisson distribution density is given by:

$$p(y \mid x; \beta) = \frac{e^{y\beta'x}e^{-e^{\beta'}}}{y!}$$

The log likelihood of the poisson distribution is :

$$\ell(\beta \mid X, Y) = \log L(\beta \mid X, Y) = \sum_{i=1}^{n} \left(y\beta' x - e^{\beta' x} - \log y! \right)$$

```
# Initialize values
n cols <- ncol(ebay data[,-1])</pre>
#remove 1st column since that is target variable and convert to matrix
# matrix of features
covariates <- as.matrix(ebay data[,-1])
labels <- as.matrix(ebay data[,1])</pre>
mu \leftarrow rep(0, n_cols)
initVal <- matrix(0, n_cols, 1)</pre>
Sigma <- as.matrix(100 * solve(t(covariates)\\'*\'\covariates))</pre>
LogPosteriorFunc <- function(betas, X, y, mu, Sigma){</pre>
  log_prior <- dmvnorm(betas, mu, Sigma, log=TRUE)</pre>
  log_likelihood <- sum(X%*%betas * y - exp(X%*%betas) -log(factorial(y)))</pre>
  res <- log_prior + log_likelihood
  return(res)
}
# Optimizer
OptimRes <- optim(initVal, LogPosteriorFunc, gr = NULL, y = labels, X = covariates,
                   mu = mu, Sigma = Sigma, method=c("BFGS"),
                   control=list(fnscale=-1), hessian=TRUE)
beta_mode <- OptimRes$par</pre>
jacobian <- OptimRes$hessian
inv_jacobian <- -solve(jacobian)</pre>
beta_draws <- as.matrix(rmvnorm(10000, mean = beta_mode, sigma = inv_jacobian))
beta_estimate <- colMeans(beta_draws)</pre>
```

```
hist(beta_draws, breaks = 50, main = 'Histogram of Posterior Draws', xlab = 'Betas')
```

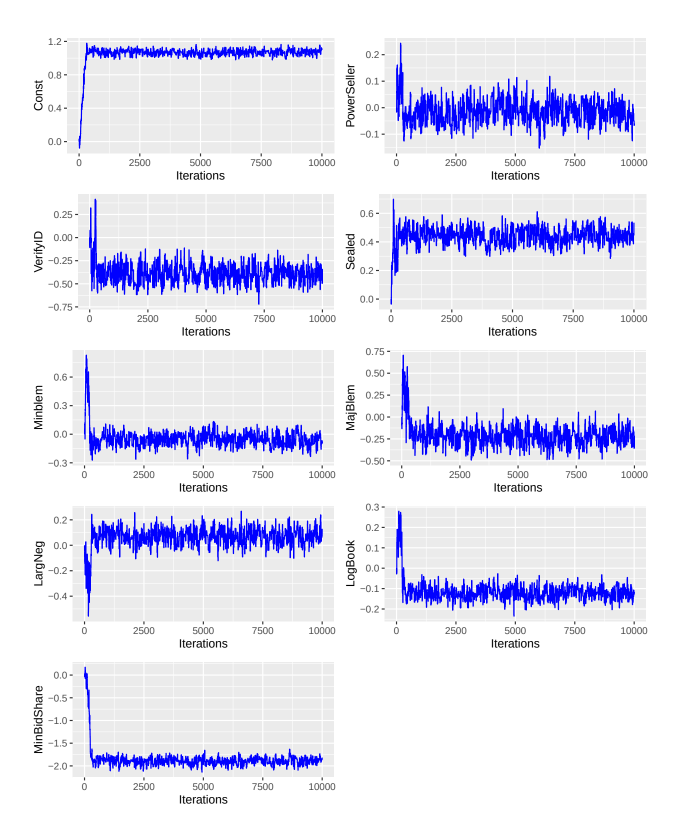
c) Let's simulate from the actual posterior of β using the Metropolis algorithm and compare the results with the approximate results in b). Program a general function that uses the Metropolis algorithm to generate random draws from an arbitrary posterior density. In order to show that it is a general function for any model, we denote the vector of model parameters by θ . Let the proposal density be the multivariate normal density mentioned in Lecture 8 (random walk Metropolis):

$$\theta_p \mid \theta^{(i-1)} \sim N\left(\theta^{(i-1)}, c \cdot \Sigma\right)$$

