Bayesian model in JAGS

Tutorial 3 for transition

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```
knitr::opts_chunk$set(echo = TRUE)
require(rjags)
#"Note that the rjags package does not include a copy of
# the JAGS you must install this separately.
# For instructions on downloading JAGS,
# see the home page at https://mcmc-jags.sourceforge.io."
```

require(mcmcplots) # to plot the trace plots

require(coda) # for gelman diagnostic

Outline

In this tutorial, we will learn how to use JAGS (Just Another Gibbs Sampler) in R to estimate unknown parameters with Markov Chain Monte Carlo (MCMC) methods in Bayesian framework. If you are interested in learning more about the theoretical knowledge of MCMC and Bayesian modeling, please check out courses "Probability and Statistics" (taught by Prof. Joseph Chang) and "Bayesian Statistics" (taught by Prof. Josh Warren).

In the first section, we will introduce a simple jags model to help understand the syntax for rjags package.

In the second section, we will learn how to build up complicated models step by step (censor data). Before reading this section, you will need to familiar yourself with the RSV_timing_explain tutorials and a research article on the relative RSV timing

In the third section, we will introduce the concept of spatial autocorrelation and how to incorporate this in jags model.

Example 1: a simple jags model to estimate the unknown parameters

First: Simulate a dataset for analysis

In this example, I simulate my own data in R in order to compare the MCMC results with our input parameters. I create a continuous outcome variable y as a function of one predictor x and an error term ϵ . I simulate 100 observations. The linear regression looks like

$$y = \beta_0 + \beta_1 x + \epsilon$$

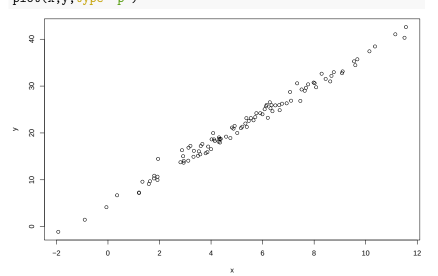
Our goal is to estimate the unknown parameters β_0 and β_1 given the observations and the linear relationship

```
n.sim=100; set.seed(123)
x=rnorm(n.sim, mean = 5, sd = 3)
epsilon=rnorm(n.sim, mean = 0, sd = 1)
beta0=5
beta1=3.2
y=beta0 + beta1 * x + epsilon
```

Visualize the observations

Question: what is the intercept and slope for this linear regression?

plot(x,y,type="p")



rjags mamual

The following is quoted from rjags document, authored by Martyn Plummer:

"JAGS is a clone of BUGS (Bayesian analysis Using Gibbs Sampling). See Lunn et al (2009) for a history of the BUGS project. Note that the rjags package does not include a copy of the JAGS library: you must install this separately. For instructions on downloading JAGS, see the home page at https://mcmc-jags.sourceforge.io.

To fully understand how JAGS works, you need to read the JAGS User Manual. The manual explains the basics of modelling with JAGS and shows the functions and distributions available in the dialect of the BUGS language used by JAGS. It also describes the command line interface. The rjags package does not use the command line interface but provides equivalent functionality using R functions."

rjags mamual (continued)

- "Analysis using the rjags package proceeds in steps: 1. Define the model using the BUGS language in a separate file.
- 2. Read in the model file using the jags.model function. This creates an object of class "jags".
- 3. Update the model using the update method for "jags" objects. This constitutes a 'burn-in' period.
- 4. Extract samples from the model object using the coda.samples function. This creates an object of class "mcmc.list" which can be used to summarize the posterior distribution. The coda package also provides convergence diagnostics to check that the output is valid for analysis (see Plummer et al 2006)."

Jags model structure (Step 1)

```
basic_mod <- "model{</pre>
 #model
  for(i in 1:n.sim){
   y[i] ~ dnorm(mu[i], tau)
# tau is the error term
   mu[i] = beta0 + beta1 * x[i]
# beta0 and beta1 are the unknown parameters
  }
 #priors; these are weakly informative priors
 beta0 \sim dnorm(0, 0.01)
# dnorm(mean, precision)
# precision = 0.01 --> var = 100
 beta1 ~ dnorm(0, 0.01)
tau ~ dgamma(0.01,0.01)
# dgamma(shape,rate)
}
```

Initialize model with known data and initial guess of unknown parameters (Step 2)

```
######## Define known data #########
datalist=list("y"=y,"x"=x,"n.sim"=n.sim)

######## Set initial values for parameter estimate ######
inits=function(){list("beta0"=rnorm(1), "beta1"=rnorm(1))}
```

Read in the model file using the jags.model function (Step 3)

