Bayesian Learning Lab 3

Mohamed Ali - Mohal954

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Gibbs sampler for a normal model

\mathbf{A}

Implement (code!) a Gibbs sampler that simulates from the joint posterior $p(\mu, \sigma^2 | lny_1, ..., lnyn)$. The full conditional posteriors are given on the slides from Lecture 7.

Evaluate the convergence of the Gibbs sampler by calculating the Inefficiency Factors (IFs) and by plotting the trajectories of the sampled Markov chains.

The Gibbs Sampling algorithm:

- 1- Choose inital values $\theta_2^0, \theta_3^0, ..., \theta_n^0$. 2- Repeat for j=1,...,N: - Draw θ_1^j from $p(\theta_1|\theta_2^{j-1},...,\theta_k^{j-1})$. - Draw θ_2^j from $p(\theta_2|\theta_1^j,...,\theta_k^{j-1})$. . . 3- Draw θ_k^j from $p(\theta_k|\theta_1^j,...,\theta_k^j)$.
- 3- Draw θ_k^j from $p(\theta_k|\theta_1^j,...,\theta_{k-1}^j)$. 4- Return draws: $\theta^1,...,\theta^N$, where $\theta^j = (\theta_1^j,...,\theta_k^j)$.

$$\bar{\theta} = 1/N \sum_{t=1}^{N} \theta^{t}$$

$$var(\bar{\theta}) = \frac{\sigma^{2}}{N}$$

 $Autocorrelated\ samples:$

$$var(\bar{\theta}) = \frac{\sigma^2}{N} (1 + 2 \sum_{k=1}^{\inf} \rho_k)$$

Convergence diagnostics:

$$var(\bar{\theta}) = \frac{\sigma^2}{N} (1 + 2 \sum_{k=1}^{\inf} \rho_k)$$

$$IF = 1 + 2 \sum_{k=1}^{\inf} \rho_k$$

$$ESS = N/IF$$

We start by building our function using the Slides notes we can define the Normal Model with Conditionally conjugate prior as:

$$\mu \sim N(\mu_0, \tau_0^2)\sigma \sim Inv - \chi^2(v_0^2, \sigma_0^2)$$

And the full conditional posteriors:

$$\mu | \sigma^2, x \sim N(\mu_n, \tau_n^2)$$

 $\sigma^2 | \mu, x \sim Inv - \chi^2(v_n, \frac{v_0 \sigma^2 + \sum_{i=1}^n (x_i - \mu)^2}{n + v_0})$

Where we have:

$$\mu_n = w\bar{x} + (1 - w)\mu_0$$

$$w = \frac{\frac{n}{\sigma^2}}{\frac{n}{\sigma^2} + \frac{1}{\tau_0^2}}$$

$$\frac{1}{\tau_n^2} = \frac{n}{\sigma^2} + \frac{1}{\tau_0^2}$$

$$v_n = v_0 + n$$

We start by defining the above variables in R, Note that as we have $p(lny_1,...,lnyn|\mu,\sigma^2) \sim N(\mu,\sigma^2)$ we will transform our variable by taking the natural log of the daily precipitation.

```
# we start by reading our dataset using the readRDS command in R
df <- data.frame(x=readRDS("Precipitation.rds"))</pre>
# in the task we looking at natural log of the daily precipitation
#lny1,lny2,lny3,...lnyn follows N(mu,siqma)
# We add a new var called logx
df$logx<- log(df$x)
sigma2 0<- var(df$logx) # Sample var</pre>
tau2_0<- 1# arbirtry initail value let it be 1
n<- nrow(df)# Sample size
mu_0<- mean(df$logx)#sample mean</pre>
w \leftarrow (n/sigma2_0)/((n/sigma2_0) + (1/tau2_0)) # value used to calculate mu_n
v_0<- 1 # arbirtry initail value let it be 1
v_n<- v_0+n
mu_n \leftarrow w*(mean(df$logx)) + (1-w)*mu_0
###### This part of the code has been modified#####
tau2_n<- 1/((n/sigma2_0)+(1/tau2_0))########### shoud be 1/tau
nDraws<- 1000
gibbsDraws <- matrix(0,nDraws,2)</pre>
```

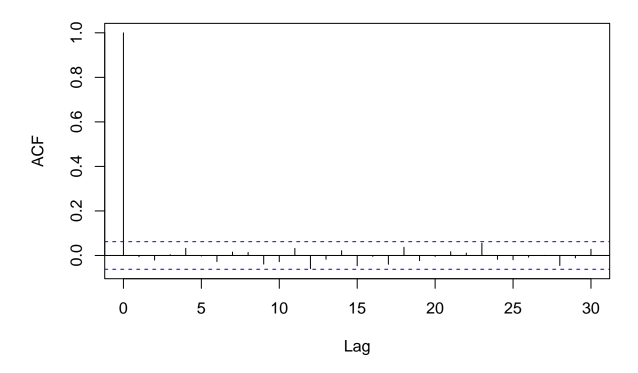
Now we write a code to simulates from the joint posterior using Gibbs sampler.

```
gibbsDraws[i,1] <- theta1

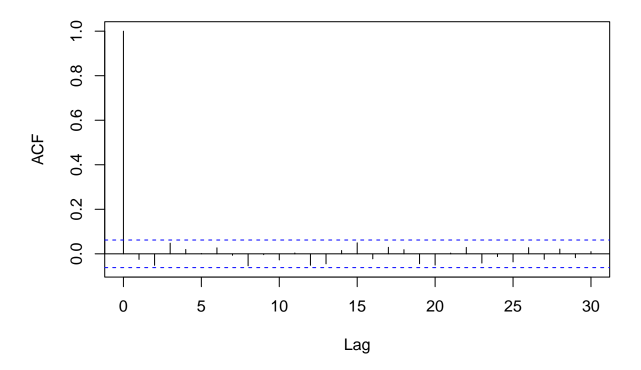
scale<- ((v_0*sigma2_0)+(sum((df$logx-theta1)^2)))/(n+v_0)
# Update theta2 ---> sigma2 given theta1
theta2 <- rinvchisq(1,v_n,scale=scale)
gibbsDraws[i,2] <- theta2
}</pre>
```

To Evaluate the convergence of the Gibbs sampler we calculate the Inefficiency Factors (IFs) $1 + 2\sum_{k=1}^{\infty} \rho_k$ - where ρ_k is the autocorrelation at lag k - and then we plot the trajectories of the sampled Markov chains.

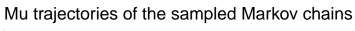
Series gibbsDraws[, 1]

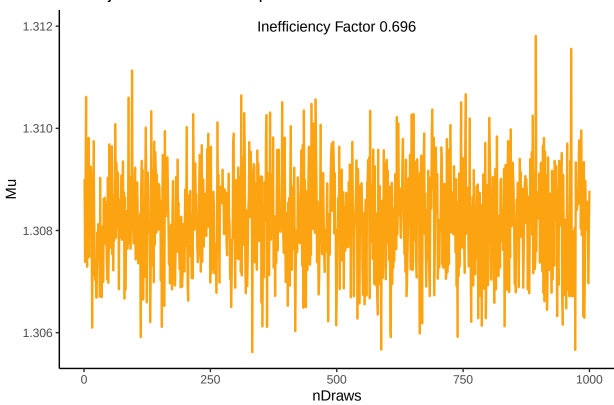


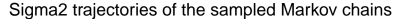
Series gibbsDraws[, 2]



- ## Warning: Using 'size' aesthetic for lines was deprecated in ggplot2 3.4.0.
- ## i Please use 'linewidth' instead.
- ## This warning is displayed once every 8 hours.
- ## Call 'lifecycle::last_lifecycle_warnings()' to see where this warning was
- ## generated.









The graph above shows the convergence of the Gibbs sampler and the Inefficiency Factors, the IF tell us number of iterations needed to get an independent sample from the posterior distribution, as we can see μ and σ^2 have 0,69 and 0,57 IF respectively, which tell us a better the convergence of the Gibbs sampler for the σ^2 .

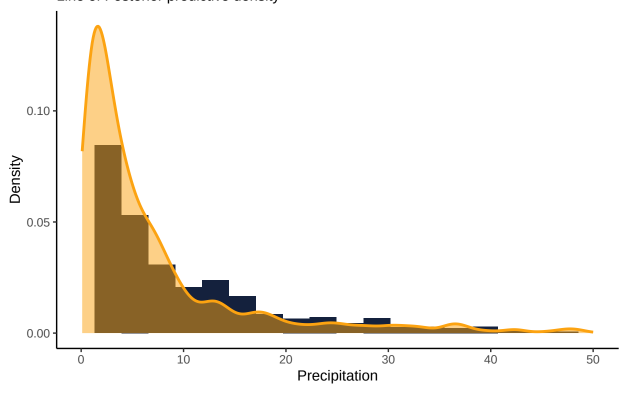
\mathbf{B}

Plot the following in one figure:

- 1) a histogram or kernel density estimate of the daily precipitation $(y_1, ..., y_n)$.
- 2) The resulting posterior predictive density $p(\tilde{y}|y_1,...,y_n)$ using the simulated posterior draws from (a). How well does the posterior predictive density agree with this data?

To do we draw a sample from rnorm with mean and sd from the posterior results we got in (a).

