Introductory Bayesian Course

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1 Background

The purpose of these course notes is to introduce participants to Bayesian analysis with R, RStudio and JAGS. It is assumed that participants are familiar with R and RStudio as covered in the Introductory R Course notes at http://www.poissonconsulting.ca/course/2014/09/12/an-introduction-to-r-course.html.

1.1 Licence

The notes, which are released under a CC BY 4.0 license, are a draft of the material to be presented at the Introductory Bayesian Course in Kelowna on November 20th-21st, 2014. They were written by Dr. Joseph Thorley R.P.Bio..

1.2 Installation

If you haven't already done so, download the the most recent version of the R base distribution binary for your platform from http://cran.r-project.org/ and install using the default options. Next download and install RStudio from http://www.rstudio.com/products/rstudio/download/ using the default options. Then, download JAGS from http://sourceforge.net/projects/mcmc-jags/files/JAGS/ and install with the default options.

To make sure you have all the required packages installed on your hard drive execute the following code at the command line

```
install.packages("devtools", quiet = TRUE)
library(devtools)
install.packages("dplyr", quiet = TRUE)
install.packages("ggplot2", quiet = TRUE)
```

```
install.packages("scales", quiet = TRUE)
install_github("poissonconsulting/tulip@v0.0.11")
install_github("poissonconsulting/datalist@v0.4")
install_github("poissonconsulting/juggler@v0.1.3")
install_github("poissonconsulting/jaggernaut@v2.0.0")
```

Then start any scripts with

```
library(dplyr)
library(ggplot2)
library(scales)
library(jaggernaut)
```

1.3 Bayesian and Frequentist Statistical Analysis

Statistical analysis uses probability models to provide bounded estimates of parameter values (θ) from the data (y).

There are two primary approaches to statistical analysis: Bayesian and frequentist. As far as a frequentist is concerned the best estimates of θ are those values that maximise the *likelihood* which is the probability of the data given the estimates, i.e., $p(y|\theta)$. A Bayesian on the other hand chooses the values with the highest posterior probability - that is to say the probability of the estimates given the data, i.e., $p(\theta|y)$.

1.3.1 Coin Flips

Consider the case where n = 10 flips of a coin produce y = 3 tails. We can model this using a binomial distribution

$$y \sim dbin(\theta, n)$$

where θ is the probability of throwing a head.

1.3.1.1 Maximum Likelihood The likelihood for the binomial model is given by the following equation

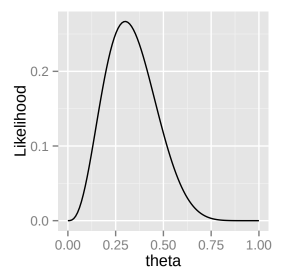
$$p(y|\theta) = \binom{n}{y} \theta^y (1-\theta)^{n-y}$$

•

The likelihood values for different values of θ are therefore as follows

```
likelihood <- function(theta, n = 10, y = 3) {
    choose(n, y) * theta^y * (1 - theta)^(n - y)
}
theta <- seq(from = 0, to = 1, length.out = 100)

qplot(theta, likelihood(theta), geom = "line", ylab = "Likelihood", xlab = "theta")</pre>
```



The estimate $(\hat{\theta})$ is the value of θ with the maximum likelihood (ML) value, which in this case is 0.3. A 95% confidence interval (CI) can then be calculated using the asymptotic normal approximation

$$\hat{\theta} \pm 1.96 \frac{1}{\sqrt{I(\hat{\theta})}}$$

where $I(\hat{\theta})$ is the expected second derivative of the log-likelihood at the estimate. This calculation is based on the assumption that the sample size is of sufficient size that the likelihood is normally distributed.

In the current case,

$$I(\hat{\theta}) = \frac{n}{\hat{\theta}(1-\hat{\theta})}$$

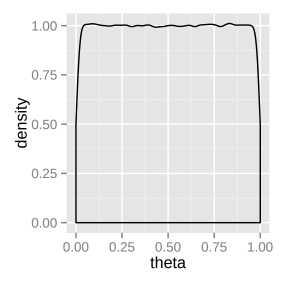
which gives a point estimate of 0.3 and lower and upper 95% confidence intervals of 0.02 and 0.58 respectively.