where $\Sigma = J_y^{-1}(\tilde{\beta})$ was obtained in b). The value c is a tuning parameter and should be an input to your Metropolis function. The user of your Metropolis function should be able to supply her own posterior density function, not necessarily for the Poisson regression, and still be able to use your Metropolis function. This is not so straightforward, unless you have come across function objects in R. The note HowToCo deRWM.pdf in Lisam describes how you can do this in R. Now, use your new Metropolis function to sample from the posterior of β in the Poisson regression for the eBay dataset. Assess MCMC convergence by graphical methods.

```
MetHas_RandomWalk <- function(nDraws,fun,mu,Sigma,c){</pre>
  #initialize matrix
  draw_matrix <- matrix(0,nrow = nDraws,ncol = n_cols)</pre>
  #initialize first row to mu
  draw_matrix[1,] <- mu</pre>
  for(i in 2:nDraws){
    # sample from multivariate normal distribution
    proposed_sample <- as.vector(rmvnorm(n = 1,mean = draw_matrix[i-1,],</pre>
                                            sigma = c*as.matrix(Sigma)))
    #print(proposed_sample)
    # IMPORTANT : the log is inside the posterior function
    log_acceptance_prob <- exp(fun(proposed_sample) - fun(draw_matrix[i-1,]))</pre>
    #random sample
    u <- runif(1)
    # calculate acceptance probability
    a <- min(1,log acceptance prob)
    if(u \le a){
      #accept sample
      draw_matrix[i,] <- proposed_sample</pre>
    }
    else{
      # stay at same values from previous draw
      draw_matrix[i,] <- draw_matrix[i-1,]</pre>
    }
  }
  return(draw_matrix)
# this function we pass to MetHas Algorithm, can be changed to another posterior density
logPostFunc <- function(theta){</pre>
  res <- dmvnorm(theta,mean = beta_estimate,sigma = inv_jacobian,log = TRUE)</pre>
```

```
if(is.na(res)){
    print(theta)
 return(res)
}
df <- MetHas_RandomWalk(nDraws = 10000,fun = logPostFunc,mu = rep(0,n_cols),</pre>
                         Sigma = inv_jacobian,c = 1)
# assign colnames
colnames(df) <- colnames(ebay_data)[2:10]</pre>
## plotting
plot_list <- list()</pre>
for (col in colnames(df)) {
 # Plot iterations vs every column
 p <- ggplot(data = as.data.frame(df), aes_string(x = 1:nrow(df), y = col)) +</pre>
    geom_line(col = 'blue') +
    labs(x = "Iterations", y = col)
  #show plot
  #print(p)
 plot_list[[col]] <- p</pre>
# Arranging in 1 fig
grid.arrange(grobs = plot_list, ncol = 2)
```



Assessing the plots of the 9 column variables vs Iterations, we see that convergence usually occurs after 1500 iterations. (burn-in period)

- d) Use the MCMC draws from c) to simulate from the predictive distribution of the number of bidders in a new auction with the characteristics below. Plot the predictive distribution. What is the probability of no bidders in this new auction?
 - Power Seller = 1
 - VerifyID = 0
 - Sealed = 1
 - MinBlem = 0
 - MajBlem = 1
 - LargNeg = 0
 - Log Book = 1.2
 - MinBidShare = 0.8

$$Mean = \lambda = e^{\beta' \cdot x}$$

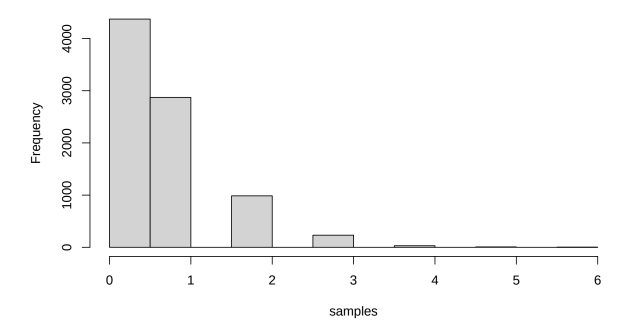
```
# extra 1 at the start for the intercept - Const
new_data <- c(1,1,0,1,0,1,0,1.2,0.8)

# lambda = e^Beta*x
# discarding first 1500 samples as burn-in period
lambda <- exp(df[-c(1:1500),] %*% new_data)

samples <- c()

for (i in 1:nrow(df[-c(1:1500),])) {
    #sample from each row of df to get the predictive distribution based on the
    #posterior betas
    samples[i] <- rpois(1,lambda = lambda[i])
}
hist(samples)</pre>
```

Histogram of samples



res <- length(samples[samples == 0])/length(samples)</pre>

The probability of have zero bidders in the new auction is 51.4352941%.

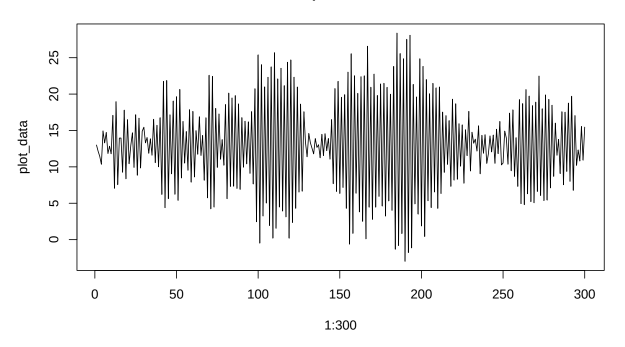
Q3 Time series models in Stan

(a)