Histogram of Daily Precipitation Line of Posterior predictive density



Metropolis Random Walk for Poisson regression

\mathbf{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?

```
##
## Call:
## glm(formula = nBids ~ ., family = poisson, data = df1[, -2])
##
## Deviance Residuals:
##
                  1Q
                       Median
                                    3Q
                                             Max
##
   -3.5800
            -0.7222
                      -0.0441
                                0.5269
                                          2.4605
##
## Coefficients:
               Estimate Std. Error z value Pr(>|z|)
## (Intercept) 1.07244
                            0.03077
                                     34.848
                                              < 2e-16 ***
## PowerSeller -0.02054
                            0.03678
                                      -0.558
                                               0.5765
## VerifyID
               -0.39452
                            0.09243
                                      -4.268 1.97e-05 ***
## Sealed
                0.44384
                            0.05056
                                      8.778
                                              < 2e-16 ***
## Minblem
               -0.05220
                            0.06020
                                     -0.867
                                               0.3859
```

```
## MajBlem
               -0.22087
                           0.09144
                                     -2.416
                                              0.0157 *
                           0.05633
                                              0.2096
## LargNeg
                0.07067
                                      1.255
               -0.12068
                           0.02896
## LogBook
                                     -4.166 3.09e-05 ***
## MinBidShare -1.89410
                           0.07124 -26.588
                                             < 2e-16 ***
##
                   0 '*** 0.001 '** 0.01 '* 0.05 '.' 0.1 ' 1
## Signif. codes:
##
##
   (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
  AIC: 3610.3
##
##
## Number of Fisher Scoring iterations: 5
```

The model reuslts shows the coefficients and their standard errors for each predictor. The "Estimate" shows the estimated effect of each predictor on the expected number of bids. A positive coefficient indicates that the predictor is associated with an increase in the expected number of bids, while a negative coefficient indicates a decrease.

The "Pr(>|z|)" column shows the p-value for each coefficient. If this value is less than 0.05, the coefficient is considered statistically significant.

The model suggests that the variables VerifyID, Sealed, LogBook, and MinBidShare have a significant effect on the expected number of bids (p-value less that 0.01 (p<.01) at $\alpha = 0.01$), while the others do not.

\mathbf{B}

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

$$\beta|y \sim N(\tilde{\beta}, J_y^{-1}(\tilde{\beta}))$$

where $\tilde{\beta}$ is the posterior mode and $J_y^{-1}(\tilde{\beta})$ is the negative Hessian at the posterior mode. $\tilde{\beta}$ and $J_y^{-1}(\tilde{\beta})$ 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.).

```
## [1] "The posterior mode is:"
##
             [,1]
                         [,2]
                                    [,3]
                                               [,4]
                                                            [,5]
                                                                        [,6]
                                                                                   [,7]
## [1,] 1.069841 -0.02051246 -0.393006 0.4435555 -0.05246627 -0.2212384 0.07069683
##
               [,8]
                         [,9]
## [1,] -0.1202177 -1.891985
## attr(,"names")
## [1] "Const"
                      "PowerSeller" "VerifyID"
                                                    "Sealed"
                                                                   "Minblem"
   [6] "MajBlem"
                      "LargNeg"
                                     "LogBook"
                                                    "MinBidShare"
## [1] "The Hessian Matrix:"
```

```
##
               [,1]
                            [,2]
                                           [,3]
                                                          [,4]
                                                                         [,5]
    [1,] -3634.2841 -1574.88862 -1.284330e+02 -5.054825e+02 -3.089573e+02
##
                    -1574.88862 -6.049186e+01 -3.260764e+02 -1.044615e+02
##
    [2,] -1574.8886
    [3,]
          -128.4330
                       -60.49186 -1.284330e+02 -6.148277e+01 -9.865299e+00
##
##
    [4,]
          -505.4825
                      -326.07643 -6.148277e+01 -5.054825e+02
                                                               1.705303e-07
          -308.9573
                      -104.46148 -9.865299e+00
##
    [5,]
                                                 1.705303e-07 -3.089573e+02
##
    [6,]
          -126.9303
                       -68.96966
                                  2.273737e-07
                                                 0.000000e+00
                                                               0.00000e+00
    [7,]
##
          -385.7170
                       -53.05278 -1.136868e-07 0.000000e+00 -3.390566e+01
##
    [8,]
          -638.9730
                        71.79073 -6.887776e+01 -1.287304e+02 -2.229706e+01
##
    [9,]
           729.8896
                       146.21556
                                 2.380017e+01 8.975677e+01
                                                               5.518223e+01
##
                   [,6]
                                 [,7]
                                              [,8]
                                                          [,9]
    [1,] -1.269303e+02 -3.857170e+02
                                        -638.97297
##
                                                    729.88956
##
    [2,] -6.896966e+01 -5.305278e+01
                                          71.79073
                                                    146.21556
##
          2.273737e-07 -1.136868e-07
                                         -68.87776
                                                     23.80017
    [4,]
          0.000000e+00 0.000000e+00
                                                     89.75677
##
                                        -128.73043
##
    [5,]
          0.000000e+00 -3.390566e+01
                                         -22.29706
                                                     55.18223
##
    [6,] -1.269303e+02 0.000000e+00
                                         -36.39914
                                                     34.16904
##
         0.000000e+00 -3.857170e+02
                                        -220.23559
                                                    115.38523
##
    [8,] -3.639914e+01 -2.202356e+02
                                      -1930.07936
                                                    534.16381
##
          3.416904e+01
                        1.153852e+02
                                         534.16381 -446.88731
   [1] "The approximate posterior standard deviation is:"
##
         Const PowerSeller
                               VerifyID
                                              Sealed
                                                         Minblem
                                                                      MajBlem
                                         0.05057448
##
    0.03074837
                0.03678418
                             0.09227871
                                                      0.06020470
                                                                   0.09146070
##
       LargNeg
                    LogBook MinBidShare
##
    0.05634767
                0.02895635
                             0.07109682
```

\mathbf{C}

Random walk Metropolis algorithm:

- Initialize θ^0 and iterate for i=1,2,...
 - 1- Sample proposal: $\theta_p | \theta^{(i-1)} \sim N(\theta^{(i-1)}, c.\Sigma)$.
 - 2- Compute the acceptance probability $(\alpha = min(1, \frac{p(\theta_p|y)}{\theta^{(i-1)}|y|}))$.
 - 3- With probability α true set $\theta^i = \theta_p$ and $\theta^i = \theta^{i-1}$.

Note

The draw backs of the MH usual find it hard to find proposal distribution and getting too small step size give us too many rejections.

Alternately we can use HM (Hamiltonian Monto Carlo), which solve this by giving distant proposal and high acceptance probabilities.

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):

$$\theta_p | \theta^{i-1} \sim N(\theta^{i-1}, c.\Sigma)$$

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. * Note that* on how to How to code up the Random Walk Metropolis Algorithm in R:

One of the input arguments of our RWMSampler function is logPostFunc. logPostFunc is a function object that computes the log posterior density at any value of the parameter vector.

This is needed when we compute the acceptance probability of the Metropolis algorithm. from (q2 a) we program the log posterior density, since logs are more stable and avoids problems with too small or large numbers (overflow).

The ratio of posterior densities in the Metropolis acceptance probability can be written as:

$$\frac{p(\theta_p|y)}{p(\theta^{i-1}|y)} = exp(\log p(\theta_p|y) - \log p(\theta^{i-1}|y))$$

The first argument our (log) posterior function is theta (theta_old,theta_new), the vector of parameters for which the posterior density is evaluated. You can of course use some other name for the variable, but it must be the first argument of your posterior density function.

Function RWMSampler

```
RWMSampler_old<- function(logPostFunc,nDraws,c,y,x,mu,Sigma){</pre>
  # First we buld our data frame of samples
  sample <- data.frame(matrix(nrow = nDraws, ncol = ncol(x)))</pre>
  colnames(sample) <- colnames(x)</pre>
  # The inital sample value c here represent a tuning parameter
  sample[1,] <- mvrnorm(1, posteriorMode, c*postCov)</pre>
  # Now we implement the Metropolis-Hastings
  #in which we generate samples from the proposal distribution in this case
  # We look at the results of the first sample
  #as theta i-1 pluged in monorm to get theta i and the we use the values in
  # our proposed logPostFunc
  counter <- 1
  i=1
  while (counter < nDraws) {</pre>
    theta_old<-as.numeric(sample[counter,])</pre>
    theta_new<-mvrnorm(1,theta_old,c*postCov)
  # We define th our accept/reject threshold
    th<-runif(1,0,1)
  # now we find the value of the target/proposed distribution
    proposed<- logPostFunc(theta_new,y = y,</pre>
                           x = x
                           mu = posteriorMode,
                           Sigma = postCov)
    target<- logPostFunc(theta_old,y = y,</pre>
                         x = x
                         mu = posteriorMode,
```

```
Sigma = postCov)
# the ratio of posterior densities in the Metropolis acceptance probability
if (th<min(1,exp(proposed-target))) {
    counter=counter+1
    sample[counter,]<-theta_new
    }
}
return(sample)
}</pre>
```

Old solution

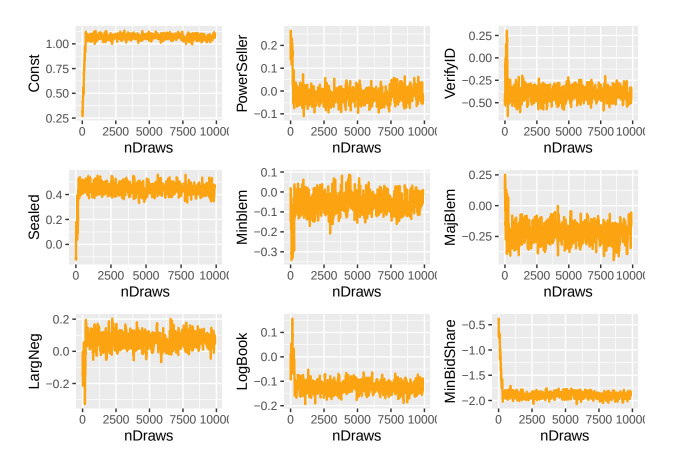
```
RWMSampler<- function(logPostFunc,nDraws,c,y,x,mu,Sigma,brnin){</pre>
  # First we buld our data frame of samples
  sample <- data.frame(matrix(0,nrow = nDraws - brnin, ncol = ncol(x)))</pre>
  colnames(sample) <- colnames(x)</pre>
  # The inital sample value c here represent a tuning parameter
  # sample[1,] <- murnorm(1, posteriorMode, c*postCov)</pre>
  # Now we implement the Metropolis-Hastings
  #in which we generate samples from the proposal distribution in this case
  # We look at the results of the first sample
  #as theta_i-1 pluged in monorm to get theta_i and the we use the values in
  # our proposed logPostFunc
  theta_old<-as.numeric(sample[1,])</pre>
  for (i in 1:nDraws) {
    theta_new<-mvrnorm(1,theta_old,c*postCov)</pre>
    # We define th our accept/reject threshold
    th<-runif(1,0,1)
    # now we find the value of the target/proposed distribution
    proposed<- logPostFunc(theta_new,y = y,</pre>
                            x = x
                            mu = posteriorMode,
                            Sigma = postCov)
    target<- logPostFunc(theta_old,y = y,</pre>
                          x = x
                          mu = posteriorMode,
                          Sigma = postCov)
    # the ratio of posterior densities in the Metropolis acceptance probability
    if (th<exp(proposed-target)) {</pre>
      theta_old<-theta_new
    if (i> brnin) {
      sample[i-brnin,]<-theta_old</pre>
    }
  }
  return(sample)
}
```