```
## Compiling model graph
## Resolving undeclared variables
## Allocating nodes
## Graph information:
## Observed stochastic nodes: 100
## Unobserved stochastic nodes: 3
## Total graph size: 406
##
## Initializing model
```

Discard burn-in period and sample for posterior distribution (Step 4)

Summarize the posterior distribution

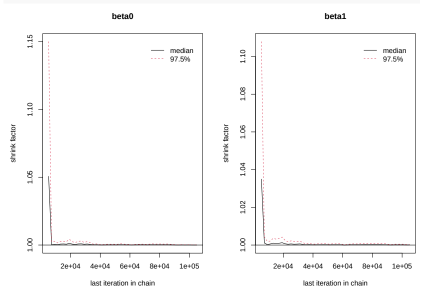
```
summary(model_sample)
```

```
##
## Iterations = 5005:105000
## Thinning interval = 5
  Number of chains = 4
  Sample size per chain = 20000
##
   1. Empirical mean and standard deviation for each variable,
     plus standard error of the mean:
##
##
##
         Mean
                   SD Naive SE Time-series SE
## beta0 4.981 0.21408 0.0007569 0.0010421
## beta1 3.183 0.03594 0.0001271 0.0001752
##
  2. Quantiles for each variable:
##
         2.5% 25% 50% 75% 97.5%
##
## beta0 4.558 4.838 4.981 5.125 5.399
## beta1 3.113 3.159 3.183 3.207 3.254
```

Check for model convergence (gelman diagnotics)

Check for model convergence (gelman plot)

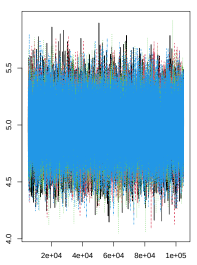
gelman.plot(model_sample)



Plot the trace plot

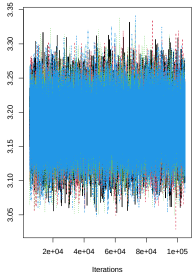
```
par(mfrow=c(1,2))
traceplot(model_sample)
```





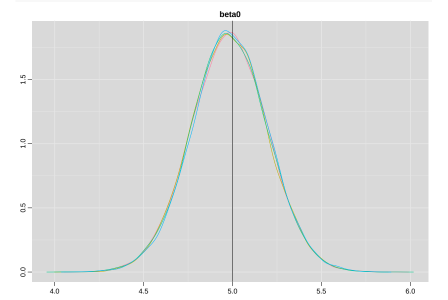
Iterations

Trace of beta1



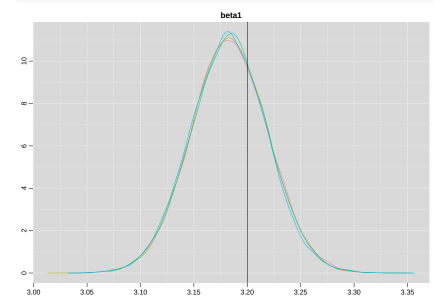
Compare the posterior density and real value for beta0

```
denplot(model_sample, parms = c("beta0"))
abline(v=5) # True value
```



Compare the posterior density and real value for beta1

```
denplot(model_sample, parms = c("beta1"))
abline(v=3.2) # True value
```



Section 2: estimate the parameters for the relative RSV timing

After we estimated the RSV timing using second derivative method (please check: RSV_timing_explain), we would like to know if some potential drivers contribute to the relative timing difference in RSV epidemic onsets before and during COVID-19 pandemic. (In reality, we manually coded the spatial autocorrelated model. As this requires taking Josh's classes, I will not go over it in the introduction tutorials.)

The model is given as:

$$Y_i = \alpha_{0i} + \beta_1 \overline{W_i} + \epsilon_i$$
$$\alpha_{0i} = \beta_0 + \mathbf{x}_i^T \boldsymbol{\lambda} + \theta_i$$

Meaning of the variables and parameters

where Y_i is the relative timing of RSV epidemic onset with respect to Florida in 2021 in state i (measured in weeks), and $\overline{w_i}$ is the average relative timing of RSV seasons in state i compared to Florida from 2016-2019. The global intercept parameter β_0 and slope parameter β_1 describe the overall similarity between a typical year's relative onset timing across all states and the relative timings observed in 2021. The observation-level random effect ϵ_i $N(0,\sigma^2)$ accounts for residual variability in the data.

The state-specific intercepts, $\alpha_{0i}=\beta_0+\mathbf{x_i}^T\boldsymbol{\lambda}+\theta_i$, represent the state-level similarity in relative timing for 2021 compared to previous seasons, where x_i is the vector of covariates that could potentially explain differences in the relative timing of RSV between 2021 and previous years. The covariates include population density, average household size, and stringency index of non-pharmaceutical interventions against COVID-19.

 θ_i represents a spatially correlated random effect, which we will learn more in detail later.

Jags model code without considering spatial autocorrelation