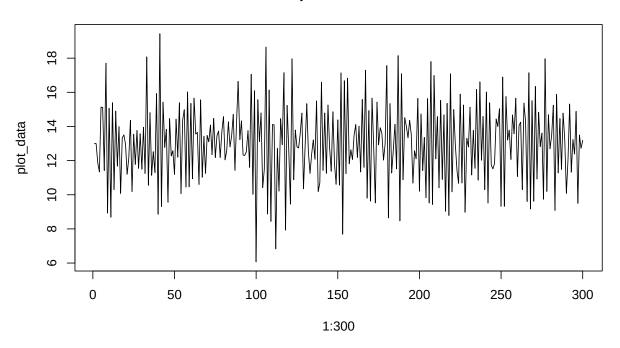
Write a function in R that simulates data from the AR(1)-process

```
#####Q3
mu <- 13
sigma_square <- 3
t <- 300
phi \leftarrow seq(from = -1, to = 1, by =0.25)
ar_func <- function(phi,mu,sigma_square,t){</pre>
  counter <- 0
  result_vector <- rep(0,t)</pre>
  result_vector[1] <- mu</pre>
  for(i in 2:t){
    epislon <- rnorm(1,0,sqrt(sigma_square))</pre>
    x_i \leftarrow mu+phi*(result_vector[i-1]-mu)+ epislon
    result_vector[i] <- x_i
  }
  return(result_vector)
test_phi_func <- function(phi,mu,sigma_square,t){</pre>
  phi_test_df <- data.frame(matrix(0, nrow = t, ncol = length(phi)))</pre>
  colnames(phi_test_df) <- phi</pre>
  for (j in 1:length(phi)) {
    phi_test <- ar_func(phi[j], mu, sigma_square, t)</pre>
    phi_test_df[, j] <- phi_test</pre>
  return(phi_test_df)
phi_df <- test_phi_func(phi,mu,sigma_square,t)</pre>
for(k in 1:length(phi)){
  plot_data <- phi_df[,k]</pre>
  plot(x=1:300, plot_data, type = "l", main = paste(" phi = ", phi[k]))
  #Sys.sleep(1)
```