New Solution

• 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.

```
## Warning: 'aes_string()' was deprecated in ggplot2 3.0.0.
## i Please use tidy evaluation idioms with 'aes()'.
## i See also 'vignette("ggplot2-in-packages")' for more information.
## This warning is displayed once every 8 hours.
## Call 'lifecycle::last_lifecycle_warnings()' to see where this warning was
## generated.
```



\mathbf{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?

```
* VerifyID = 0

* Sealed = 1

* MinBlem = 0

* MajBlem = 1

* LargNeg = 0

* LogBook = 1.2

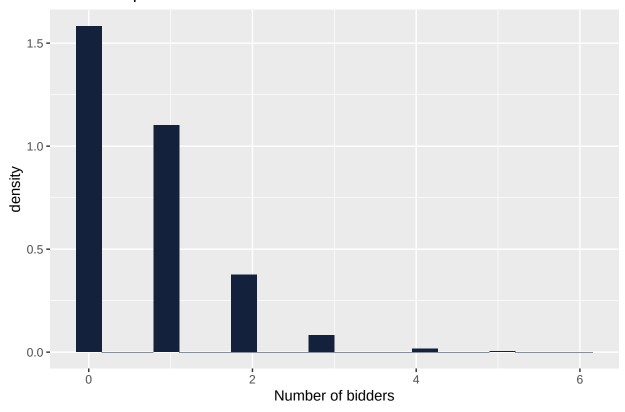
* MinBidShare = 0.8
```

* PowerSeller = 1

```
#First we estimate the betas from our RWMSampler function
betas<- as.matrix(res)
# Input data
x_new <- as.matrix(c(1,1,0,1,0,1,0,1.2,0.8))
##### This code has been modified
# calculating the bet for Poisson since we have our y follow poisson
beta_pois<-exp(betas %*% x_new)
# Finding number of bidders using poisson function
nbiders<-data.frame(x=rpois(length(beta_pois),beta_pois))</pre>
```

Plotting the predictive distribution. What is the probability of no bidders in this new auction?

Plot of the predictive distribution for Number of bidders



Time series models in Stan

\mathbf{A}

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

 $x_t = \mu + \phi(x_{t-1} - \mu) + \epsilon_t$, $\epsilon_t \sim N(0, +\sigma^2)$ for given values of μ , ϕ and σ^2 .

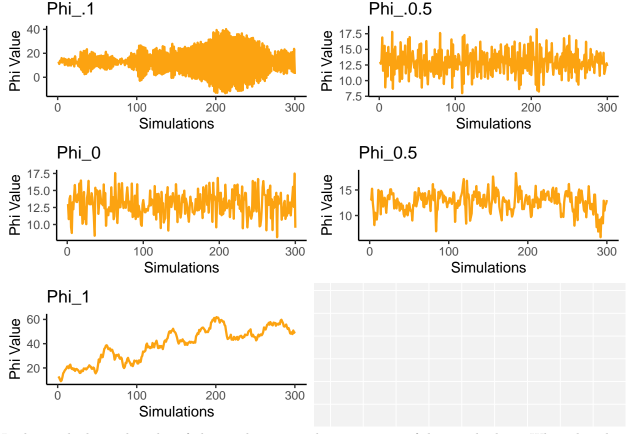
Start the process at $x1 = \mu$ and then simulate values for x_t for t = 1, 2, 3, ..., T and return the vector x1:T containing all time points.

Use μ = 13; σ^2 = 3 and T = 300 and look at some different realizations (simulations) of $x_{1:t}$ for values of ϕ between -1 and 1 (this is the interval of ϕ where the AR(1)-process is stationary). Include a plot of at least one realization in the report.

What effect does the value of ϕ have on $x_{1:t}$?

```
ar_process<- function(phi,mu, sigma,t){
    x_t<-c()
    x_t[1]<-mu
    for(i in 2:t){
        e<- rnorm(1,0,sqrt(3))
        x_t[i]<-mu+(phi*(x_t[i-1]-mu))+e
    }
    return(x_t)
}</pre>
```

Now we Include a plot of at least one realization in the report. What effect does the value of ϕ have on $x_{1:t}$?



In the graph above, the value of phi greatly impacts the convergence of the sample chain. When phi is less than 0, it can lead to instability and hinder the convergence to a stationary distribution. This instability is evident when the chains appear to wander around or exhibit erratic behavior, indicating that the sampler has not yet converged.

However, it is notable that the chains demonstrate good mixing, meaning they efficiently explore the parameter space and cover a wide range of plausible values.

To achieve convergence, it may be necessary to increase the number of iterations.

It is important to note that when phi is equal to 1, the convergence appears to be reached more quickly compared to when phi is less than 0.

\mathbf{B}

Use your function from a) to simulate two AR(1)-processes, $x_{1:T}$ with $\phi = 0.2$ and $y_{1:T}$ with $\phi = 0.95$. Now, treat your simulated vectors as synthetic data, and treat the values of , μ and σ^2 as unknown parameters. Implement Stancode that samples from the posterior of the three parameters, using suitable non-informative priors of your choice. Notes from: (https://mc-stan.org/docs/stan-users-guide/autoregressive.html). We have A first-order autoregressive model (AR(1)) with normal noise takes each point y_n in a sequence y to be generated according to:

$$y_n \sim N(\alpha + \beta y_{n-i}, \sigma)$$

That is, the expected value of y_n is $\alpha + \beta y_{n-i}$, with noise scaled as σ .

With improper flat priors on the regression coefficients α and β and on the positively-constrained noise scale (σ) , the Stan program for the AR(1) model is as follows:

```
vector[N] y;
}
parameters {
  real alpha;
  real beta;
  real<lower=0> sigma;
}
model {
  for (n in 2:N) {
    y[n] ~ normal(alpha + beta * y[n-1], sigma);
  }
}
```

Adding the the Slicing for efficiency in which we silce the vector we can rewrite the above as:

```
data {
  int<lower=0> N;
  vector[N] y;
}
parameters {
  real alpha;
  real beta;
  real<lower=0> sigma;
}
model {
    y[2:N] ~ normal(alpha + beta * y[1:(N - 1)], sigma);
}
```

From lec notes we have Gibbs sampling for AR process:

If we have AP(p) process:

$$x_t = \mu + \phi_1(x_{t-1} - \mu) + \dots + \phi_p(x_{t-p} - \mu) + \epsilon, \ \epsilon_t \sim N(0, \sigma^2)$$

let $\phi = (\phi_1, ..., \phi_p)'$, we have prior:

 $\mu \sim Normal$ $\phi \sim Multivariate\ Normal$ $\sigma^2 \sim Scaled\ inverse\ \chi$

And the posterior can be simulated by Gibbs sampling:

$$\mu | \phi \ \sigma^2 \sim Normal$$

 $\phi | \mu \ \sigma^2 \sim Multivariate \ Normal$
 $\sigma^2 | \mu \ \phi \sim Scaled \ inverse \ \chi$

StanModel

```
##### Example from notes #####
# library(rstan)
y=c(4,5,6,4,0,2,5,3,8,6,10,8)
# N=length(y)
# StanModel = '
# data {
   int<lower=0> N; // Number of observations
   int<lower=0> y[N]; // Number of flowers
# }
# parameters {
  real mu;
#
   real<lower=0> sigma2;
# }
# model {
  mu ~ normal(0,100); // Normal with mean 0, st.dev. 100
#
   sigma2 ~ scaled_inv_chi_square(1,2); // Scaled-inv-chi2 with nu 1,sigma 2
#
   for(i in 1:N){
#
     y[i] ~ normal(mu,sqrt(sigma2));
#
# }'
#
# data \leftarrow list(N=N, y=y)
# warmup <- 1000
# niter <- 2000
\# fit <- stan(model\_code=StanModel, data=data, warmup=warmup, iter=niter, chains=4)
# # Print the fitted model
# print(fit, diqits_summary=3)
# # Extract posterior samples
# postDraws <- extract(fit)</pre>
# # Do traceplots of the first chain
\# par(mfrow = c(1,1))
\#\ plot(postDraws\$mu[1:(niter-warmup)], type="l", ylab="mu", main="Traceplot")
# # Do automatic traceplots of all chains
# traceplot(fit)
# # Bivariate posterior plots
# pairs(fit)
# #################################
StanModel = '
data {
  int<lower=0> N; // Number of observations
  vector[N] y;
}
parameters {
  real mu;
 real<lower=0> sigma2;
 real<lower=-1, upper=1> phi;
 //To enforce the estimation of a
  //stationary AR(1) process, the slope
  //coefficient beta may be constrained with bounds as follows.
}
model {
mu ~ normal(0,100); // Normal with mean 0, st.dev. 100
```

```
sigma2 ~ scaled_inv_chi_square(1,2); // Scaled-inv-chi2 with nu 1,sigma 2
// Changing the model to be y_i-mu
y[2:N] ~ normal(mu + phi * (y[1:(N - 1)]-mu), sqrt(sigma2^2));
}'
```

Model results

Part i Report the posterior mean, 95% credible intervals and the number of effective posterior samples for the three inferred parameters for each of the simulated AR(1)-process. Are you able to estimate the true values?