1.3.1.2 Posterior Probability The posterior probability on the other hand is given by Bayes rule which states that

$$p(\theta|y) \propto p(y|\theta)p(\theta)$$

where $p(\theta)$ is the prior probability.

Bayesians consider this an advantage because prior information can be incorporated into an analysis while frequentists consider it subjective. In most cases Bayesians use *low-information* priors which have very small effects on the posteriors. For example a uniform distribution with a lower limit of 0 and an upper limit of 1 (dunif(0,1)) is commonly used for probabilities.



As it is generally not possible to calculate the posterior probability, the posterior probability distribution is sampled using Markov Chain Monte Carlo (MCMC) algorithms such as Gibbs Sampling.

1.3.1.2.1 Gibbs Sampling Consider the case where the parameters $\theta = (\theta_1, \theta_2, \dots, \theta_k)$ then Gibbs Sampling proceed as follows

Step 1 Choose starting initial values for $\theta_1^{(0)}$ and $\theta_2^{(0)}$

Step 2 Sample
$$\theta_1^{(1)}$$
 from $p(\theta_1|\theta_2^{(0)}, y)$
Sample $\theta_2^{(1)}$ from $p(\theta_2|\theta_1^{(1)}, y)$

Step 3 *Iterate* step 2 thousands (or millions) of times to obtain a sample from $p(\theta|y)$.

Typically this is performed for two or three independent chains.

1.4 JAGS and BUGS

Programming an efficient MCMC algorithm for a particular model is outside the scope of most research projects. Fortunately, JAGS (which stands for Just Another Gibbs Sampler) can take a dataset and a model specified in the simple but flexible BUGS language (which stands for Bayesian Analysis Using Gibbs Sampling) and perform MCMC sampling for us.

In order to do this we will use the jaggernaut package to talk to the standalone JAGS program via the rjags package.

First we need to specify the underlying probability model in the BUGS language and save it as an object of class jags_model.

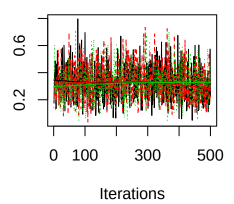
```
model1 <- jags_model("model {
  theta ~ dunif(0, 1)
    y ~ dbin(theta, n)
}")</pre>
```

then we call JAGS using jaggernaut in the default report mode to generate samples from θ 's posterior probability distribution.

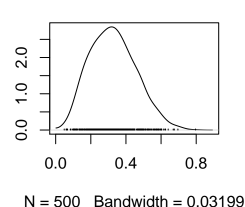
```
data <- data.frame(n = 10, y = 3)
analysis1 <- jags_analysis(model1, data = data)</pre>
```

plot(analysis1)

Trace of theta



Density of theta



coef(analysis1)

```
## estimate lower upper sd error significance
## theta 0.3271032 0.1074594 0.5983072 0.13028 75 0
```

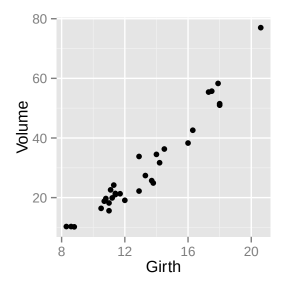
The model output indicates that the point estimate (in this case the mean of the samples) is 0.33 and the 95% credible interval (in this case the 2.25th and 97.75th percentiles) is 0.11 to 0.6. The model output also indicates that the posterior probability distribution has a standard deviation (sd) of 0.13. The significance and error values are discussed later.

Exercise 1 Previous studies indicate that the coin was definitely biased towards tails. Modify the prior distribution accordingly and rerun the above model. How does the posterior distribution change?

1.5 Black Cherry Trees

The trees data set in the dataset package provides information on the girth and volume of 31 black cherry trees.