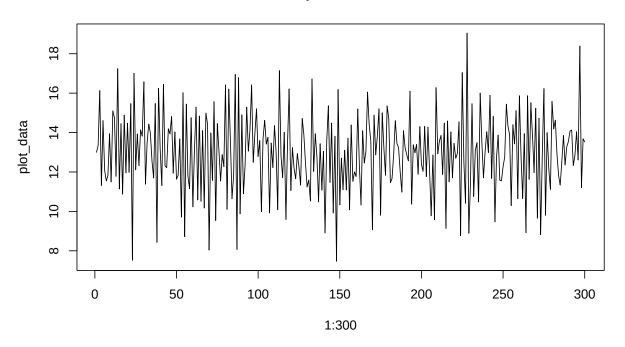
phi = −1



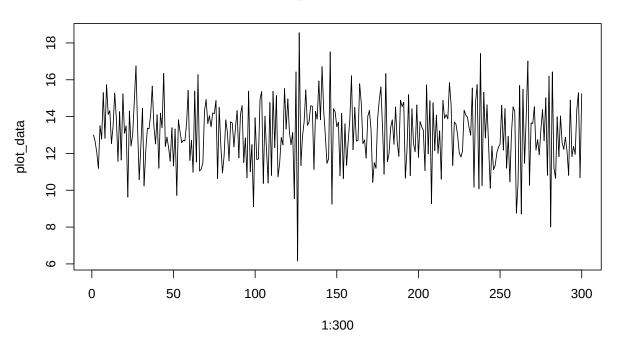
phi = -0.75

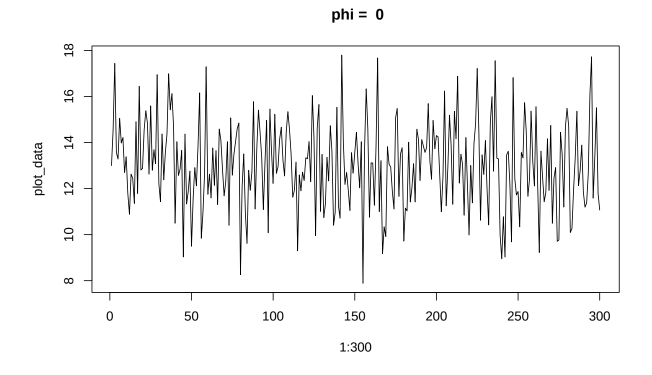


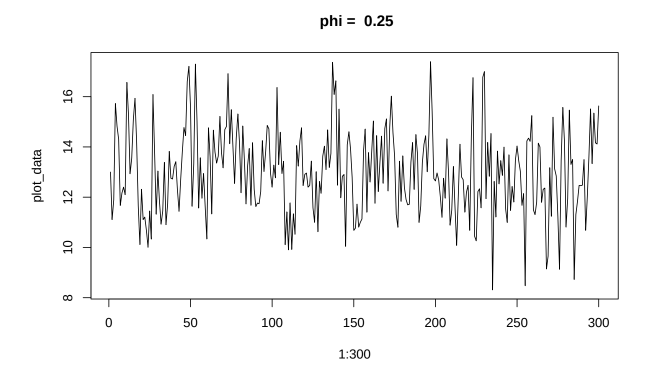
phi = -0.5



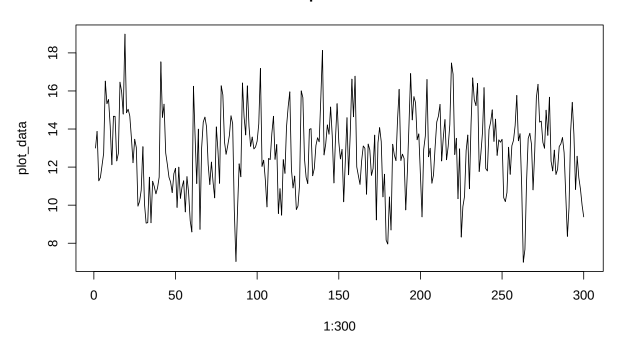




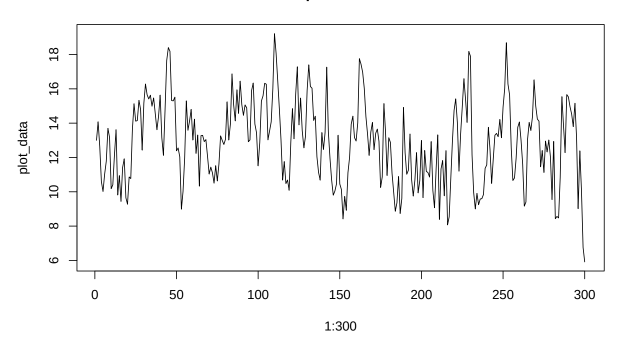




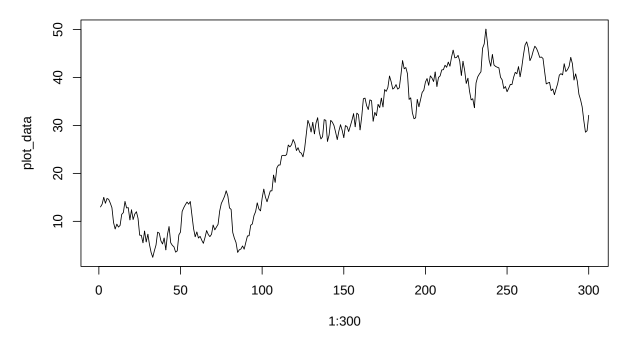
phi = 0.5











As we can observed with different ϕ , as ϕ increase, the gap between peaks are getting larger (not as fluctuate as lower ϕ). This indicate that the data tends to have higher autocorrelation, which means each data point is more correlated with preceding data.

(b

Rstan

```
library(rstan)
set.seed(12345)
x <- ar_func(phi=0.2, mu, sigma_square, t)</pre>
y <- ar_func(phi=0.95, mu, sigma_square, t)
stan_code <- "
data {
  int<lower=0> T;
                           // Number of time points
  vector[T] x;
  vector[T] y;
parameters {
  real mu_x;
  real mu_y;
  real phi_x;
  real phi_y;
  real sigma_x;
  real sigma_y;
}
```

```
model {
  // After some research, it is common to use a flat prior or a vague prior as
  // non-informative prior
  // Therefore, we pick normal distribution with higher variance as prior.
  mu_x ~ normal(0, 50);
  mu_y ~ normal(0, 50);
  phi_x ~ normal(0, 10);
  phi_y ~ normal(0, 10);
  sigma_x ~ normal(0, 50);
  sigma_y ~ normal(0, 50);
 x[2:T] \sim normal(mu_x + phi_x * (x[1:(T - 1)] - mu_x), sigma_x);
 y[2:T] \sim normal(mu_y + phi_y * (y[1:(T - 1)] - mu_y), sigma_y);
}
data_list <- list(</pre>
 T = t,
 x = x
 y = y
# Set the MCMC settings
niter <- 5000
warmup <- 500
# Compile the Stan model
model <- stan_model(model_code = stan_code)</pre>
# Fit the Stan model to the data
# Set the control to avoid too many divergent after warmup
control <- list(adapt_delta = 0.90, stepsize = 0.0001)</pre>
fit <- sampling (model, data = data_list, warmup = warmup, iter = niter, chains = 5, control=control, refre
#refresh = 0 to mute the printout
summary(fit)$summary
##
                  mean
                            se_mean
                                            sd
                                                        2.5%
                                                                     25%
            13.1736454 0.0009810772 0.12894646
                                               12.92120195
                                                              13.0866009
## mu_x
            14.7662220 0.4216001425 9.66646731 -12.59440997
## mu_y
                                                              12.9360194
             0.1886121 0.0004666932 0.05768479
## phi_x
                                                0.07426121
                                                               0.1492954
## phi_y
             0.9713154 0.0003255917 0.01843371
                                                  0.93472438
                                                               0.9583243
## sigma_x
           1.7892259 0.0005562130 0.07331174
                                               1.65097259
                                                               1.7390022
## sigma_y 1.7228587 0.0006095451 0.07117800
                                               1.58966456
                                                               1.6741621
          -634.2138672 0.0264929349 1.83419194 -638.47628106 -635.3056310
## lp__
##
                   50%
                                75%
                                           97.5%
                                                      n_eff
                                                               Rhat
## mu x
            13.1734372
                        ## mu y
            15.5310107 18.0162790 34.1507642
                                                   525.6953 1.006105
                                     0.2991747 15277.7597 1.000192
## phi_x
             0.1889787
                         0.2277933
```

