Hands-on training session 1

Introduction, revision and running basic models

Matt Denwood Giles Innocent 2020-02-19

Course Outline and Practicalities

Overview

Date/time:

- 19th February 2020
- **1**4.00 15.30

Teachers:

- Matt Denwood (presenter)
- Giles Innocent

Revision

Bayes Rule

Bayes' theorem is at the heart of Bayesian statistics:

$$P(\theta|Y) = \frac{P(\theta) \times P(Y|\theta)}{P(Y)}$$

3

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$$P(\theta|Y) = \frac{P(\theta) \times P(Y|\theta)}{P(Y)}$$

Where: θ is our parameter value(s);

Y is the data that we have observed;

 $P(\theta|Y)$ is the posterior probability of the parameter value(s);

 $P(\theta)$ is the prior probability of the parameters;

 $P(Y|\theta)$ is the likelihood of the data given the parameters value(s);

P(Y) is the probability of the data, integrated over parameter space.

• In practice we usually work with the following:

$$P(\theta|Y) \propto P(\theta) \times P(Y|\theta)$$

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$$P(\theta|Y) \propto P(\theta) \times P(Y|\theta)$$

- Our Bayesian posterior is therefore always a combination of the likelihood of the data, and the parameter priors
- But for more complex models the distinction between what is 'data' and 'parameters' can get blurred!

MCMC

- A way of obtaining a numerical approximation of the posterior
- Highly flexible
- Not inherently Bayesian but most widely used in this context
- Assessing convergence is essential, otherwise we may not be summarising the true posterior
- Our chains are correlated so we need to consider the effective sample size

Preparation

- Any questions so far? Anything unclear?
- Do we all have R and JAGS installed?
- Can we all access the teaching material from GitHub?

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Any problems: see us during the first practical session!

of MCMC

Session 1a: Theory and application

MCMC in Practice

- We can write a Metropolis algorithm but this is complex and inefficient
- There are a number of general purpose languages that allow us to define the problem and leave the details to the software:
 - WinBUGS/OpenBUGS
 - Bayesian inference Using Gibbs Sampling
 - JAGS
 - Just another Gibbs Sampler
 - Stan
 - Named in honour of Stanislaw Ulam, pioneer of the Monte Carlo method

JAGS

- JAGS uses the BUGS language
 - This is a declarative (non-procedural) language
 - The order of statements does not matter
 - The compiler converts our model syntax into an MCMC algorithm with appropriately defined likelihood and prior
 - You can only define each variable once!!!

JAGS

- JAGS uses the BUGS language
 - This is a declarative (non-procedural) language
 - The order of statements does not matter
 - The compiler converts our model syntax into an MCMC algorithm with appropriately defined likelihood and prior
 - You can only define each variable once!!!
- Different ways to run JAGS from R:
 - rjags, runjags, R2jags, jagsUI
- See http://runjags.sourceforge.net/quickjags.html
 - This is also in the GitHub folder

A simple JAGS model might look like this:

```
model{
      # Likelihood part:
      Positives ~ dbinom(prevalence, TotalTests)
3
5
      # Prior part:
      prevalence ~ dbeta(2, 2)
6
      # Hooks for automatic integration with R:
8
      #data# Positives, TotalTests
      #monitor# prevalence
10
      #inits# prevalence
11
12
```

There are two model statements:

Positives ~ dbinom(prevalence, TotalTests)

 states that the number of Positive test samples is Binomially distributed with probability parameter prevalence and total trials TotalTests There are two model statements:

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prevalence ~ dbeta(2,2)
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 states that our prior probability distribution for the parameter prevalence is Beta(2,2)

These are very similar to the likelihood and prior functions defined in the preparatory exercise

The other lines in this model:

```
#data# Positives, TotalTests
#monitor# prevalence
#inits# prevalence
```

are automated hooks that are only used by runjags

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- #data# Positives, TotalTests
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Compared to our Metropolis algorithm, this JAGS model is:

- Eaiser to write and understand
- More efficient (lower autocorrelation)
- Faster to run

To run this model, copy/paste the code above into a new text file called "basicjags.bug" in the same folder as your current working directory. Then run:

