Applied Bayesian Analysis : NCSU ST 540

Midterm2

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Test section - VAR(1) in JAGS

This section is a test section where we generate and fit a vector autoregressive model - $VAR(1) \in \mathbf{R}^6$ given by

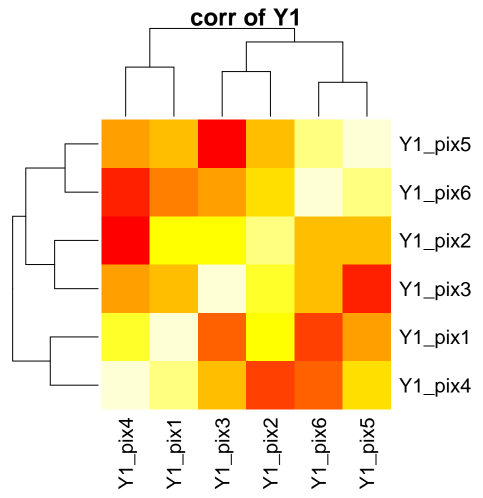
$$y_t = \nu + \rho * y_{t-1} + \epsilon$$

$$\epsilon \sim N(0, \Sigma)$$

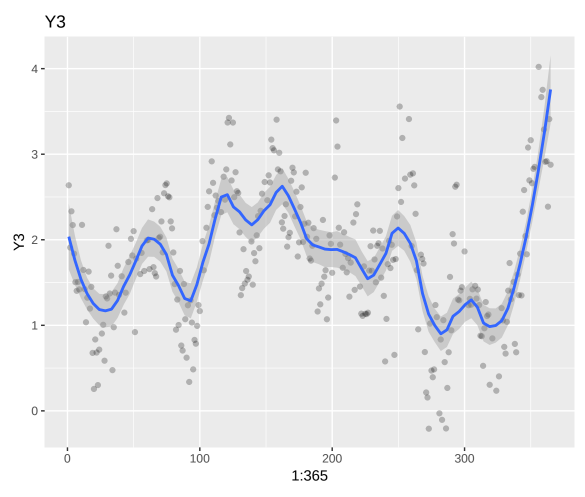
We use the y1 data to calculate a NaN firendly sample covariance and then we find the nearest positive semidefinite matrix to use to generate data for the model.

Notes - imputation is not working for this model - this section seeks to find the parameters that best explain the data

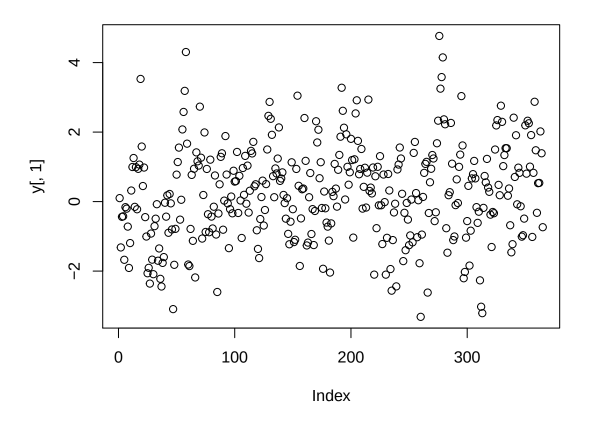
```
library(rjags)
library(coda)
library(modeest)
library(MASS)
load("E2.RData")
DEBUG <- TRUE
if (DEBUG) {
    nSamples <- 5000
    n.chains <- 1
} else {
    nSamples <- 40000
    n.chains <- 8
}
cor.y1 <- cor(Y1, use = "pairwise.complete.obs")</pre>
cov.y1 <- cov(Y1, use = "pairwise.complete.obs")</pre>
heatmap(cor.y1, main = "corr of Y1")
```



```
ggplot(data.frame(Y3 = Y3), aes(x = 1:365,
    y = Y3)) + geom_point(alpha = 0.25) +
    geom_smooth(method = "loess", span = 0.22) +
    ggtitle("Y3")
```



```
N <- nrow(Y1)</pre>
p = 6
Y1.scaled <- scale(Y1)
# Try to simulate using the corr
#install.packages("Matrix")
library("Matrix")
sig <- nearPD(cov.y1)</pre>
N = 365
Sigma = sig$mat
rho = .6
nu = matrix(rep(.1,6), p, 1)
y = matrix(NA, N,p)
y[1,] = nu
for(t in 2:N)
 y[t,] = mvrnorm(1, nu + rho * y[t-1,], Sigma)
plot(y[,1])
```



```
# Jags code to fit the model to the simulated data
model_code = '
model
{
  # Likelihood
  for (t in 2:N)
    y[t, ] ~ dmnorm(mu[t, ], precisionAR)
    mu[t, 1:p] \leftarrow nu + rho * y[t-1,]
  precisionAR ~ dwish(I, p+1)
  Sigma <- inverse(precisionAR)</pre>
  # Priors
  rho \sim dunif(-1, 1)
  for(i in 1:p)
  {
    nu[i] ~ dnorm(0, 0.01)
  }
```