1.0021845 3205.3759 1.003303

phi_y

0.9713980

0.9854067

```
## sigma_x   1.7867845   1.8374897   1.9404140 17372.5969 1.000122
## sigma_y   1.7208153   1.7689548   1.8698400 13635.7782 1.000495
## lp__   -633.9305759 -632.8203154 -631.5496630   4793.2427 1.001257

post_mean <- summary(fit)$summary[, "mean"]
interval_025 <- summary(fit)$summary[, "25%"]
interval_975 <- summary(fit)$summary[, "97.5%"]
n_eff <- summary(fit)$summary[, "n_eff"]
Rhat <- summary(fit)$summary[, "Rhat"]</pre>
```

(i) Based on the summary, we obtain:

```
Posterior mean:
```

```
##
                                      phi_x
           mu x
                         mu_y
                                                    phi_y
                                                                sigma_x
                                                                              sigma_y
##
     13.1736454
                   14.7662220
                                  0.1886121
                                                0.9713154
                                                              1.7892259
                                                                            1.7228587
##
           lp__
## -634.2138672
```

95% credible interval are 2.5%:

```
##
           mu_x
                         mu_y
                                      phi_x
                                                    phi_y
                                                                sigma_x
                                                                              sigma_y
##
     13.0866009
                   12.9360194
                                  0.1492954
                                                0.9583243
                                                              1.7390022
                                                                            1.6741621
##
           lp__
## -635.3056310
```

97.5%:

```
sigma_x
##
           mu_x
                         mu_y
                                      phi_x
                                                    phi_y
                                                                              sigma_y
##
     13.4247425
                   34.1507642
                                  0.2991747
                                                1.0021845
                                                              1.9404140
                                                                            1.8698400
           lp__
## -631.5496630
```

number of effective posterior sample:

```
## mu_x mu_y phi_x phi_y sigma_x sigma_y lp__
## 17274.7783 525.6953 15277.7597 3205.3759 17372.5969 13635.7782 4793.2427
```

```
The true values are \mu_x = \mu_y = 13, \phi_y = 0.95, \phi_x = 0.2, \sigma_x = \sigma_y = \sqrt{3} \approx 1.73
```

For posterior mean, all the inferred parameter are very close to their true value. Hence we can say that we are able to estimate the true value. In terms of 95% credible interval and number of effective posterior sample, we would say that for estimated μ_x , ϕ_y , σ_x and σ_y has narrower credible intervals, while estimated μ_x , ϕ_x , σ_x and σ_y has larger number of effective posterior samples. This indicates the estimate of these inferred parameters are more precise.

(ii) For each of the two data sets, evaluate the convergence of the samplers and plot the joint posterior of μ and ϕ . Comments?

We can use Rhat to evaluate the convergence. The Rhat are as below:

```
## mu_x mu_y phi_x phi_y sigma_x sigma_y lp__
## 1.000210 1.006105 1.000192 1.003303 1.000122 1.000495 1.001257
```

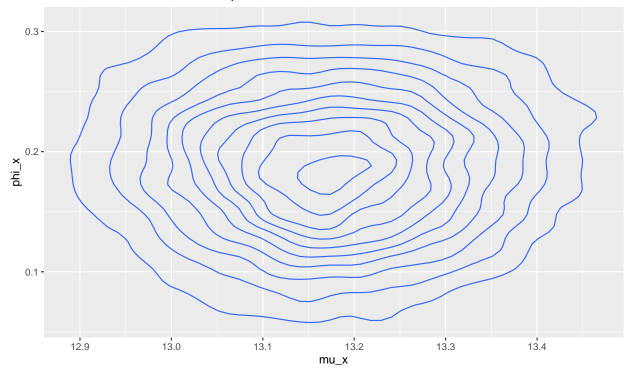
Since all inferred parameters has Rhat close to 1, this imply our sampler converged well.

Below are the plot of the joint posterior of μ and ϕ

For x, the shape of joint distribution are rather symmetric along two axis, this means μ and ϕ has higher correlation. For y, the different shape means, mu and phi has lower correlation hence make it harder to understand the relationship.