Giving $x_{1:T}$ with $\phi = 0.2$ and $y_{1:T}$ with $\phi = 0.95$, the model results can be shown as below.

1- For $x_{1:T}$ with $\phi = 0.2$

```
# Simulate of x
ar_x<-ar_process(.2, 13, 3, 300)

#From lec notes we have the stanmodel function defined as
y=ar_x
N=length(y)

data <- list(N=N, y=y)
warmup <- 1000
niter <- 2000
fitx <- stan(model_code=StanModel,data=data,warmup=warmup,iter=niter,chains=4)</pre>
```

```
##
## SAMPLING FOR MODEL '8874c61302b41c9e7aa22f87c38effdb' NOW (CHAIN 1).
## Chain 1:
## Chain 1: Gradient evaluation took 0 seconds
## Chain 1: 1000 transitions using 10 leapfrog steps per transition would take 0 seconds.
## Chain 1: Adjust your expectations accordingly!
## Chain 1:
## Chain 1:
## Chain 1: Iteration: 1 / 2000 [ 0%]
                                            (Warmup)
## Chain 1: Iteration: 200 / 2000 [ 10%]
                                            (Warmup)
## Chain 1: Iteration: 400 / 2000 [ 20%]
                                            (Warmup)
## Chain 1: Iteration: 600 / 2000 [ 30%]
                                            (Warmup)
## Chain 1: Iteration: 800 / 2000 [ 40%]
                                            (Warmup)
## Chain 1: Iteration: 1000 / 2000 [ 50%]
                                            (Warmup)
## Chain 1: Iteration: 1001 / 2000 [ 50%]
                                            (Sampling)
## Chain 1: Iteration: 1200 / 2000 [ 60%]
                                            (Sampling)
## Chain 1: Iteration: 1400 / 2000 [ 70%]
                                            (Sampling)
## Chain 1: Iteration: 1600 / 2000 [ 80%]
                                            (Sampling)
## Chain 1: Iteration: 1800 / 2000 [ 90%]
                                            (Sampling)
## Chain 1: Iteration: 2000 / 2000 [100%]
                                            (Sampling)
## Chain 1:
## Chain 1: Elapsed Time: 0.159 seconds (Warm-up)
## Chain 1:
                           0.069 seconds (Sampling)
                           0.228 seconds (Total)
## Chain 1:
```

```
## Chain 1:
##
## SAMPLING FOR MODEL '8874c61302b41c9e7aa22f87c38effdb' NOW (CHAIN 2).
## Chain 2:
## Chain 2: Gradient evaluation took 0 seconds
## Chain 2: 1000 transitions using 10 leapfrog steps per transition would take 0 seconds.
## Chain 2: Adjust your expectations accordingly!
## Chain 2:
## Chain 2:
## Chain 2: Iteration:
                        1 / 2000 [ 0%]
                                            (Warmup)
## Chain 2: Iteration: 200 / 2000 [ 10%]
                                            (Warmup)
## Chain 2: Iteration: 400 / 2000 [ 20%]
                                            (Warmup)
## Chain 2: Iteration: 600 / 2000 [ 30%]
                                            (Warmup)
## Chain 2: Iteration: 800 / 2000 [ 40%]
                                            (Warmup)
## Chain 2: Iteration: 1000 / 2000 [ 50%]
                                            (Warmup)
## Chain 2: Iteration: 1001 / 2000 [ 50%]
                                            (Sampling)
## Chain 2: Iteration: 1200 / 2000 [ 60%]
                                            (Sampling)
## Chain 2: Iteration: 1400 / 2000 [ 70%]
                                            (Sampling)
## Chain 2: Iteration: 1600 / 2000 [ 80%]
                                            (Sampling)
## Chain 2: Iteration: 1800 / 2000 [ 90%]
                                            (Sampling)
## Chain 2: Iteration: 2000 / 2000 [100%]
                                            (Sampling)
## Chain 2:
## Chain 2: Elapsed Time: 0.309 seconds (Warm-up)
## Chain 2:
                           0.073 seconds (Sampling)
## Chain 2:
                           0.382 seconds (Total)
## Chain 2:
##
## SAMPLING FOR MODEL '8874c61302b41c9e7aa22f87c38effdb' NOW (CHAIN 3).
## Chain 3:
## Chain 3: Gradient evaluation took 0 seconds
## Chain 3: 1000 transitions using 10 leapfrog steps per transition would take 0 seconds.
## Chain 3: Adjust your expectations accordingly!
## Chain 3:
## Chain 3:
## Chain 3: Iteration:
                        1 / 2000 [ 0%]
                                            (Warmup)
## Chain 3: Iteration: 200 / 2000 [ 10%]
                                            (Warmup)
## Chain 3: Iteration: 400 / 2000 [ 20%]
                                            (Warmup)
## Chain 3: Iteration: 600 / 2000 [ 30%]
                                            (Warmup)
## Chain 3: Iteration: 800 / 2000 [ 40%]
                                            (Warmup)
## Chain 3: Iteration: 1000 / 2000 [ 50%]
                                            (Warmup)
## Chain 3: Iteration: 1001 / 2000 [ 50%]
                                            (Sampling)
## Chain 3: Iteration: 1200 / 2000 [ 60%]
                                            (Sampling)
## Chain 3: Iteration: 1400 / 2000 [ 70%]
                                            (Sampling)
## Chain 3: Iteration: 1600 / 2000 [ 80%]
                                            (Sampling)
## Chain 3: Iteration: 1800 / 2000 [ 90%]
                                            (Sampling)
## Chain 3: Iteration: 2000 / 2000 [100%]
                                            (Sampling)
## Chain 3:
## Chain 3: Elapsed Time: 0.082 seconds (Warm-up)
## Chain 3:
                           0.078 seconds (Sampling)
## Chain 3:
                           0.16 seconds (Total)
## Chain 3:
## SAMPLING FOR MODEL '8874c61302b41c9e7aa22f87c38effdb' NOW (CHAIN 4).
## Chain 4:
```

```
## Chain 4: Gradient evaluation took 0 seconds
## Chain 4: 1000 transitions using 10 leapfrog steps per transition would take 0 seconds.
## Chain 4: Adjust your expectations accordingly!
## Chain 4:
## Chain 4:
## Chain 4: Iteration:
                        1 / 2000 [ 0%]
                                            (Warmup)
## Chain 4: Iteration: 200 / 2000 [ 10%]
                                            (Warmup)
## Chain 4: Iteration: 400 / 2000 [ 20%]
                                            (Warmup)
## Chain 4: Iteration: 600 / 2000 [ 30%]
                                            (Warmup)
## Chain 4: Iteration: 800 / 2000 [ 40%]
                                            (Warmup)
## Chain 4: Iteration: 1000 / 2000 [ 50%]
                                            (Warmup)
## Chain 4: Iteration: 1001 / 2000 [ 50%]
                                            (Sampling)
## Chain 4: Iteration: 1200 / 2000 [ 60%]
                                            (Sampling)
## Chain 4: Iteration: 1400 / 2000 [ 70%]
                                            (Sampling)
## Chain 4: Iteration: 1600 / 2000 [ 80%]
                                            (Sampling)
## Chain 4: Iteration: 1800 / 2000 [ 90%]
                                            (Sampling)
## Chain 4: Iteration: 2000 / 2000 [100%]
                                            (Sampling)
## Chain 4:
## Chain 4: Elapsed Time: 0.092 seconds (Warm-up)
## Chain 4:
                           0.072 seconds (Sampling)
## Chain 4:
                           0.164 seconds (Total)
## Chain 4:
# Print the fitted model
print(fitx,digits_summary=3)
## Inference for Stan model: 8874c61302b41c9e7aa22f87c38effdb.
## 4 chains, each with iter=2000; warmup=1000; thin=1;
## post-warmup draws per chain=1000, total post-warmup draws=4000.
##
                                                                  75%
##
                                      2.5%
                                                25%
                                                                         97.5%
              mean se mean
                              sd
                                                         50%
## mu
            13.137
                     0.002 0.111
                                    12.917
                                             13.064
                                                      13.137
                                                               13.210
                                                                         13.356
                     0.001 0.064
                                              1.518
## sigma2
           1.563
                                    1.443
                                                       1.560
                                                                1.605
                                                                          1.692
## phi
             0.204
                     0.001 0.056
                                    0.094
                                              0.166
                                                       0.204
                                                                0.241
                                                                          0.311
          -284.604
                     0.027 1.209 -287.783 -285.164 -284.296 -283.733 -283.249
## lp__
          n eff Rhat
##
## mu
           3594 1.000
## sigma2 4056 0.999
           4246 1.000
## phi
## lp__
           2058 1.000
##
## Samples were drawn using NUTS(diag_e) at Mon May 29 21:52:07 2023.
## For each parameter, n_eff is a crude measure of effective sample size,
## and Rhat is the potential scale reduction factor on split chains (at
## convergence, Rhat=1).
samples_x <-extract(fitx, pars = c("mu", "sigma2", "phi"))</pre>
# Compute the mean
mean_x <- sapply(samples_x, mean)</pre>
# 95% credible intervals
intv_x <- sapply(samples_x,function(samples) quantile(samples,c(0.025, 0.975)))
# effective posterior samples
eff_samp_x <- summary(fitx)$summary[1:3,"n_eff"]</pre>
```

```
##
                Mean Credible_Interval_Lower Credible_Interval_Upper
## mu
                                12.91693642
          13.1371909
                                                          13.3561165
## sigma2 1.5625755
                                 1.44261786
                                                           1.6921687
           0.2035987
                                  0.09371789
                                                           0.3106965
## phi
##
          Effective_Samples
## mu
                   3593.851
## sigma2
                   4055.560
## phi
                   4246.170
```

Based on the table above, the simulations performed using the Metropolis algorithm have provided estimates for the parameters μ , σ , and ϕ .

The estimated values are found to be approximately 11.6, 1.6, and 0.11, respectively.

These estimates are close to the true values used to simulate the AR(1) process, indicating that the sampling algorithm has been not effective in capturing the underlying parameter values. This suggests that the simulation has not successfully captured the characteristics of the data and has produced reliable estimates for the parameters of interest.