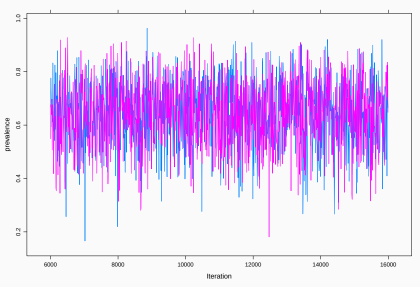
```
results <- run.jags('basicjags.bug', n.chains=2, burnin=5000,

⇒ sample=10000)
```

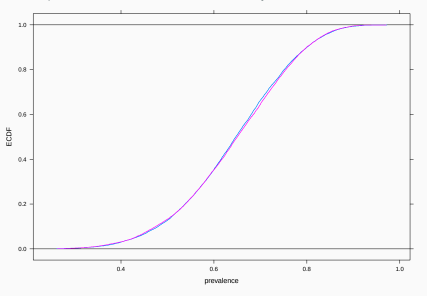
First check the plots for convergence:

```
plot(results)
```

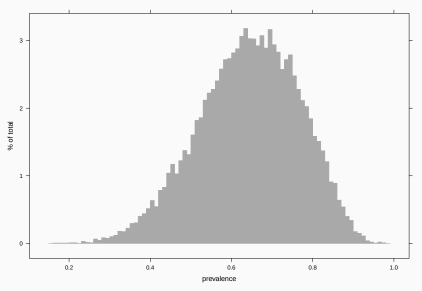
Trace plots: the two chains should be stationary:



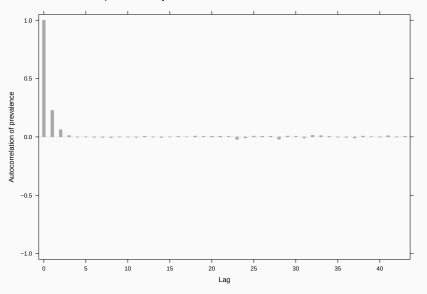
ECDF plots: the two chains should be very close to each other:



Histogram of the combined chains should appear smooth:



Autocorrelation plot tells you how well behaved the model is:



Then check the effective sample size (SSeff) and Gelman-Rubin statistic (psrf):

```
results
   ##
   ## JAGS model summary statistics from 20000 samples (chains = 2;
       adapt+burnin = 6000):
   ##
                Lower95 Median Upper95 Mean
   ##
                                                    SD
                                                          Mode
   ## prevalence 0.40498 0.64856 0.87278 0.64301 0.12317 0.66466
   ##
                    MCerr MC%ofSD SSeff AC.10 psrf
   ##
   ## prevalence 0.0011071 0.9 12377 -0.00013892 1
9
   ##
10
   ## Total time taken: 0.5 seconds
```

Reminder: we want psrf < 1.05 and SSeff > 1000

Exercise

- Run this model yourself in JAGS
- Change the initial values for the two chains and make sure it doesn't affect the results
- Reduce the burnin length does this make a difference?
- Change the sample length does this make a difference?

Optional Exercise

- Change the number of chains to 1 and 4
 - Remember that you will also need to change the initial values
 - What affect does having different numbers of chains have?
- Try using the run.jags argument method='parallel' what affect does this have?

Session 1b: Working with basic

models (apparent prevalence)

Other runjags options

There are a large number of other options to runjags. Some highlights:

- The method can be parallel or background or bgparallel
- You can use extend.jags to continue running an existing model (e.g. to increase the sample size)
- You can use coda::as.mcmc.list to extract the underlying MCMC chains
- Use the summary() method to extract summary statistics
 - See ?summary.runjags and ?runjagsclass for more information

Using embedded character strings

 For simple models we might not want to bother with an external text file. Then we can do:

```
mt <- "
    model{
      Positives ~ dbinom(prevalence, TotalTests)
      prevalence ~ dbeta(2, 2)
5
      #data# Positives, TotalTests
6
      #monitor# prevalence
      #inits# prevalence
10
1.1
    results <- run.jags(mt, n.chains=2)
12
```

But I would advise that you stick to using a separate text file!