```
Nospatial model <- "model{
for(i in 1:n){
Y[i] ~ dnorm(mu[i], inv var)
# for the observed timing (we log transform Y already)
mu[i] <- alpha 0[i] + beta1*log W[i]</pre>
# log W is the log transform average relative timing of RSV
alpha 0[i]= beta0+x1[i]*lambda1+x2[i]*lambda2+lambda3*x3[i]
# this is alpha_0i = beta0 + X_i*lambda (coviariates)
beta0~dnorm(0,0.0001)
beta1~dnorm(0,0.0001)
lambda1~dnorm(0,0.0001)
lambda2~dnorm(0,0.0001)
lambda3~dnorm(0,0.0001)
```

```
Load the data and run the jags model to sample posteriors
   ## loading data from the working directory
   data_nospatial <- readRDS("./Nospatial_data.rds")</pre>
   ### Read in the model file using the jags.model function ##
   non spatial <- jags.model(textConnection(Nospatial model),</pre>
                         data=data nospatial,n.chains = 4)
   ## Warning in jags.model(textConnection(Nospatial_model), data =
   ## Unused variable "Z" in data
   ## Compiling model graph
   ##
         Resolving undeclared variables
         Allocating nodes
   ##
   ## Graph information:
         Observed stochastic nodes: 0
   ##
   ##
         Unobserved stochastic nodes: 53
   ##
         Total graph size: 491
```

##

Initializing model

update(non spatial, 10000)

######### burn-in period ##########

Summarize the posterior distribution

summary(non_spatial_sample)

##

```
##
## Tterations = 10020:110000
## Thinning interval = 20
## Number of chains = 4
  Sample size per chain = 5000
##
  1. Empirical mean and standard deviation for each variable,
     plus standard error of the mean:
##
##
##
             Mean
                       SD Naive SE Time-series SE
                                         0.71204
## beta0 -0.85052 101.086 0.71478
                                         0.71214
## beta1 -0.07134 100.709 0.71212
## inv_var 0.94198 9.622 0.06804
                                         0.06979
## lambda1 -0.48336 99.914 0.70650
                                         0.70685
## lambda2 0.27092 99.471 0.70336
                                     0.70785
  lambda3 1.50128 99.755 0.70538 0.70540
##
  2. Quantiles for each variable:
```

Check for model convergence (gelman diagnostic)

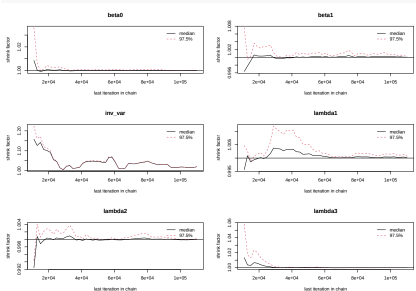
inv_var did not converge well. This indicates that we should either prolong the iterations or re-parameterize the model. Using gelman.plot (next slides), we could tell that prolong iterations may help convergence.

```
gelman.diag(non_spatial_sample)
```

```
## Potential scale reduction factors:
##
##
           Point est. Upper C.I.
##
  beta0
                 1.00
                            1.00
                 1.00
                            1.00
## beta1
  inv var
                1.02
                            1.02
  lambda1
                1.00
                            1.00
##
## lambda2
                1.00
                            1.00
## lambda3
                 1.00
                            1.00
##
## Multivariate psrf
##
## 1
```

Gelman plot

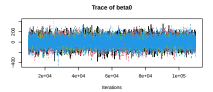
par(mfrow=c(3,2))
gelman.plot(non_spatial_sample)

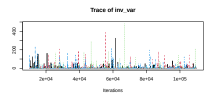


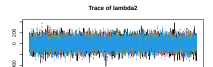
Trace plots

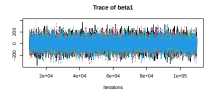
note: inv_var is precision = the inverse of variance

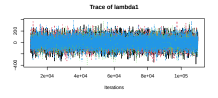
par(mfrow=c(3,2))
traceplot(non_spatial_sample)

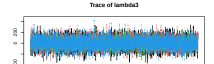






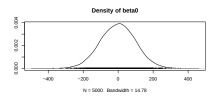


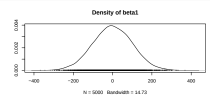


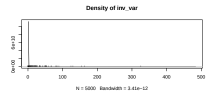


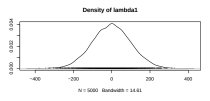
Density plot of the posterior distributions

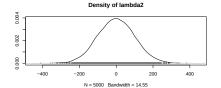
par(mfrow=c(3,2))
densplot(non_spatial_sample)

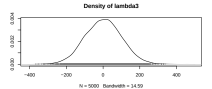












Section 2.1: right censored survival analysis

In the early phase of an epidemic, we may not yet observe the onset point. For example, if you consider an onset threshold and the cases are still below the threshold, the epidemic has not taken off. In this case, we need to use right censored survival analysis.

Click to know more about right censored survival analysis

Model structure for right censored survival analysis