2- For $y_{1:T}$ with $\phi = 0.95$

```
# Simulate of y
ar_y<-ar_process(.95, 13, 3, 300)
#From lec notes we have the stanmodel function defined as
#y=ar_y
N=length(y)
data <- list(N=N, y=y)</pre>
warmup <- 1000
niter <- 2000
fity <- stan(model_code=StanModel,data=data, warmup=warmup,iter=niter,chains=4)
##
## SAMPLING FOR MODEL '8874c61302b41c9e7aa22f87c38effdb' NOW (CHAIN 1).
## Chain 1:
## Chain 1: Gradient evaluation took 0 seconds
## Chain 1: 1000 transitions using 10 leapfrog steps per transition would take 0 seconds.
## Chain 1: Adjust your expectations accordingly!
## Chain 1:
## Chain 1:
## Chain 1: Iteration: 1 / 2000 [ 0%]
                                            (Warmup)
## Chain 1: Iteration: 200 / 2000 [ 10%]
                                            (Warmup)
## Chain 1: Iteration: 400 / 2000 [ 20%]
                                            (Warmup)
## Chain 1: Iteration: 600 / 2000 [ 30%]
                                            (Warmup)
## Chain 1: Iteration: 800 / 2000 [ 40%]
                                            (Warmup)
## Chain 1: Iteration: 1000 / 2000 [ 50%]
                                            (Warmup)
## Chain 1: Iteration: 1001 / 2000 [ 50%]
                                            (Sampling)
```

```
## Chain 1: Iteration: 1200 / 2000 [ 60%]
                                            (Sampling)
## Chain 1: Iteration: 1400 / 2000 [ 70%]
                                            (Sampling)
## Chain 1: Iteration: 1600 / 2000 [ 80%]
                                            (Sampling)
## Chain 1: Iteration: 1800 / 2000 [ 90%]
                                            (Sampling)
## Chain 1: Iteration: 2000 / 2000 [100%]
                                            (Sampling)
## Chain 1:
## Chain 1: Elapsed Time: 0.098 seconds (Warm-up)
## Chain 1:
                           0.066 seconds (Sampling)
                           0.164 seconds (Total)
## Chain 1:
## Chain 1:
##
## SAMPLING FOR MODEL '8874c61302b41c9e7aa22f87c38effdb' NOW (CHAIN 2).
## Chain 2:
## Chain 2: Gradient evaluation took 0 seconds
## Chain 2: 1000 transitions using 10 leapfrog steps per transition would take 0 seconds.
## Chain 2: Adjust your expectations accordingly!
## Chain 2:
## Chain 2:
## Chain 2: Iteration:
                          1 / 2000 [ 0%]
                                            (Warmup)
## Chain 2: Iteration: 200 / 2000 [ 10%]
                                            (Warmup)
## Chain 2: Iteration: 400 / 2000 [ 20%]
                                            (Warmup)
## Chain 2: Iteration: 600 / 2000 [ 30%]
                                            (Warmup)
## Chain 2: Iteration: 800 / 2000 [ 40%]
                                            (Warmup)
## Chain 2: Iteration: 1000 / 2000 [ 50%]
                                            (Warmup)
## Chain 2: Iteration: 1001 / 2000 [ 50%]
                                            (Sampling)
## Chain 2: Iteration: 1200 / 2000 [ 60%]
                                            (Sampling)
## Chain 2: Iteration: 1400 / 2000 [ 70%]
                                            (Sampling)
## Chain 2: Iteration: 1600 / 2000 [ 80%]
                                            (Sampling)
## Chain 2: Iteration: 1800 / 2000 [ 90%]
                                            (Sampling)
## Chain 2: Iteration: 2000 / 2000 [100%]
                                            (Sampling)
## Chain 2:
## Chain 2: Elapsed Time: 0.326 seconds (Warm-up)
## Chain 2:
                           0.068 seconds (Sampling)
## Chain 2:
                           0.394 seconds (Total)
## Chain 2:
## SAMPLING FOR MODEL '8874c61302b41c9e7aa22f87c38effdb' NOW (CHAIN 3).
## Chain 3:
## Chain 3: Gradient evaluation took 0 seconds
## Chain 3: 1000 transitions using 10 leapfrog steps per transition would take 0 seconds.
## Chain 3: Adjust your expectations accordingly!
## Chain 3:
## Chain 3:
## Chain 3: Iteration:
                          1 / 2000 [ 0%]
                                            (Warmup)
## Chain 3: Iteration: 200 / 2000 [ 10%]
                                            (Warmup)
## Chain 3: Iteration: 400 / 2000 [ 20%]
                                            (Warmup)
                        600 / 2000 [ 30%]
## Chain 3: Iteration:
                                            (Warmup)
## Chain 3: Iteration:
                        800 / 2000 [ 40%]
                                            (Warmup)
## Chain 3: Iteration: 1000 / 2000 [ 50%]
                                            (Warmup)
## Chain 3: Iteration: 1001 / 2000 [ 50%]
                                            (Sampling)
## Chain 3: Iteration: 1200 / 2000 [ 60%]
                                            (Sampling)
## Chain 3: Iteration: 1400 / 2000 [ 70%]
                                            (Sampling)
## Chain 3: Iteration: 1600 / 2000 [ 80%]
                                            (Sampling)
## Chain 3: Iteration: 1800 / 2000 [ 90%]
                                            (Sampling)
```

```
## Chain 3: Iteration: 2000 / 2000 [100%]
## Chain 3:
## Chain 3: Elapsed Time: 0.139 seconds (Warm-up)
                           0.076 seconds (Sampling)
## Chain 3:
## Chain 3:
                           0.215 seconds (Total)
## Chain 3:
## SAMPLING FOR MODEL '8874c61302b41c9e7aa22f87c38effdb' NOW (CHAIN 4).
## Chain 4:
## Chain 4: Gradient evaluation took 0 seconds
## Chain 4: 1000 transitions using 10 leapfrog steps per transition would take 0 seconds.
## Chain 4: Adjust your expectations accordingly!
## Chain 4:
## Chain 4:
## Chain 4: Iteration:
                         1 / 2000 [ 0%]
                                            (Warmup)
## Chain 4: Iteration:
                        200 / 2000 [ 10%]
                                            (Warmup)
                        400 / 2000 [ 20%]
## Chain 4: Iteration:
                                            (Warmup)
## Chain 4: Iteration:
                        600 / 2000 [ 30%]
                                            (Warmup)
## Chain 4: Iteration: 800 / 2000 [ 40%]
                                            (Warmup)
## Chain 4: Iteration: 1000 / 2000 [ 50%]
                                            (Warmup)
## Chain 4: Iteration: 1001 / 2000 [ 50%]
                                            (Sampling)
## Chain 4: Iteration: 1200 / 2000 [ 60%]
                                            (Sampling)
## Chain 4: Iteration: 1400 / 2000 [ 70%]
                                            (Sampling)
## Chain 4: Iteration: 1600 / 2000 [ 80%]
                                            (Sampling)
## Chain 4: Iteration: 1800 / 2000 [ 90%]
                                            (Sampling)
## Chain 4: Iteration: 2000 / 2000 [100%]
                                            (Sampling)
## Chain 4:
## Chain 4: Elapsed Time: 0.162 seconds (Warm-up)
## Chain 4:
                           0.071 seconds (Sampling)
## Chain 4:
                           0.233 seconds (Total)
## Chain 4:
# Print the fitted model
print(fity,digits_summary=3)
## Inference for Stan model: 8874c61302b41c9e7aa22f87c38effdb.
## 4 chains, each with iter=2000; warmup=1000; thin=1;
## post-warmup draws per chain=1000, total post-warmup draws=4000.
##
##
                                      2.5%
                                                25%
                                                         50%
                                                                  75%
                                                                          97.5%
              mean se_mean
                              sd
## mu
            13.137
                     0.002 0.114
                                    12.920
                                             13.060
                                                      13.139
                                                                13.213
                                                                         13.360
## sigma2
             1.565
                     0.001 0.065
                                    1.449
                                              1.519
                                                       1.563
                                                                1.608
                                                                          1.703
                     0.001 0.057
                                    0.093
## phi
             0.200
                                              0.160
                                                       0.199
                                                                 0.238
                                                                          0.314
## lp__
          -284.672
                     0.027 1.235 -287.725 -285.222 -284.368 -283.762 -283.242
##
          n_eff Rhat
           3853 1.001
## mu
## sigma2
           4391 0.999
           4325 1.000
## phi
           2037 1.001
## lp__
##
## Samples were drawn using NUTS(diag_e) at Mon May 29 21:52:09 2023.
## For each parameter, n_eff is a crude measure of effective sample size,
## and Rhat is the potential scale reduction factor on split chains (at
## convergence, Rhat=1).
```

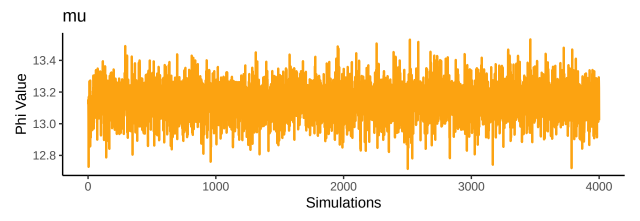
```
##
                Mean Credible_Interval_Lower Credible_Interval_Upper
## mu
                                 12.91954935
          13.1370383
                                                           13.3604494
## sigma2 1.5653937
                                   1.44876753
                                                            1.7027939
           0.2002713
                                   0.09332073
                                                            0.3138576
## phi
##
          Effective_Samples
## mu
                   3593.851
## sigma2
                   4055.560
## phi
                   4246.170
```

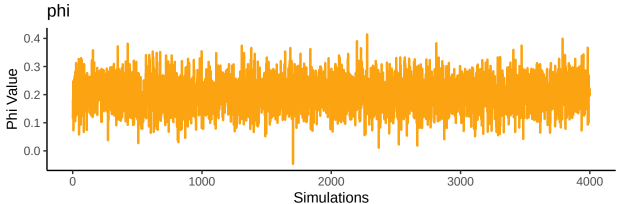
In the table above, it is observed that the second simulated AR(1) process exhibits some differences compared to the first one. While the estimates for σ and ϕ are relatively accurate, the estimate for μ appears to deviate slightly from the true value. This discrepancy suggests that the sampling algorithm may have encountered challenges in accurately capturing the true mean parameter.