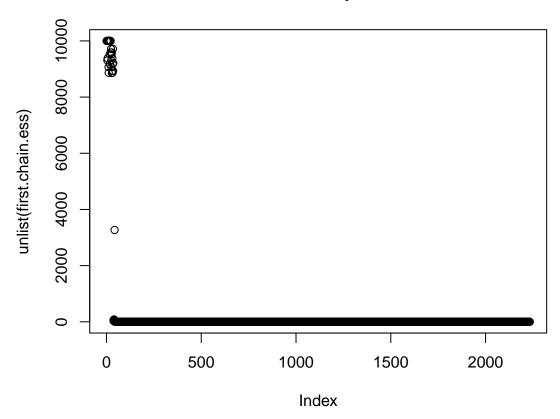
```
# Missing data model for y - not working in JAGS
  #for(i in 1:N)
  #{
    #y[i,1:p]~dmnorm(x_mn[],x_prec[,])
  # Priors for missing-data model parameters
  for(j in 1:p)
  {
    x_mn[j]~dnorm(0,0.01)
  x_prec[1:p,1:p]~dwish(R[,],k)
  x_cov[1:p,1:p]<-inverse(x_prec[,])</pre>
  k < -p+0.1
  for(j1 in 1:p)
    for(j2 in 1:p)
      R[j1,j2] \leftarrow 0.1*equals(j1,j2)
    }
  }
}
# Set up the data
model_data = list(N = N, p = p, y = y, I = diag(p))
# Choose the parameters to watch
model_parameters = c("nu", "rho", "Sigma", "y")
model <- jags.model(textConnection(model_code),data = model_data,n.chains = n.chains)#Compile.
## Compiling model graph
##
      Resolving undeclared variables
##
      Allocating nodes
## Graph information:
      Observed stochastic nodes: 364
##
##
      Unobserved stochastic nodes: 15
##
      Total graph size: 1207
##
## Initializing model
update(model, nSamples, progress.bar="none"); # Burnin
if (FALSE)
₹
  out.coda <- coda.samples(model, variable.names=model_parameters,n.iter=2*nSamples)
  save(out.coda,file = "out.coda_JAGS_VAR1.RData")
}else{
  load("out.coda_JAGS_VAR1.RData")
}
```

```
#plot(out.coda )

if(n.chains > 1)
{
    gelman.srf <-gelman.diag(out.coda)
    plot(gelman.srf$psrf,main = "Gelman Diagnostic")
}

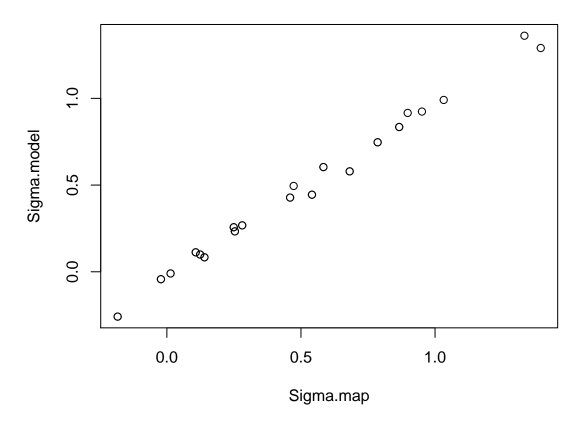
chains.ess <- lapply(out.coda,effectiveSize)
first.chain.ess <- chains.ess[1]
plot(unlist(first.chain.ess), main="Effective Sample Size")</pre>
```

