# Tutorial on Bayesian Statistics: Replicating Sections of BDA3

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#### 1 - Replicating example from pg 66-67

Chapter 3 of BDA presents a first exercise of simulation (pgs 66-67) where from a sample of 66 obs with mean 26.2 and standard deviation 10.8, a posterior for the mean is simulated assuming a joint normal likelihood and uninformative prior (where  $\sigma^2 | y \sim Inv - \chi^2(n-1, s^2)$  and  $\mu | \sigma^2, y \sim N(\bar{y}, \sigma^2/n)$ ).

```
# Draw 1000 random numbers from a Inverse chi-squared
f.sim.sigma = function(draws) (n-1)*(se^2)/rchisq(draws,n-1)
f.sim.norm = function(draws) rnorm(draws,ybar,(f.sim.sigma(draws)/(n-1))^.5)
sim.norm = f.sim.norm(1000)
ci.mu = sim.norm[order(sim.norm)][c(25,975)]
print(ci.mu)
```

## [1] 23.45 28.83

```
# And now repeating the similation excercise R = 100 times:
R = 100
R.sim.norm = sapply(1:R, function(x) f.sim.norm(1000))
ci.mu = t(apply(R.sim.norm,2,function(x) x[order(x)][c(25,975)]))
print(c(median_low=median(ci.mu[,1]), median_high=median(ci.mu[,2]), p10_low =quantile(ci.mu[,1],c(.1),
## median_low median_high p10_low.10% p90_up.90%
```

29.04

#### **2** - Simulating values from a postetior normal with $\sigma^2$ and $\mu$ unknown and conjugate prior.

23.37

The following is not an exercise in the book (that I'm aware of), but I though that it would be interesting to work on this. After going over the model derived in pg 67-69 in BDA3, I will simulate values from the joint posterior (and hopefully leave the code flexible enough to play later on with the hyper-parameters).

With the prior density:

23.53

##

28.87

$$p(\mu, \sigma^2) \propto \sigma^{-1}(\sigma^2)^{(\nu_0/2+1} exp\left(-\frac{1}{2\sigma^2}[\nu_0\sigma_0^2 + \kappa_0(\mu_0 - \mu)^2]\right)$$

And a likelihood:

$$p(y|\mu,\sigma^2) \propto \sigma^{-n} exp\left(-\frac{1}{2\sigma^2}[(n-1)s^2 + n(\bar{y}-\mu)^2]\right)$$

We obtain the following joint posterior:

$$p(\mu, \sigma^{2}|y) \propto \sigma^{-1}(\sigma^{2})^{(\nu_{0}/2+1}exp\left(-\frac{1}{2\sigma^{2}}[\nu_{0}\sigma_{0}^{2} + \kappa_{0}(\mu_{0} - \mu)^{2}]\right) \times \\ \times (\sigma^{2})^{-n/2}exp\left(-\frac{1}{2\sigma^{2}}[(n-1)s^{2} + n(\bar{y} - \mu)^{2}]\right) \\ = N - Inv - \chi^{2}(\mu_{n}, \sigma_{n}^{2}; \nu_{n}, \sigma_{n}^{2}) \\ \mu_{n} = \frac{\kappa_{0}}{\kappa_{0} + n}\mu_{0} + \frac{n}{\kappa_{0} + n}\bar{y} \\ \kappa_{n} = \kappa_{0} + n \\ \nu_{n} = \nu_{0} + n \\ \nu_{n}\sigma_{n}^{2} = \nu_{0}\sigma_{0}^{2} + (n-1)s^{2} + \frac{\kappa_{0}n}{\kappa_{0} + n}(\bar{y} - \mu_{0})^{2}.$$

Using the fact that  $p(\mu, \sigma^2|y) = p(\mu|\sigma^2, y)p(\sigma^2|y)$  with:

$$\mu | \sigma^2, y \sim N(\mu_n, \sigma_n^2 / \kappa_n)$$
  
 $\sigma^2 | y \sim Inv - \chi^2(\nu_n, \sigma_n^2) \ labelpost.sig$ 

The simulation algorithm is as follows:

Draw a random number from the conditional posterior of sigma (equation \ref{post.sig})

Using that value of  $\sigma^2$ , draw a random number for the conditional posterior of  $\mu$  (equation \ref{post.mu})