```
censormodel <- "model{</pre>
for(i in 1:n){
is.censor[i] ~ dinterval(Z[i],t.cen)
# whether the data is censor and the censoring time
Z[i] ~ dlnorm(mu[i], inv var)
# for the observed timing (we did not log transform Z)
# if we already log transform Z then we use dnorm
mu[i] <- alpha_0[i] + beta1*log_W[i]</pre>
# log_W is the log transform average relative timing of RSV
alpha_0[i] = beta0+x1[i] *lambda1+x2[i] *lambda2
# this is alpha_0i = beta0 + X_i*lambda (coviariates)
```

Model structure for right censored survival analysis (continued)

Since the model running process is the same, I will skip the model fitting process in this tutorial (it takes time to knit the PDF). To run the model, you need to combine the two chunks of model structures and change the eval from False to True.

Same process to sample posteriors of right censored model

```
## loading data from the working directory
data censor <- readRDS("./censor data.rds")</pre>
### Read in the model file using the jags.model function ##
censor_model_jags <- jags.model(textConnection(censormodel),</pre>
                      data=data censor, n.chains = 4)
######### burn-in period ##########
update(censor model jags, 10000)
####### sample for posterior distribution #########
censor model sample <- coda.samples(censor_model_jags,</pre>
                             c("beta0", "beta1",
                             "lambda1", "lambda2",
                             "lambda3", "inv var"),
                              100000, thin=20)
```

Section 3: spatial autocorrelation

Very often, the disease incidence is not randomly distributed across space. Observations in nearby localities tend to be more similar than observations further away, also known as positive spatial autocorrelation.

Prof. Duncan Lee has developed many statistical tools and tutorials to help evaluate spatial autocorrelation in public health. Please refer to the following materials to have an complete overview on Bayesian spatial model for public health and choosing conditional autoregressive priors:

- ▶ Bayesian Disease Mapping for Public Health
- Overview of Spatial Incidence Models
- CARBayes: An R Package for Bayesian Spatial Modeling with Conditional Autoregressive Priors
- ➤ Spatio-Temporal Areal Unit Modeling in R with Conditional Autoregressive Priors Using the CARBayesST Package

A generalized linear mixed effect model (GLMM) framework

To produce accurate estimates of disease incidence, statistical models must take into account the full range of features present in the data, including non-normal distributions of count data, the impact of explanatory variables, and spatial correlation. One solution is to utilize generalized linear mixed models (GLMMs), which integrate a generalized linear model (GLM) with normally distributed random effects that account for both spatial correlation and overdispersion.

An example model structure is given as:

$$Y_i \sim Poisson(\mu_i)$$

$$\log \mu_i = \log E_i + x_i' \alpha + \phi_i$$

where Y_i is the observed number of disease in region i, μ_i is the conditional mean, E_i is the expected number of disease in region i, x_i is the explanatory variables (covariates) of region i, α is the coefficients of the corresponding explanatory variables, and ϕ_i accounts for the random effects, which can be spatially correlated.

Conditional autoregressive (CAR) priors for spatial modeling

To model spatial autocorrelation, four CAR priors are commonly used, including the intrinsic models, BYM models, Leroux models, and Cressie models. In **Duncan Lee's paper**, he systematically compared these four different priors and concluded that Leroux model is the best from both theoretical and practical standpoints. For short, Leroux model is appealing because it uses a single set of random effects to model the varying strengths of spatial autocorrelation $\rho,$ where $\rho=0$ means spatially independent while ρ near 1 indicates strong spatial correlation.

Leroux priors for spatial modeling

The Leroux version of spatial random effects is given as:

$$\phi_i | \phi_{-i}, \rho, \tau^2 \sim MVN(\frac{\rho \sum_{j=1}^n w_{ij} \phi_i}{\rho \sum_{j=1}^n w_{ij} + 1 - \rho}, \frac{\tau^2}{\rho \sum_{j=1}^n w_{ij} + 1 - \rho})$$

where $\rho \in [0,1)$ represents the spatial correlation that was estimated from the data; $\rho=0$ corresponds to spatial independence, simplifying the model to an independent random effects model, and ρ near 1 indicates strong spatial correlation. The w_{ij} are variables that contain information about spatial proximity between two locations; w_{ij} equals one if regions i and j share a common border and equal zero otherwise (w_{ii} =0 for all i). The total variance of spatial random effects is described by τ^2 .