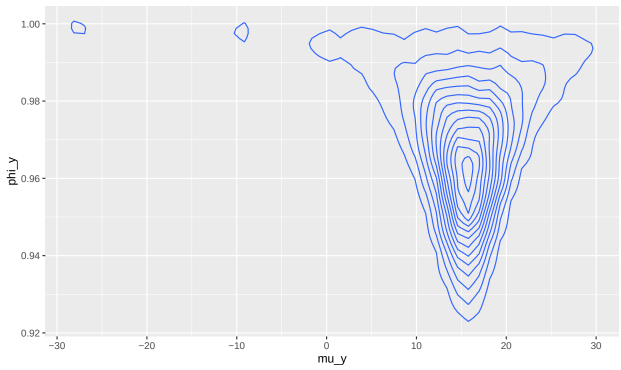
```
posterior_x <- extract(fit, pars = c("mu_x", "phi_x"))
posterior_df_x <- data.frame(posterior_x)
ggplot(data = posterior_df_x, aes(x = mu_x, y = phi_x)) +
    stat_density_2d() +
    xlab("mu_x") +
    ylab("phi_x") +
    ggtitle("Joint distribution of mu_x and phi_x")</pre>
```

Joint distribution of mu_x and phi_x



```
posterior_y <- extract(fit, pars = c("mu_y", "phi_y"))
posterior_df_y <- data.frame(posterior_y)
ggplot(data = posterior_df_y, aes(x = mu_y, y = phi_y)) +
    stat_density_2d()+
    xlab("mu_y") +
    ylab("phi_y") +
    ggtitle("Joint distribution of mu_y and phi_y")</pre>
```