Setting the RNG seed

• If we want to get numerically replicable results we need to add .RNG.name and .RNG.seed to the initial values, and an additional #modules# lecuyer hook to our basicjags.bug file:

```
model{
Positives ~ dbinom(prevalence, TotalTests)
prevalence ~ dbeta(2, 2)

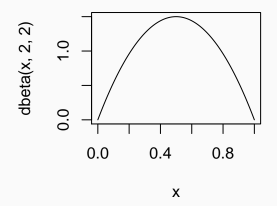
#data# Positives, TotalTests
#monitor# prevalence
#inits# prevalence, .RNG.name, .RNG.seed
#modules# lecuyer
}
```

```
1   .RNG.name <- "lecuyer::RngStream"
2   .RNG.seed <- list(chain1=1, chain2=2)
3   results <- run.jags('basicjags.bug', n.chains=2)</pre>
```

A different prior

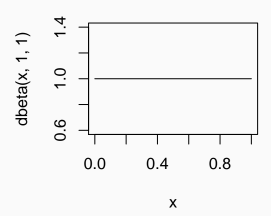
• A quick way to see the distribution of a prior:

```
curve(dbeta(x, 2, 2), from=0, to=1)
```



• A minimally informative prior might be:

curve(dbeta(x, 1, 1), from=0, to=1)



• Let's change the prior we are using to dbeta(1,1):

```
model{
model{
positives ~ dbinom(prevalence, TotalTests)

prevalence ~ dbeta(1, 1)

# Hooks for automatic integration with R:
# data# Positives, TotalTests
#monitor# prevalence
#inits# prevalence

}
```

An Equivalent Model

We could equivalently specify an observation-level model:

```
model{
      # Likelihood part:
      for(i in 1:TotalTests){
        Status[i] ~ dbern(prevalence)
 5
6
      # Prior part:
      prevalence ~ dbeta(1, 1)
8
10
      # Hooks for automatic integration with R:
      #data# Status, TotalTests
11
      #monitor# prevalence
12
      #inits# prevalence
1.3
14
```

• But we need the data in a different format: a vector of 0/1 rather than total positives!

A GLM Model

```
model{
      # Likelihood part:
      for(i in 1:TotalTests){
        Status[i] ~ dbern(predicted[i])
        logit(predicted[i]) <- intercept</pre>
5
6
7
      # Prior part:
8
      intercept ~ dnorm(0, 10^-6)
9
10
      # Derived parameter:
1.1
      prevalence <- ilogit(intercept)</pre>
12
13
      # Hooks for automatic integration with R:
14
      #data# Status, TotalTests
15
      #monitor# intercept, prevalence
16
      #inits# intercept
17
18
```

- This is the start of a generalised linear model, where we could add covariates at individual animal level.
- We introduce a new distribution dnorm() notice this is mean and precision, not mean and sd!
- For a complete list of the distributions available see:
 - https://sourceforge.net/projects/mcmcjags/files/Manuals/4.x/
 - This document is also provided on the GitHub repository
- However, notice that the prior is specified differently...

Exercise

- Run the original version, the observation-level version, and the GLM version of the model and compare results with the same data
- Now try a larger sample size: e.g. 70 positives out of 100 tests
 are the posteriors from the two models more or less similar than before?
- Now try running the GLM model with a prior of dnorm(0, 0.33) (and the original data) - does this make a difference?

Optional Exercise

Another way of comparing different priors is to run different models with no data - as there is no influence of a likelihood, the posterior will then be identical to the priors (and the model will run faster).

One way to do this is to make all of the response data (i.e. either Positives or Status) missing. Try doing this for the following three models, and compare the priors for prevalence:

- The original model with prior prevalence ~ dbeta(1,1)
- The GLM model with prior intercept ~ dnorm(0, 10^-6)
- The GLM model with prior intercept ~ dnorm(0, 0.33)

models (imperfect test)