There could be several reasons for this discrepancy. It is possible that the non-informative priors chosen for the parameters might not have been appropriate, leading to a bias in the estimation of μ . Additionally, the specific characteristics of the second AR(1) process, such as its underlying dynamics and data distribution, might have posed challenges for the sampling algorithm in accurately estimating μ .

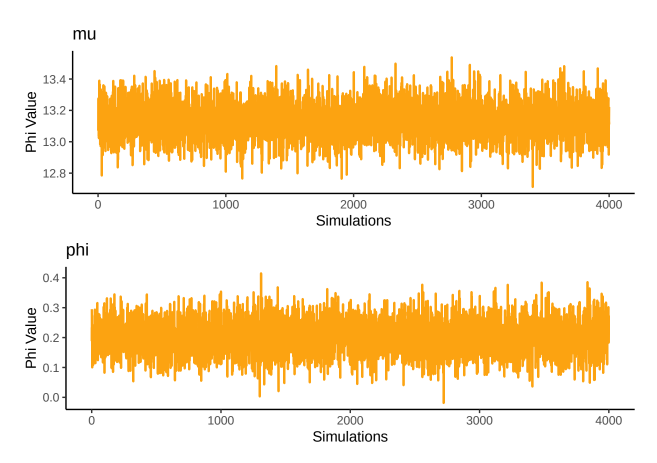
Part ii For each of the two data sets, evaluate the convergence of the samplers and plot the joint posterior of μ and ϕ . Comments?

First sample: For $x_{1:T}$ with $\phi = 0.2$





Second sample: For $y_{1:T}$ with $\phi = 0.95$



Examining the above results, it is evident that the posterior estimates for both μ and ϕ exhibit convergence for each of the two data sets. This convergence indicates that the Markov chain has explored the parameter space sufficiently and reached a stable distribution.

References:

1- Stan Model

```
#### In case of normal
# library(rstan)
y=c(4,5,6,4,0,2,5,3,8,6,10,8)
# N=length(y)
# StanModel =
# data {
# int<lower=0> N; // Number of observations
# int<lower=0> y[N]; // Number of flowers
# }
# parameters {
# real mu;
# real<lower=0> sigma2;
# }
# model {
# mu ~ normal(0,100); // Normal with mean 0, st.dev. 100
# sigma2 ~ scaled_inv_chi_square(1,2); // Scaled-inv-chi2 with nu 1,sigma 2
```

```
# for(i in 1:N){
# y[i] ~ normal(mu,sqrt(sigma2));
# }
# }'
######## Multilevel normal
# StanModel <- '
# data {
# int<lower=0> N; // Number of observations
# int<lower=0> y[N]; // Number of flowers
# int<lower=0> P; // Number of plants
# }
# transformed data {
# int<lower=0> M; // Number of months
# M = N / P;
# }
# parameters {
# real mu;
# real<lower=0> sigma2;
# real mup[P];
# real sigmap2[P];
# }
# model {
# mu ~ normal(0,100); // Normal with mean 0, st.dev. 100
# sigma2 ~ scaled_inv_chi_square(1,2); // Scaled-inv-chi2 with nu 1, sigma 2
# for(p in 1:P){
# mup[p] ~ normal(mu,sqrt(sigma2));
# for(m in 1:M) {
\# y[M*(p-1)+m] \sim normal(mup[p], sqrt(sigmap2[p]));
# }
# }
# }'
#########Possion
# StanModel <- '
# data {
# int<lower=0> N; // Number of observations
# int<lower=0> y[N]; // Number of flowers
\# int<lower=0> P; // Number of plants
# }
# transformed data {
# int<lower=0> M; // Number of months
# M = N / P;
# }
# parameters {
# real mu;
# real<lower=0> sigma2;
# real mup[P];
# }
# model {
# mu ~ normal(0,100); // Normal with mean 0, st.dev. 100
# sigma2 ~ scaled_inv_chi_square(1,2); // Scaled-inv-chi2 with nu 1, sigma 2
# for(p in 1:P){
# mup[p] ~ lognormal(mu,sqrt(sigma2)); // Log-normal
# for(m in 1:M) {
```

```
# y[M*(p-1)+m] ~ poisson(mup[p]); // Poisson
# }
# }
# }'
```

1- Example code from lec notes

```
### The original Code:
# Direct vs Gibbs sampling for bivariate normal distribution
# Initial setting
mu1 <- 1
mu2 <- 1
rho <- 0.9
mu <- c(mu1,mu2)
Sigma = matrix(c(1,rho,rho,1),2,2)
nDraws <- 500 # Number of draws
library(MASS) # To access the murnorm() function
# Direct sampling from bivariate normal distribution
directDraws <- mvrnorm(nDraws, mu, Sigma)</pre>
# Gibbs sampling
gibbsDraws <- matrix(0,nDraws,2)</pre>
theta2 <- 0 # Initial value for theta2
for (i in 1:nDraws){
  # Update theta1 given theta2
  theta1 <- rnorm(1, mean = mu1 + rho*(theta2-mu2), sd = sqrt(1-rho**2))
  gibbsDraws[i,1] <- theta1</pre>
  # Update theta2 given theta1
  theta2 <- rnorm(1, mean = mu2 + rho*(theta1-mu1), sd = sqrt(1-rho**2))
  gibbsDraws[i,2] <- theta2</pre>
}
a_Direct <- acf(directDraws[,1])</pre>
a_Gibbs <- acf(gibbsDraws[,1])</pre>
IF_Gibbs <- 1+2*sum(a_Gibbs$acf[-1])</pre>
par(mfrow=c(2,4))
# DIRECT SAMPLING
plot(1:nDraws, directDraws[,1], type = "l",col="blue") # traceplot of direct draws
hist(directDraws[,1],col="blue") # histogram of direct draws
) # Cumulative mean value of theta1, direct draws
cusumData <- cumsum(directDraws[,1])/seq(1,nDraws</pre>
plot(1:nDraws, cusumData, type = "l", col="blue")
```

- 2- Stan Development Team. (2021). Stan user's guide. Retrieved from https://mc-stan.org/docs/2_27/stan-users-guide/index.html.
- 3- Bertil Wegmann (2023). Bayesian Learning [Lecture notes]. 732A73, Department of Computer and Information Science,LiU University.