Joint distribution of mu_y and phi_y



Appendix

```
rm(list=ls())
knitr::opts_chunk$set(echo = TRUE)
knitr::opts_chunk$set(fig.align = "center")
knitr::opts_chunk$set(fig.width=8, fig.height=5)
knitr::opts_chunk$set(warning=FALSE)
library(mvtnorm)
library(ggplot2)
library(gridExtra)
set.seed(12345)
#####01
library(ggplot2)
library(mvtnorm)
#library(extraDistr)
precipitation <- readRDS("Precipitation.rds")</pre>
log_prec <- log(precipitation)</pre>
mu_post <- function(mu_0,tau_0_sqr,sigma_square_current,y,n){</pre>
  tau_n_sqr = 1 / (1/tau_0_sqr + n/sigma_square_current)
  mu_n <- tau_n_sqr * (mu_0/tau_0_sqr + sum(y)/sigma_square_current)</pre>
  mupost <- rnorm(1, mu_n, sqrt(tau_n_sqr))</pre>
  return(mupost)
}
sigma_square_post <- function(mu_current, v_0, sigma_square_0, y,n, use_myinvchi=TRUE) {
  v n <- v 0 + n
  elem1 <- v_0*sigma_square_0
  elem2 <- sum((y - mu_current)^2)</pre>
  elem3 \leftarrow n+v_0
  elem_comb <- (elem1+elem2)/elem3</pre>
  if (use_myinvchi){
    sigma_square_post <- my_inv_chi(v_n,elem_comb)</pre>
  }
  else {
    sigma_square_post <- rinvchisq(1, v_n, elem_comb) #Requires package extraDistr
  return(sigma_square_post)
}
my_inv_chi<- function(df,tau_sqr) {</pre>
  X <- rchisq(1,df)</pre>
  inv_chi <- (df*tau_sqr)/X</pre>
  return(inv_chi)
}
#init
```

```
mu_0 <- 0
tau_0_sqr <- 1
sigma_square_0 <- 1
v_0 <- 1 #degree of freedom for chi square
gibbs_sampler <- function(nstep, data, mu_0, tau_0_sqr, v_0, sigma_square_0) {
  # Init parameters
 mu_current <- 0</pre>
  sigma_square_current <- 1</pre>
 mu_samples <- rep(0,nstep)</pre>
  sigma_square_samples <- rep(0,nstep)</pre>
  for (i in 1:nstep) {
    mu_current <- mu_post(mu_0, tau_0_sqr, sigma_square_current, y=data, length(data))</pre>
    #print(mu_current)
    sigma_square_current <- sigma_square_post(mu_current, v_0, sigma_square_0, y=data,
                                          length(data),use_myinvchi=TRUE)
    #print(sigma_square_current)
    mu_samples[i] <- mu_current</pre>
    sigma_square_samples[i] <- sigma_square_current</pre>
 output_df <- data.frame(mu_sample = mu_samples, sigma_sample = sigma_square_samples)</pre>
 return(output_df)
}
sample_gibbs <- gibbs_sampler(nstep=10000, data=log_prec, mu_0, tau_0_sqr, v_0, sigma_square_0)</pre>
my_acf <- acf(sample_gibbs$mu_sample,plot = F) # getting the autocorrelation
if_mu <- 1 + 2 * sum(my_acf$acf[-1])</pre>
my_acf <- acf(sample_gibbs$sigma_sample,plot = F)</pre>
if_sigma \leftarrow 1 + 2 * sum(my_acf_acf[-1])
ggplot(data=sample_gibbs, aes(x = 1:length(mu_sample), y = mu_sample)) +
 geom_line()+labs(title = "mu sample")
ggplot(data=sample_gibbs, aes(x = 1:length(sigma_sample), y = sigma_sample)) +
 geom_line() +labs(title = "sigma sample")
# Using Gibbs sample's mu and sigma to sample posterior prediction
post_pred_samples <- rnorm(n = 10000, mean = sample_gibbs$mu_sample,</pre>
                            sd = sqrt(sample_gibbs$sigma_sample))
ggplot(data = data.frame(y = log_prec), aes(x = y)) +
  geom_histogram(aes(y =after_stat(density)), color = "black", binwidth = 0.2) +
  geom_density(data = data.frame(y = post_pred_samples), aes(x = y, y = after_stat(density)),
               color = "red",linewidth = 2) +
  labs(x = "Log- Daily Precipitation ", y = "Density")
######Q2######
ebay_data <- read.table("eBayNumberOfBidderData.dat", header = TRUE)</pre>
```

```
model1 <- glm(formula = nBids ~ .,data = ebay_data[,-2],family = 'poisson')</pre>
summary(model1)
# Initialize values
n_cols <- ncol(ebay_data[,-1])</pre>
#remove 1st column since that is target variable and convert to matrix
# matrix of features
covariates <- as.matrix(ebay_data[,-1])</pre>
labels <- as.matrix(ebay_data[,1])</pre>
mu <- rep(0, n_cols)
initVal <- matrix(0, n_cols, 1)</pre>
Sigma <- as.matrix(100 * solve(t(covariates));*%covariates))</pre>
LogPosteriorFunc <- function(betas, X, y, mu, Sigma){</pre>
  log_prior <- dmvnorm(betas, mu, Sigma, log=TRUE)</pre>
  log_likelihood <- sum(X%*%betas * y - exp(X%*%betas) -log(factorial(y)))
 res <- log_prior + log_likelihood</pre>
  return(res)
}
# Optimizer
OptimRes <- optim(initVal, LogPosteriorFunc, gr = NULL, y = labels, X = covariates,
                   mu = mu, Sigma = Sigma, method=c("BFGS"),
                   control=list(fnscale=-1), hessian=TRUE)
beta_mode <- OptimRes$par</pre>
jacobian <- OptimRes$hessian
inv_jacobian <- -solve(jacobian)</pre>
beta_draws <- as.matrix(rmvnorm(10000, mean = beta_mode, sigma = inv_jacobian))
beta_estimate <- colMeans(beta_draws)</pre>
hist(beta_draws,breaks = 50,main = 'Histogram of Posterior Draws',xlab = 'Betas')
MetHas_RandomWalk <- function(nDraws,fun,mu,Sigma,c){</pre>
  #initialize matrix
  draw_matrix <- matrix(0,nrow = nDraws,ncol = n_cols)</pre>
  #initialize first row to mu
  draw_matrix[1,] <- mu</pre>
  for(i in 2:nDraws){
    # sample from multivariate normal distribution
    proposed_sample <- as.vector(rmvnorm(n = 1,mean = draw_matrix[i-1,],</pre>