```
Create a spatial adjacency matrix
Step 1. Set up – Shapefiles
(I learned this from Kayoko Shioda!)
```

Load a shapefile. Make sure each polygon on shapefile
corresponds to only one state.
require(rgdal) # for working with shapefiles

require(rgdal) # for working with shapefiles
require(ggplot2) # for plot and fortify
CT1map = readOGR(dsn=path.expand('./WGS84'), layer= 'zipct_37800')

OGR data source with driver: ESRI Shapefile
Source: "/Users/zhezheng/Library/CloudStorage/Box-Box/RSV_tut
with 282 features
It has 11 fields

Integer64 fields read as strings: ALAND10 AWATER10
CTmap.df <- as.data.frame(CT1map)
CT1map_fort <- fortify(CT1map,region="ZCTA5CE10")
ZCTA5CE10 is CT zipcode ID
The ID variable is then the zipcode/area unit</pre>

Sort both the shapefile and the data file by ID.

CT1map_fort\$zipcode <- CT1map_fort\$id

How to create a spatial adjacency matrix Step 2. Create a neighbor matrix from the sorted shapefile

```
library(spdep) # for neighbor evaluation
# Neighbors can either be Queen
# (any zip that touches another zip - even at corners)
# or Rook neighbors (share full sides -- not corners)
# The snap dist is governing the NA in neighbor matrix
# Can remove SNAP statement:
# if Queen = F then single point touch not allowed
# and Rook distance used.
# When data is missing -- there's more non-zero links involved
# Try to make snap distance as small as possible
# while making sure all zipcode have at least one neighbor
neighb <- poly2nb(CTmap.sort, queen = T, snap = sqrt(0.001))</pre>
# Check average number of links -- increase snap distance
```

Check average number of links -- increase snap distance # if it says there is a zip with zero neighbors neighb # Avg Number of links = 8.18705

How to create a spatial adjacency matrix

Step 3. Create a W matrix for Leroux model

```
# For Leroux model, neighbor matrix needs to have O
# if not neighboring, -1 if neighbor,
# and number of neighbors for each zip on diagonal), so...
w.mat <- ifelse(neighb.mat==1,-1,neighb.mat)
num.neigh <- colSums(neighb.mat)</pre>
# Count the number of neighbors for each state
for (i in 1:nrow(w.mat)) {
  # Put the number of neighbors on diagonal
  for (j in 1:ncol(w.mat)) {
    if (i==j) {
      w.mat[i,j] <- num.neigh[i]</pre>
```

Spatial model with censored data

```
censor spatial model <- "model{</pre>
for(i in 1:n){
is.censor[i] ~ dinterval(Z[i],t.cen)
Z[i] ~ dlnorm(mu[i], inv_var)
mu[i] <- beta0[i] + beta1*log W[i]</pre>
beta0[1:n] ~ dmnorm(intercepts, Leroux)
for(i in 1:n){
 intercepts[i] <- b0+x1[i]*b1+x2[i]*b2}
beta1~dnorm(0,0.0001)
b0~dnorm(0,0.0001)
b1~dnorm(0,0.0001)
b2~dnorm(0,0.0001)
```

Spatial model with censored data (continued)

```
"model{# delete this one line of code
# when combine the two chunks. Separating the
# codes helps to show them in slides
Leroux <- inv.tau*(rho*w.mat + (1 - rho)*Id.mat)</pre>
# calculate spatial correlated random effects
  inv.tau ~ dgamma(0.01, 0.01)
# note the w.mat here is equal to
# diag(rowSums(neighbors_mat)) - neighbors_mat
  rho~ dunif(0,1)
inv_var ~ dgamma(0.01, 0.01)}"
```