Algebraically, the linear regression of Volume against Girth can be defined as follows

$$Volume_i = \alpha + \beta * Girth_i + \epsilon_i$$

where α is the intercept and β is the slope and the error terms (ϵ_i) are drawn from a normal distribution with an standard deviation of σ .

The model can be defined as follows in the BUGS language where \leftarrow indicates a *deterministic* as opposed to *stochastic* node (which is indicated by \sim).

```
model1 <- jags_model("model {
    alpha ~ dnorm(0, 50^-2)
    beta ~ dnorm(0, 10^-2)
    sigma ~ dunif(0, 10)

for(i in 1:length(Volume)) {
    eMu[i] <- alpha + beta * Girth[i]
    Volume[i] ~ dnorm(eMu[i], sigma^-2)
    }
}")</pre>
```

The standard deviations of the normal distributions are raised to the power of -2 because (for historical reasons) Bayesians quantify variation in terms of the precision (τ) as opposed to the variance (σ^2) or standard deviation (σ) where $\tau = 1/\sigma^2$.

1.5.1 Parallel Chains

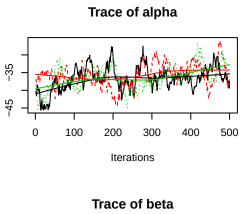
To reduce the analysis time, MCMC chains can be run on parallel processes. In jaggernaut this achieved using the registerDoParallel function and by setting the parallel option to be TRUE. This only needs to be done once at the start of a session.

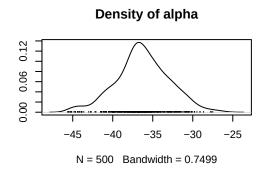
```
registerDoParallel()
opts_jagr(parallel = TRUE)
```

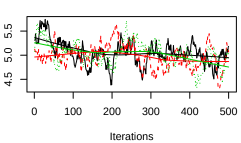
The resultant trace plots and coefficients for the trees analysis are as follows.

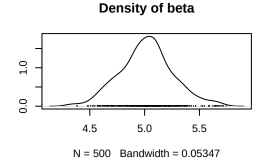
```
data(trees)
analysis1 <- jags_analysis(model1, data = trees)</pre>
```

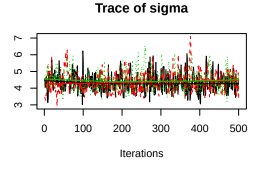
plot(analysis1)

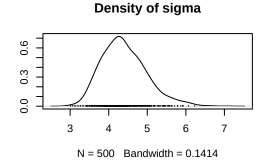












coef(analysis1)

##		estimate	lower	upper	sd	error	significance
##	alpha	-36.262113	-43.597545	-29.717354	3.36420	19	0
##	beta	5.017273	4.546653	5.539521	0.24571	10	0
##	sigma	4.405349	3.462012	5.779424	0.58384	26	0

Exercise 2 What do you notice about the trace plots? The output of auto_corr(analysis1) and cross_cor(analysis1) might give you some clues.

1.5.2 Chain Mixing

Cross-correlations between parameters, which cause poor chain mixing (i.e., high auto-correlation), can sometimes be eliminated or reduced by reparameterising the model. In the current model, Girth can be centered, i.e., Girth - mean(Girth), using the following code select_data(model1) <- c("Volume", "Girth+").

Exercise 3 What is the effect of centring Girth on the trace plots?

Note if you ever want to examine the actual data being passed to JAGS set the modify_data term of your jags_model object to be a simple function that prints and returns its one argument

```
modify_data(model1) <- function (data) { print(data); data }</pre>
```

1.5.2.1 glm Module In the case of generalized linear mixed models the JAGS glm module uses block updating to free the user of the need to centre predictor variables. It currently only works on parameters that have a normal prior distribution. The glm module can be loaded using the following code. The list.models() function lists the currently loaded modules.

```
library(rjags)
list.modules()
load.module("glm")
list.modules()
unload.module("glm")
list.modules()
```

For further information see the JAGS User Manual.