                                           sigma = c*as.matrix(Sigma)))
    #print(proposed_sample)
    # IMPORTANT : the log is inside the posterior function
    log_acceptance_prob <- exp(fun(proposed_sample) - fun(draw_matrix[i-1,]))</pre>
    #random sample
    u <- runif(1)
    # calculate acceptance probability
```

```
a <- min(1,log_acceptance_prob)</pre>
    if(u \le a){
      #accept sample
      draw_matrix[i,] <- proposed_sample</pre>
    }
    else{
      # stay at same values from previous draw
      draw_matrix[i,] <- draw_matrix[i-1,]</pre>
    }
  }
  return(draw_matrix)
# this function we pass to MetHas Algorithm, can be changed to another posterior density
logPostFunc <- function(theta){</pre>
  res <- dmvnorm(theta, mean = beta_estimate, sigma = inv_jacobian, log = TRUE)
  if(is.na(res)){
    print(theta)
  return(res)
}
df <- MetHas_RandomWalk(nDraws = 10000,fun = logPostFunc,mu = rep(0,n_cols),</pre>
                         Sigma = inv_jacobian,c = 1)
# assign colnames
colnames(df) <- colnames(ebay_data)[2:10]</pre>
## plotting
plot_list <- list()</pre>
for (col in colnames(df)) {
  # Plot iterations vs every column
  p <- ggplot(data = as.data.frame(df), aes_string(x = 1:nrow(df), y = col)) +
    geom_line(col = 'blue') +
    labs(x = "Iterations", y = col)
  #show plot
  #print(p)
  plot_list[[col]] <- p</pre>
# Arranging in 1 fig
grid.arrange(grobs = plot_list, ncol = 2)
# extra 1 at the start for the intercept - Const
new_data \leftarrow c(1,1,0,1,0,1,0,1.2,0.8)
\# lambda = e^Beta*x
# discarding first 1500 samples as burn-in period
lambda <- exp(df[-c(1:1500),] %*% new_data)</pre>
```

```
samples <- c()</pre>
for (i in 1:nrow(df[-c(1:1500),])) {
  #sample from each row of df to get the predictive distribution based on the
  #posterior betas
  samples[i] <- rpois(1,lambda = lambda[i])</pre>
}
hist(samples)
res <- length(samples[samples == 0])/length(samples)</pre>
#####Q3
mu <- 13
sigma_square <- 3
t <- 300
phi \leftarrow seq(from = -1, to = 1, by = 0.25)
ar_func <- function(phi,mu,sigma_square,t){</pre>
  counter <- 0
  result_vector <- rep(0,t)
  result_vector[1] <- mu
  for(i in 2:t){
    epislon <- rnorm(1,0,sqrt(sigma_square))</pre>
    x_i <- mu+phi*(result_vector[i-1]-mu)+ epislon</pre>
    result_vector[i] <- x_i
  return(result_vector)
}
test_phi_func <- function(phi,mu,sigma_square,t){</pre>
  phi_test_df <- data.frame(matrix(0, nrow = t, ncol = length(phi)))</pre>
  colnames(phi_test_df) <- phi</pre>
  for (j in 1:length(phi)) {
    phi_test <- ar_func(phi[j], mu, sigma_square, t)</pre>
    phi_test_df[, j] <- phi_test</pre>
  return(phi_test_df)
phi_df <- test_phi_func(phi,mu,sigma_square,t)</pre>
for(k in 1:length(phi)){
  plot_data <- phi_df[,k]</pre>
  plot(x=1:300, plot_data, type = "l", main = paste(" phi = ", phi[k]))
  #Sys.sleep(1)
}
library(rstan)
set.seed(12345)
```

```
x <- ar_func(phi=0.2, mu, sigma_square, t)</pre>
y <- ar_func(phi=0.95, mu, sigma_square, t)
stan_code <- "
data {
 int<lower=0> T;
                     // Number of time points
 vector[T] x;
 vector[T] y;
}
parameters {
 real mu_x;
 real mu_y;
 real phi_x;
 real phi_y;
 real sigma_x;
 real sigma_y;
model {
 // After some research, it is common to use a flat prior or a vague prior as
 // non-informative prior
 // Therefore, we pick normal distribution with higher variance as prior.
 mu_x ~ normal(0, 50);
 mu_y ~ normal(0, 50);
  phi_x ~ normal(0, 10);
  phi_y ~ normal(0, 10);
  sigma_x ~ normal(0, 50);
  sigma_y ~ normal(0, 50);
 x[2:T] \sim normal(mu_x + phi_x * (x[1:(T - 1)] - mu_x), sigma_x);
 y[2:T] \sim normal(mu_y + phi_y * (y[1:(T - 1)] - mu_y), sigma_y);
data_list <- list(</pre>
 T = t,
 x = x
 y = y
# Set the MCMC settings
niter <- 5000
warmup <- 500
# Compile the Stan model
model <- stan_model(model_code = stan_code)</pre>
# Fit the Stan model to the data
# Set the control to avoid too many divergent after warmup
control <- list(adapt_delta = 0.90, stepsize = 0.0001)</pre>
```

```
fit <- sampling (model, data = data_list, warmup = warmup, iter = niter, chains = 5, control=control, refre
#refresh = 0 to mute the printout
summary(fit)$summary
post_mean <- summary(fit)$summary[, "mean"]</pre>
interval_025 <- summary(fit)$summary[, "25%"]</pre>
interval_975 <- summary(fit)$summary[, "97.5%"]</pre>
n eff <- summary(fit)$summary[, "n eff"]</pre>
Rhat <- summary(fit)$summary[, "Rhat"]</pre>
post_mean
interval_025
interval_975
n_eff
Rhat
posterior_x <- extract(fit, pars = c("mu_x", "phi_x"))</pre>
posterior_df_x <- data.frame(posterior_x)</pre>
ggplot(data = posterior_df_x, aes(x = mu_x, y = phi_x)) +
  stat_density_2d() +
  xlab("mu_x") +
  ylab("phi_x") +
  ggtitle("Joint distribution of mu_x and phi_x")
posterior_y <- extract(fit, pars = c("mu_y", "phi_y"))</pre>
posterior_df_y <- data.frame(posterior_y)</pre>
ggplot(data = posterior_df_y, aes(x = mu_y, y = phi_y)) +
  stat_density_2d()+
  xlab("mu_y") +
  ylab("phi_y") +
  ggtitle("Joint distribution of mu_y and phi_y")
```