Code Appendix

```
set.seed(123456)
knitr::opts chunk$set(echo = TRUE)
library(LaplacesDemon)
library(mvtnorm)
library(MASS)
library(ggplot2)
library(gridExtra)
library(rstan)
\# we start by reading our dataset using the readRDS command in R
df <- data.frame(x=readRDS("Precipitation.rds"))</pre>
# in the task we looking at natural log of the daily precipitation
#lny1, lny2, lny3,...lnyn follows N(mu, sigma)
# We add a new var called logx
df$logx<- log(df$x)</pre>
sigma2_0<- var(df$logx) # Sample var
tau2_0<- 1# arbirtry initail value let it be 1
n<- nrow(df)# Sample size</pre>
mu 0<- mean(df$logx)#sample mean
w<- (n/sigma2_0)/((n/sigma2_0) + (1/tau2_0))# value used to calculate mu_n
v_0<- 1 # arbirtry initail value let it be 1
v_n<- v_0+n
```

```
mu_n \leftarrow w*(mean(df$logx))+ (1-w)*mu_0
##### This part of the code has been modified#####
tau2_n<- 1/((n/sigma2_0)+(1/tau2_0))############ shoud be 1/tau
nDraws<- 1000
gibbsDraws <- matrix(0,nDraws,2)</pre>
### From lec notes we can use this part of the code
 ##### This part of the code has been modified#####
w<- (n/scale)/((n/scale) + (1/tau2_0)) ############
 mu_n<- w*(mean(df$logx))+ (1-w)*mu_0 ########### We need to update those parameters as well
 # Update theta1 ----> mu given theta2
 theta1 <- rnorm(1, mean = mu_n, sd = (tau2_n))
 gibbsDraws[i,1] <- theta1</pre>
 scale \leftarrow ((v_0*sigma2_0)+(sum((df$logx-theta1)^2)))/(n+v_0)
 # Update theta2 ----> sigma2 given theta1
 theta2 <- rinvchisq(1,v_n,scale=scale)</pre>
 gibbsDraws[i,2] <- theta2</pre>
## Finding the acf plots and value
a_Gibbs_mu <- acf(gibbsDraws[,1])</pre>
a_Gibbs_sigma <- acf(gibbsDraws[,2])</pre>
IF_Gibbs_mu <- 1+2*sum(a_Gibbs_mu$acf[-1])</pre>
IF_Gibbs_sigma <- 1+2*sum(a_Gibbs_sigma$acf[-1])</pre>
### From lec notes we can use this part of the code
plot_df<- as.data.frame(cbind(1:nDraws,gibbsDraws))</pre>
# Plot for Mu trajectories of the sampled Markov chains
p1<-ggplot(plot_df,aes(x=V1))+geom_line(aes(y=V2), color='#FCA311', size=.8)+
 annotate(geom = "text", x = 500, y = 1.312,
         label = paste0("Inefficiency Factor ",format(round(IF_Gibbs_mu, 3),
                                                  nsmall = 3)))+
 labs(title = 'Mu trajectories of the sampled Markov chains',
      x= 'nDraws', y='Mu')+ theme_classic()
# Plot for Sigma trajectories of the sampled Markov chains
p2<-ggplot(plot_df,aes(x=V1))+geom_line(aes(y=V3), color='#FCA311', size=.8)+
 annotate(geom = "text", x = 500, y = 2,
         label = paste0("Inefficiency Factor ",format(round(IF_Gibbs_sigma, 3), nsmall = 3)))+
 labs(title = 'Sigma2 trajectories of the sampled Markov chains',
      x= 'nDraws', y='Sigma')+ theme_classic()
p1
p2
```

```
# mean and Sd from the results of the Gibbs sampler
#plot_df$V2 represent mean and plot_df$V3 Var
# The resulting posterior predictive density
post_pred<- rnorm(1000, mean = plot_df$V2, sd = sqrt(plot_df$V3))</pre>
y <- exp(post_pred)</pre>
# Now we plot histogarm of the data and the posterior predictive denstity
#data frame to store the density values
de_df<- data.frame(x=y)</pre>
ggplot() +
  geom_histogram(data=df,aes(x = x,y=..density..),linetype=1,
                fill='#14213D',bins = 20)+
  geom_density(data=de_df,aes(x=y), color='#FCA311', size=1,
              fill='#FCA311',alpha=.5)+
 labs(title = "Histogram of Daily Precipitation",
      subtitle = "Line of Posterior predictive density",
      x = "Precipitation",
      y = "Density") + xlim(0.1,50)+ theme_classic()
#First we upload our data
df1<- read.table("eBayNumberOfBidderData.dat",header = T)</pre>
#We run a possion model using the function glm in r,
#note that we exclude the intercept and we use family possion
model <- glm(nBids ~ ., data = df1[,-2], family = poisson)</pre>
summary(model)
### Prior and data inputs ###
Covs <- c(2:10) # Select which covariates/features to include
standardize <- F # If TRUE, covariates/features are standardized to mean 0 and variance 1
Nobs <- dim(df1)[1] # number of observations
y <- df1nBids # y=1 if the women is working, otherwise y=0.
x <- as.matrix(df1[,Covs]) # Covs matrix 7*7
Xnames <- colnames(x)</pre>
# Standraizing the covs matrix
if (standardize){
  Index <- 2:(length(Covs)-1)</pre>
 x[,Index] <- scale(x[,Index])
Npar \leftarrow dim(x)[2]
# Setting up the prior
mu <- c(rep(0,Npar)) # Prior mean vector</pre>
Sigma <- 100*solve(t(x)%*%x) # Prior covariance matrix
# Functions that returns the log posterior for the logistic and
#probit regression.
# First input argument of this function must be the parameters we optimize on,
# i.e. the regression coefficients beta.
logPossion <- function(beta,y,x,mu,Sigma){</pre>
  logLik <- -sum(exp(x%*%beta)) + sum(y%*%(x%*%beta))# <-----
  #We change on this line
```

```
# The first term, -sum(exp(Xbeta)), calculates the sum of the exponential of
 # the linear predictor Xbeta over all observations. This term represents the
 # log-likelihood contribution from the expected
 #counts in the Poisson distribution.
 # The second term, sum(y(Xbeta)), calculates the sum of the
 #observed response variable
 # y multiplied by the linear predictor Xbeta over all
 #observations. This term represents
 # the log-likelihood contribution from
 #the observed counts in the Poisson distribution.
 # By subtracting the first term from the second term,
 #we obtain the log-likelihood of
 # the Poisson regression model given the data
 #and the regression coefficients.
 #if (abs(logLik) == Inf) logLik = -20000;
 # Likelihood is not finite, stear the optimizer away from here!
 if (abs(logLik) == Inf){
   logLik <- -20000
 logPrior <- dmvnorm(beta, mu, Sigma, log=TRUE)</pre>
 return(logLik + logPrior)
# Select the initial values for beta
initVal <- matrix(0,1,Npar)</pre>
# The argument control is a list of options to the
#optimizer optim, where fnscale=-1 means that we minimize
# the negative log posterior. Hence, we maximize the log posterior.
OptimRes <- optim(initVal,</pre>
                 logPossion,
                 gr=NULL,
                 y=y,
                 x=x,
                 mu=mu,
                 Sigma=Sigma,
                 method=c("BFGS"),
                 control=list(fnscale=-1),
                 hessian=TRUE)
# Printing the results to the screen
names(OptimRes$par) <- Xnames # Naming the coefficient by covariates
# Computing approximate standard deviations.
approxPostStd <- sqrt(diag(solve(-OptimRes$hessian)))</pre>
names(approxPostStd) <- Xnames # Naming the coefficient by covariates</pre>
print('The posterior mode is:')
print(OptimRes$par)
print('The Hessian Matrix:')
print(OptimRes$hessian)
print('The approximate posterior standard deviation is:')
print(approxPostStd)
```

```
RWMSampler_old<- function(logPostFunc,nDraws,c,y,x,mu,Sigma){</pre>
  # First we buld our data frame of samples
  sample <- data.frame(matrix(nrow = nDraws, ncol = ncol(x)))</pre>
  colnames(sample) <- colnames(x)</pre>
  # The inital sample value c here represent a tuning parameter
  sample[1,] <- mvrnorm(1, posteriorMode, c*postCov)</pre>
  # Now we implement the Metropolis-Hastings
  #in which we generate samples from the proposal distribution in this case
  # We look at the results of the first sample
  #as theta_i-1 pluged in munorm to get theta_i and the we use the values in
  # our proposed logPostFunc
  counter <- 1
  i = 1
  while (counter < nDraws) {</pre>
    theta_old<-as.numeric(sample[counter,])</pre>
    theta_new<-mvrnorm(1,theta_old,c*postCov)</pre>
  # We define th our accept/reject threshold
    th<-runif(1,0,1)
  # now we find the value of the target/proposed distribution
    proposed<- logPostFunc(theta_new,y = y,</pre>
                           x = x,
                           mu = posteriorMode,
                           Sigma = postCov)
    target<- logPostFunc(theta_old,y = y,</pre>
                         x = x
                         mu = posteriorMode,
                         Sigma = postCov)
    # the ratio of posterior densities in the Metropolis acceptance probability
    if (th<min(1,exp(proposed-target))) {</pre>
      counter=counter+1
      sample[counter,]<-theta_new</pre>
    }
return(sample)
RWMSampler<- function(logPostFunc,nDraws,c,y,x,mu,Sigma,brnin){</pre>
  # First we buld our data frame of samples
  sample <- data.frame(matrix(0,nrow = nDraws - brnin, ncol = ncol(x)))</pre>
  colnames(sample) <- colnames(x)</pre>
  # The inital sample value c here represent a tuning parameter
  # sample[1,] <- murnorm(1, posteriorMode, c*postCov)</pre>
  # Now we implement the Metropolis-Hastings
  #in which we generate samples from the proposal distribution in this case
  # We look at the results of the first sample
  #as theta_i-1 pluged in monorm to get theta_i and the we use the values in
  # our proposed logPostFunc
  theta_old<-as.numeric(sample[1,])</pre>
  for (i in 1:nDraws) {
    theta_new<-mvrnorm(1,theta_old,c*postCov)</pre>
    # We define th our accept/reject threshold
    th<-runif(1,0,1)
```

```
# now we find the value of the target/proposed distribution
    proposed<- logPostFunc(theta_new,y = y,</pre>
                            x = x
                            mu = posteriorMode,
                            Sigma = postCov)
    target<- logPostFunc(theta_old,y = y,</pre>
                          x = x,
                          mu = posteriorMode,
                          Sigma = postCov)
    # the ratio of posterior densities in the Metropolis acceptance probability
    if (th<exp(proposed-target)) {</pre>
      theta_old<-theta_new
    }
    if (i> brnin) {
      sample[i-brnin,]<-theta_old</pre>
    }
  }
  return(sample)
nDraws=10000
c=.5
brnin=100 ##adding this new parameter for buirnin interval
posteriorMode=OptimRes$par