Session 1c: Basics of latent-class

Imperfect tests

- Up to now we have ignored issues of diagnostic test sensitivity and specificity
- Usually, however, we do not have a perfect test, so we do not know how many are truly positive or truly negative, rather than just testing positive or negative.
- But we know that:

$$Prev_{obs} = (Prev_{true} \times Se) + ((1 - Prev_{true}) \times (1 - Sp))$$

$$\implies Prev_{true} = \frac{Prev_{obs} - (1 - Sp)}{Se - (1 - Sp)}$$

Model Specification

 We can incorporate the imperfect sensitivity and specicifity into our model:

```
model{
1
      Positives ~ dbinom(obsprev, TotalTests)
      obsprev <- (prevalence * se) + ((1-prevalence) * (1-sp))
      prevalence ~ dbeta(1, 1)
5
      se ~ dbeta(1, 1)
      sp ~ dbeta(1, 1)
9
      #data# Positives, TotalTests
      #monitor# prevalence, obsprev, se, sp
10
      #inits# prevalence, se, sp
11
12
```

And run it:

```
prevalence <- list(chain1=0.05, chain2=0.95)
se <- list(chain1=0.5, chain2=0.99)
sp <- list(chain1=0.5, chain2=0.99)
Positives <- 70
TotalTests <- 100
results <- run.jags('basicimperfect.bug', n.chains=2)</pre>
```

Finished running the simulation

[Remember to check convergence and effective sample size!]

What do these results tell us?

	Lower95	Median	Upper95	SSeff	psrf
prevalence	0.058	0.515	0.996	1918	1.002
obsprev	0.604	0.694	0.778	20215	1.000
se	0.152	0.702	1.000	1527	1.002
sp	0.000	0.319	0.861	1590	1.002

What do these results tell us?

Lower95	Median	Upper95	SSeff	psrf
0.058	0.515	0.996	1918	1.002
0.604	0.694	0.778	20215	1.000
0.152	0.702	1.000	1527	1.002
0.000	0.319	0.861	1590	1.002
	0.058 0.604 0.152	0.058 0.515 0.604 0.694 0.152 0.702	0.058 0.515 0.996 0.604 0.694 0.778 0.152 0.702 1.000	0.058 0.515 0.996 1918 0.604 0.694 0.778 20215 0.152 0.702 1.000 1527

- We can estimate the observed prevalence quite well
- But not the prevalence, se or sp!
 - The model is unidentifiable.

Priors

- We cannot estimate se, sp and prevalence simultaneously
 - We need strong priors for se and sp
- We can use the PriorGen package to generate Beta priors based on published results, for example:

```
PriorGen::findbeta(themean=0.9, percentile = 0.975,

percentile.value = 0.8)

## [1] "The desired Beta distribution that satisfies the

specified conditions is: Beta( 41.82 4.65 )"

## [1] "Here is a plot of the specified distribution."

## [1] "Descriptive statistics for this distribution are:"

## Min. 1st Qu. Median Mean 3rd Qu. Max.

## 0.6425 0.8748 0.9060 0.9001 0.9326 0.9911

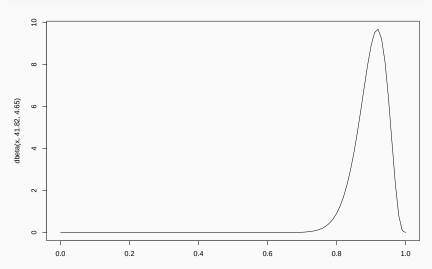
## [1] "Verification: The percentile value 0.8 corresponds to the

0.025 th percentile"
```

```
qbeta(c(0.025, 0.5, 0.975), 41.82, 4.65)
```

1 **##** [1] 0.7999144 0.9056630 0.9677283

curve(dbeta(x, 41.82, 4.65), from=0, to=1)



Exercise

- Find beta distribution priors for:
 - Sensitivity = 0.9 (95% CI: 0.85 0.95)
 - Specificity = 0.95 (95%CI: 0.92-0.97)
- Look at these distributions using curve and qbeta
- Modify the imperfect test model using these priors and re-estimate prevalence

Optional Exercise

- Run the same model with se and sp fixed to the mean estimate
 - How does this affect CI for prevalence?
- Run the same model with se and sp fixed to 1
 - How does this affect estimates and CI for prevalence?

Summary

- Using JAGS / runjags allows us to work with MCMC more easily, safely and efficiently than writing our own sampling algorithms
- But we must never forget to check convergence and effective sample size!
- More complex models become easy to implement
 - For example imperfect diagnostic tests
- But just because a model can be defined does not mean that it will be useful for our data
 - We need to be realistic about the information available in the data, what parameters are feasible to estimate, and where we will need to use strong priors