Effective Sample Size



```
chain <- out.coda[[1]]
posterior.means <- list()
posterior.modes <- list()
for( i in 1:length(colnames(chain)) )
{
    colname <- colnames(chain)[i]
    samples <- chain[,i]
    posterior.means[colname] <-mean(samples)</pre>
```

```
posterior.modes[colname] <-mlv(samples)$M
}
# plot(posterior.means, posterior.modes) # Nice and unimodal
Sigma.map <- unlist(posterior.means)[1:36]
Sigma.model<- unlist(sig$mat)
plot(Sigma.map,Sigma.model)</pre>
```



```
rho.map <- unlist(posterior.means)[43]
nu.map <- unlist(posterior.means)[37:42]
# y.map <- matrix(unlist(posterior.means)[44:2233],ncol=6, byrow=FALSE)
# plot(y[,1]-y.map[,1])</pre>
```

Missing data imputation in Openbugs

Here we try to implement missing data imputation in OpenBugs. We use the data set y1 and fit at multivariate model to the data

$$\theta[t] \sim N(Y1[t], \Sigma)$$

we use BUGS and are imputing the missing data in Y1.

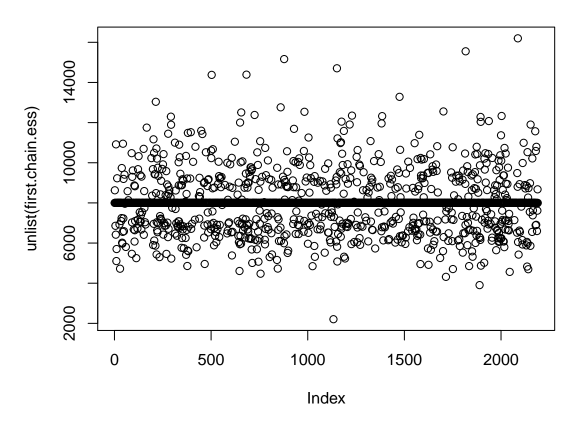
Notes: - This does not take into account the temporal correlation - Openbugs was VERY HARD TO FIND on the internet - their website is down. - We can not yetI'm happ port the VAR(1) model toOpenBugs for further refinement.

```
rm(list = ls())
setwd("d:/brucebcampbell-git/bayesian-learning-with-R")
load("E2.RData")
library(R2OpenBUGS)
library(rjags)
library(coda)
library(modeest)
N \leftarrow nrow(Y1)
p = 6
x \leftarrow scale(Y1)
mlr_model2 <- function() {</pre>
    for (i in 1:N) {
        theta[i, 1:p] ~ dmnorm(x[i, 1:p],
            precision2[, ])
    }
    # Prior for likelihood parameters: mu2,
    # precision2, rho
    rho \sim dunif(-1, 1)
    for (j in 1:p) {
        mu2[j] ~ dnorm(0, 0.01)
    precision2[1:p, 1:p] ~ dwish(R[, ],
        k)
    # Missing data model for x
    for (i in 1:N) {
        x[i, 1:p] ~ dmnorm(x_mn[], x_prec[,
            ])
    }
    # Priors for missing-data model
```

```
# parameters
    for (j in 1:p) {
        x_mn[j] ~ dnorm(0, 0.01)
    x_prec[1:p, 1:p] ~ dwish(R[, ], k)
    x_cov[1:p, 1:p] <- inverse(x_prec[,</pre>
        ])
    k < -p + 0.1
    for (j1 in 1:p) {
        for (j2 in 1:p) {
             R[j1, j2] \leftarrow 0.1 * equals(j1,
                 j2)
        }
    }
}
n.chains = 1
nSamples = 10000
stacks_dat \leftarrow list(x = x, p = 6, N = 365)
mlr_inits <- function() {</pre>
    list(rho = 0)
}
if (FALSE) {
    samps <- bugs(data = stacks_dat, inits = mlr_inits,</pre>
        parameters.to.save = c("theta"),
        model.file = mlr_model2, codaPkg = TRUE,
        n.chains = n.chains, n.burnin = 2000,
        n.iter = nSamples, n.thin = 10,
        DIC = F)
    out.coda <- read.bugs(samps)</pre>
    save(out.coda, file = "out.coda_BUGS_MVN.RData")
    load("out.coda_BUGS_MVN.RData")
}
if (n.chains > 1) {
    gelman.srf <- gelman.diag(out.coda)</pre>
    count.coeff.gt <- sum(gelman.srf$psrf >
         1.1)
    count.coeff.gt
}
chains.ess <- lapply(out.coda, effectiveSize)</pre>
```

```
first.chain.ess <- chains.ess[1]
plot(unlist(first.chain.ess), main = "Effective Sample Size")</pre>
```