Exercise 4 What is the effect of loading the glm module (without manually centring Girth) on the trace plots?

1.5.3 Predictions and Residuals

Many researchers estimate fitted values, predictions and residuals by monitoring additional nodes in their model code. The disadvantages of this approach are that:

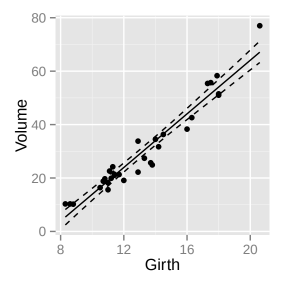
- the model code becomes more complicated.
- the MCMC sampling takes longer.
- adding derived parameters requires a model rerun.
- the table of parameter estimates becomes unwiedly.

jaggernaut overcomes these problems by allowing derived parameters to be defined in a separate chunk of BUGS code as demonstrated below.

```
dcode <- "data {
  for(i in 1:length(Volume)) {
    prediction[i] <- alpha + beta * Girth[i]
  }
  residual <- (Volume - prediction) / sigma
}"

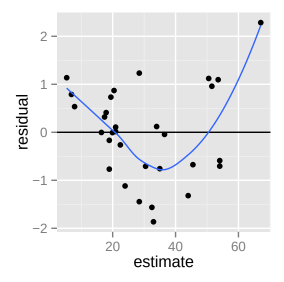
prediction <- predict(analysis1, newdata = "Girth", derived_code = dcode)</pre>
```

```
gp <- ggplot(data = prediction, aes(x = Girth, y = estimate))
gp <- gp + geom_point(data = dataset(analysis1), aes(y = Volume))
gp <- gp + geom_line()
gp <- gp + geom_line(aes(y = lower), linetype = "dashed")
gp <- gp + geom_line(aes(y = upper), linetype = "dashed")
gp <- gp + scale_y_continuous(name = "Volume")</pre>
print(gp)
```



```
fitted <- fitted(analysis1, derived_code = dcode)
fitted$residual <- residuals(analysis1, derived_code = dcode)$estimate

qplot(estimate, residual, data = fitted) + geom_hline(yintercept = 0) +
    geom_smooth(se = FALSE)</pre>
```



1.5.3.1 Allometry As discussed in the R course notes the relationship between Volume and Girth is expected to be allometric because the cross-sectional area at an given point scales to the square of the girth (circumference).

Expressed as an allometric relationship the model becomes

$$Volume_i = \alpha * Girth_i^{\beta} * \epsilon_i$$

which can be reparameterised as a linear regression by log transforming Volume and Girth.

$$log(Volume_i) = \alpha + \beta * log(Girth_i) + \epsilon_i$$

Variables can be log transformed in the model code or in the select_data argument, i.e., select_data(model1) <- c("log(Volume)", "log(Girth)").

Exercise 5 Fit the above allometric model to the trees data set. Is the model fit improved?

Exercise 6 Does adding log transformed Height improve the model?

1.5.4 Significance Values

The significance value in the jaggernaut table of coefficients is twice the probability that the posterior distribution spans zero. As such it represents the Bayesian equivalent of a frequentist two-sided p-value (Greenland and Poole 2013). By definition parameters that represent standard deviations, which must be greater than zero, will have a significance value of 0.

1.5.5 Error Values

The error value in the table of coefficients is the *percent relative error*, which is half the credible interval as a percent of the point estimate, i.e.,

$$error = (upper - lower) * 0.5/estimate * 100$$

Standard deviations with a uniform prior distribution that is not updated by the data have error values of 0.95. As a general rule, I question the informativeness of parameters representing standard deviations which have an error value ≥ 0.8 .

References

Greenland, Sander, and Charles Poole. 2013. "Living with P Values: Resurrecting a Bayesian Perspective on Frequentist Statistics." *Epidemiology* 24 (1): 62–68. doi:10.1097/EDE.0b013e3182785741.