postCov=solve(-OptimRes$hessian)
res <- RWMSampler(logPostFunc = logPossion,
                                  nDraws = nDraws,
                                    c=c,
                                    y = y,
                                    x = x
                                    mu = posteriorMode,
                                    Sigma = postCov,
                                    brnin = brnin )
### Changing the plot to be trace plot instead of density
names <- colnames (res)
p_fun<- function(coln){</pre>
  plt <- ggplot(res,aes_string(1:(nDraws-brnin),y = coln)) +</pre>
    geom_line( color='#FCA311', size=.8)+
      labs(x= 'nDraws', y=coln)
  # We look
  plt
}
plot(arrangeGrob(grobs = lapply(names, p_fun)))
#First we estimate the betas from our RWMSampler function
betas<- as.matrix(res)</pre>
# Input data
x_new \leftarrow as.matrix(c(1,1,0,1,0,1,0,1.2,0.8))
##### This code has been modified
# calculating the bet for Poisson since we have our y follow poisson
beta_pois<-exp(betas %*% x_new)</pre>
# Finding number of bidders using poisson function
nbiders<-data.frame(x=rpois(length(beta_pois),beta_pois))</pre>
```

```
ggplot(nbiders, aes(x = x)) +
  geom_histogram(aes(y=..density..),linetype=1,fill='#14213D',bins = 20)+
    labs(x = 'Number of bidders', y = 'density',
           title ='Plot of the predictive distribution for Number of bidders')
ar_process<- function(phi,mu, sigma,t){</pre>
  x_t<-c()
  x_t[1] <-mu
  for(i in 2:t){
    e<- rnorm(1,0,sqrt(3))
    x_t[i] \leftarrow mu + (phi * (x_t[i-1] - mu)) + e
  return(x_t)
phi < -seq(-1,1,by=.5)
res<- list()
for (i in phi) {
  res1 <- ar_process(i, 13, 3, 300)
  res[[paste0("Phi_", i)]] <- res1</pre>
}
res<- data.frame(res)
names<-colnames(res)</pre>
p_fun<- function(coln){</pre>
  plt \leftarrow ggplot(res,aes(x = 1:300)) +
    geom_line(aes_string(y = coln),color='#FCA311', size=.8)+
  labs(title = coln,
       x= 'Simulations', y='Phi Value')+ theme_classic()
  plt
plot(arrangeGrob(grobs = lapply(names, p_fun)))
data {
  int<lower=0> N;
  vector[N] y;
parameters {
  real alpha;
  real beta;
  real<lower=0> sigma;
model {
  for (n in 2:N) {
    y[n] ~ normal(alpha + beta * y[n-1], sigma);
}
data {
  int<lower=0> N;
  vector[N] y;
parameters {
  real alpha;
  real beta;
  real<lower=0> sigma;
```

```
}
model {
   y[2:N] \sim normal(alpha + beta * y[1:(N - 1)], sigma);
##### Example from notes #####
# library(rstan)
# y=c(4,5,6,4,0,2,5,3,8,6,10,8)
# N=length(y)
# StanModel = '
# data {
   int<lower=0> N; // Number of observations
    int<lower=0> y[N]; // Number of flowers
# }
# parameters {
# real mu;
#
  real<lower=0> sigma2;
# }
# model {
# mu ~ normal(0,100); // Normal with mean 0, st.dev. 100
   sigma2 ~ scaled_inv_chi_square(1,2); // Scaled-inv-chi2 with nu 1,sigma 2
#
  for(i in 1:N){
     y[i] ~ normal(mu,sqrt(sigma2));
#
# }'
#
# data \leftarrow list(N=N, y=y)
# warmup <- 1000
# niter <- 2000
\# fit <- stan(model\_code=StanModel, data=data, warmup=warmup, iter=niter, chains=4)
# # Print the fitted model
# print(fit, digits_summary=3)
# # Extract posterior samples
# postDraws <- extract(fit)</pre>
# # Do traceplots of the first chain
\# par(mfrow = c(1,1))
# plot(postDraws$mu[1:(niter-warmup)], type="l", ylab="mu", main="Traceplot")
# # Do automatic traceplots of all chains
# traceplot(fit)
# # Bivariate posterior plots
# pairs(fit)
# ##################################
StanModel = '
data {
 int<lower=0> N; // Number of observations
 vector[N] y;
}
parameters {
 real mu;
 real<lower=0> sigma2;
 real<lower=-1, upper=1> phi;
 //To enforce the estimation of a
  //stationary AR(1) process, the slope
```

```
//coefficient beta may be constrained with bounds as follows.
model {
  mu ~ normal(0,100); // Normal with mean 0, st.dev. 100
  sigma2 ~ scaled_inv_chi_square(1,2); // Scaled-inv-chi2 with nu 1,sigma 2
  // Changing the model to be y_i-mu
 y[2:N] ~ normal(mu + phi * (y[1:(N - 1)]-mu), sqrt(sigma2^2));
# Simulate of x
ar_x<-ar_process(.2, 13, 3, 300)
#From lec notes we have the stanmodel function defined as
y=ar_x
N=length(y)
data <- list(N=N, y=y)
warmup <- 1000
niter <- 2000
fitx <- stan(model_code=StanModel,data=data,warmup=warmup,iter=niter,chains=4)</pre>
# Print the fitted model
print(fitx,digits_summary=3)
samples_x <-extract(fitx, pars = c("mu", "sigma2", "phi"))</pre>
# Compute the mean
mean_x <- sapply(samples_x, mean)</pre>
# 95% credible intervals
intv_x <- sapply(samples_x,function(samples) quantile(samples,c(0.025, 0.975)))
# effective posterior samples
eff_samp_x <- summary(fitx)$summary[1:3,"n_eff"]</pre>
result <- data.frame( Mean = mean_x,
                      Credible_Interval_Lower = intv_x[1, ],
                      Credible_Interval_Upper = intv_x[2, ],
                      Effective_Samples = eff_samp_x)
print(result)
# Simulate of y
ar_y<-ar_process(.95, 13, 3, 300)
#From lec notes we have the stanmodel function defined as
#y=ar_y
N=length(y)
data <- list(N=N, y=y)</pre>
warmup <- 1000
niter <- 2000
fity <- stan(model_code=StanModel,data=data, warmup=warmup,iter=niter,chains=4)</pre>
# Print the fitted model
print(fity,digits_summary=3)
samples_y <-extract(fity, pars = c("mu", "sigma2", "phi"))</pre>
# Compute the mean
mean_y <- sapply(samples_y, mean)</pre>
# 95% credible intervals
intv_y <- sapply(samples_y,function(samples) quantile(samples,c(0.025, 0.975)))
# effective posterior samples
```

```
eff_samp_y <- summary(fitx)$summary[1:3,"n_eff"]</pre>
result <- data.frame( Mean = mean_y,
                      Credible_Interval_Lower = intv_y[1, ],
                      Credible_Interval_Upper = intv_y[2, ],
                      Effective_Samples = eff_samp_y)
print(result)
df x <-as.data.frame(extract(fitx, pars = c("mu", "phi")))</pre>
names<-colnames(df_x)</pre>
ln<-length(df_x[,1])</pre>
p_fun<- function(coln){</pre>
  plt \leftarrow ggplot(df_x,aes(x = 1:ln)) +
    geom_line(aes_string(y = coln),color='#FCA311', size=.8)+
    labs(title = coln,
         x= 'Simulations', y='Phi Value')+ theme_classic()
 plt
}
plot(arrangeGrob(grobs = lapply(names, p_fun)))
df_y <-as.data.frame(extract(fity, pars = c("mu", "phi")))</pre>
names<-colnames(df_y)</pre>
ln<-length(df_y[,1])</pre>
p_fun<- function(coln){</pre>
  plt \leftarrow ggplot(df_y,aes(x = 1:ln)) +
    geom_line(aes_string(y = coln),color='#FCA311', size=.8)+
    labs(title = coln,
         x= 'Simulations', y='Phi Value')+ theme_classic()
 plt
plot(arrangeGrob(grobs = lapply(names, p_fun)))
#### In case of normal
# library(rstan)
 # y=c(4,5,6,4,0,2,5,3,8,6,10,8) 
# N=length(y)
# StanModel = '
# data {
# int<lower=0> N; // Number of observations
# int<lower=0> y[N]; // Number of flowers
# }
# parameters {
# real mu;
# real<lower=0> sigma2;
# }
# model {
# mu ~ normal(0,100); // Normal with mean 0, st.dev. 100
# sigma2 ~ scaled_inv_chi_square(1,2); // Scaled-inv-chi2 with nu 1,sigma 2
# for(i in 1:N){
# y[i] ~ normal(mu,sqrt(siqma2));
# }
# }'
######## Multilevel normal
# StanModel <- '
```

```
# data {
# int<lower=0> N; // Number of observations
# int<lower=0> y[N]; // Number of flowers
# int<lower=0> P; // Number of plants
# }
# transformed data {
# int<lower=0> M; // Number of months
# M = N / P;
# }
# parameters {
# real mu;
# real<lower=0> sigma2;
# real mup[P];
# real sigmap2[P];
# }
# model {
# mu ~ normal(0,100); // Normal with mean 0, st.dev. 100
# sigma2 ~ scaled_inv_chi_square(1,2); // Scaled-inv-chi2 with nu 1, sigma 2
# for(p in 1:P){
# mup[p] ~ normal(mu,sqrt(sigma2));
# for(m in 1:M) {
\# y[M*(p-1)+m] \sim normal(mup[p], sqrt(sigmap2[p]));
# }
# }
# }'
#########Possion
# StanModel <- '
# data {
# int<lower=0> N; // Number of observations
# int<lower=0> y[N]; // Number of flowers
# int<lower=0> P; // Number of plants
# }
# transformed data {
# int<lower=0> M; // Number of months
# M = N / P;
# }
# parameters {
# real mu;
# real<lower=0> sigma2;
# real mup[P];
# }
# model {
# mu ~ normal(0,100); // Normal with mean 0, st.dev. 100
# sigma2 ~ scaled_inv_chi_square(1,2); // Scaled-inv-chi2 with nu 1, sigma 2
# for(p in 1:P){
# mup[p] ~ lognormal(mu,sqrt(sigma2)); // Log-normal
# for(m in 1:M) {
\# y[M*(p-1)+m] \sim poisson(mup[p]); // Poisson
# }
# }
# }'
### The original Code:
```

```
# Direct vs Gibbs sampling for bivariate normal distribution
# Initial setting
mu1 <- 1
mu2 <- 1
rho <- 0.9
mu <- c(mu1,mu2)
Sigma = matrix(c(1,rho,rho,1),2,2)
nDraws <- 500 # Number of draws
library(MASS) # To access the murnorm() function
# Direct sampling from bivariate normal distribution
directDraws <- mvrnorm(nDraws, mu, Sigma)</pre>
# Gibbs sampling
gibbsDraws <- matrix(0,nDraws,2)</pre>
theta2 <- 0 # Initial value for theta2
for (i in 1:nDraws){
  # Update theta1 given theta2
  theta1 <- rnorm(1, mean = mu1 + rho*(theta2-mu2), sd = sqrt(1-rho**2))
  gibbsDraws[i,1] <- theta1</pre>
  # Update theta2 given theta1
  theta2 <- rnorm(1, mean = mu2 + rho*(theta1-mu1), sd = sqrt(1-rho**2))
  gibbsDraws[i,2] <- theta2</pre>
}
a_Direct <- acf(directDraws[,1])</pre>
a_Gibbs <- acf(gibbsDraws[,1])</pre>
IF_Gibbs <- 1+2*sum(a_Gibbs$acf[-1])</pre>
par(mfrow=c(2,4))
# DIRECT SAMPLING
plot(1:nDraws, directDraws[,1], type = "l",col="blue") # traceplot of direct draws
hist(directDraws[,1],col="blue") # histogram of direct draws
) # Cumulative mean value of theta1, direct draws
cusumData <- cumsum(directDraws[,1])/seq(1,nDraws</pre>
plot(1:nDraws, cusumData, type = "l", col="blue")
barplot(height = a_Direct$acf[-1],col="blue") # acf for direct draws
# GIBBS SAMPLING
plot(1:nDraws, gibbsDraws[,1], type = "l",col="red") # traceplot of Gibbs draws
hist(gibbsDraws[,1],col="red") # histogram of Gibbs draws
# Cumulative mean value of theta1, Gibbs draws
cusumData = cumsum(gibbsDraws[,1])/seq(1,nDraws)
```