Effective Sample Size



```
chain <- out.coda[[1]]
posterior.means <- list()

posterior.modes <- list()

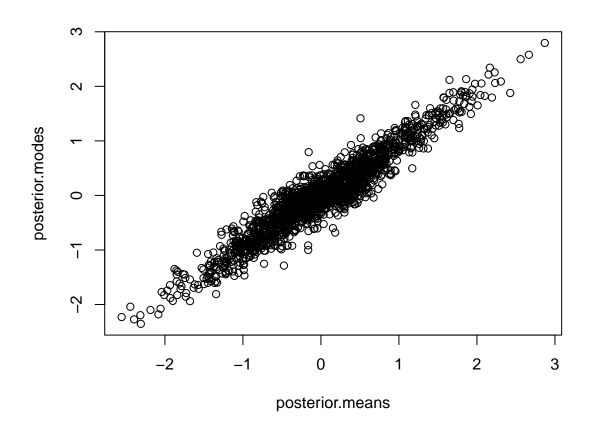
for (i in 1:(365 * 6)) {
    colname <- colnames(chain)[i]

    samples <- chain[, i]

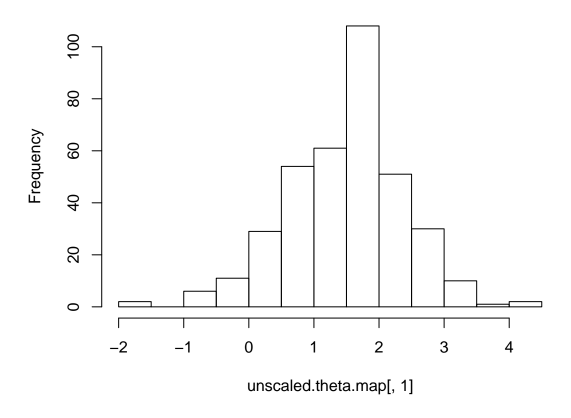
    posterior.means[i] <- mean(samples)

    posterior.modes[i] <- mlv(samples)$M
}

plot(posterior.means, posterior.modes)</pre>
```



Histogram of unscaled.theta.map[, 1]



hist(Y1[, 1])

Histogram of Y1[, 1]