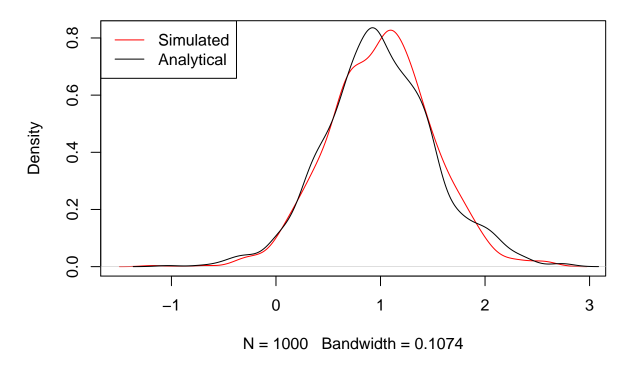
The following code implements this algorithm.

```
# Hyperparameters
p.mu_0
p.sigma2_0 = 1
p.nu_0
p.kappa_0 = 3
# Data
            = 10
n
            = 1.3
y_bar
            = 2
s
# Simulation
           = ((p.kappa_0)/(p.kappa_0 + n)) * p.mu_0 + ((n)/(p.kappa_0 + n)) * y_bar
mu_n
          = p.kappa_0 + n
kappa_n
            = p.nu_0 + n
nu_n
            = (p.nu_0*p.sigma2_0 + (n-1)*s^{2} + ((p.kappa_0*n)/(p.kappa_0 + n)) * (y_bar - p.mu_0)^{{t}}
sigma2_n
set.seed(142857)
# Draw 1000 random numbers from a Inverse chi-squared
f.mu.post = function(draws) {
  sigma2.cond = (nu_n)*(sigma2_n)/rchisq(draws,nu_n)
              = rnorm(draws,mu_n,(sigma2.cond/(kappa_n))^.5)
  return (mu.cond)
  }
sim.mu.post= f.mu.post(1000)
plot(density(sim.mu.post), main = "Draws from the marginal of the mean", col="red")
```

legend("topleft",col = c("red","black") ,legend=c("Simulated","Analytical"),lty = c(1,1))

lines(density(mu\_n +((sigma2\_n/kappa\_n)^.5)\*rt(1000,nu\_n)))

## Draws from the marginal of the mean



The plot above should be in 3D  $(\sigma^2, \mu, p(\sigma^2, \mu|y))$  but still don't know how to do that in R.

It also happens that for this particular problem there is a close form solution for the marginal of  $\mu$ . (remember that  $p(\mu|y) = \int p(\mu|\sigma^2, y)p(\sigma^2|y)d\sigma^2$ ):

$$\mu | \sigma^2, y \sim t_{\nu_n}(\mu | \mu_n, \sigma_n^2 / \kappa_n)$$

I'm still confused with the following why is that to simulate the marginal of  $\mu$  ( $\mu | \sigma^2, y$ ) I just need to draw from the conditional posterior of the mean (equation \ref{post.mu}). More generally, my confusion is the following: why is that to simulate  $h(x) = \int g(x)dF(x)$  I just need to draw  $x^s$  from dF(x) and evaluate  $g(x^s)$ . I know from MC simulation that by the LLN  $E_{dF}[g(x)] = \int g(x)dF(x) \approx \sum_s g(x^s)/S$ , but I'm having trouble seen why is that this applies to all points in the distribution (and not just the mean).

My own explanation so far: as MC can be used to approximate the mean using the LLN, the same technique can be used to approximate the whole distribution using CLT.

Simulating numbers from a Dirichlet distribution

```
draws = 1000
#This are the parameters of the gamma used in generating the final Dirichlet.
gamma.scale = c(728,538,138)
aux1 = sapply(gamma.scale,function(x) rgamma(100,x))
aux1 = aux1 / apply(aux1,1,sum)
```

#### 3 - Replicating Simulation Example 3.7 from BDA3 (pg 74+)

We are interested in modeling the dose-response of a certain drug  $(x_i)$  over the number of dead's  $(y_i)$  in a group of trial animals  $(n_i)$ . We have 4 observations. Defining  $\theta_i$  as the true death rate for each dosage, we can model this phenomena using a binomial distribution  $(y_i \sim Bin(n_i, \theta_i))$ . To model the dose-response relationship we start by looking at  $\theta_i = \alpha + \beta x_i$  but we realize that this model predicts values out of range  $(\theta, \alpha)$  as a probability has to be between 0 and 1). Hence we apply the logit  $(log(\theta_i/(1-\theta_i)))$  transformation. This implies the following likelihood.

$$p(y_i|\alpha,\beta,n_i,x_i) \propto [logit^{-1}(\alpha+\beta x_i)]^{y_i}[1-logit^{-1}(\alpha+\beta x_i)]^{n_i-y_i}$$

Question: Why don't we apply transformation method (multiply by the inverse of the Jacobian) to the likelyhood above?

**Answer:** Because we are not applying a transformation to  $\theta$  we are only replacin theta by

Using a non-informative prior  $(p(\alpha, \beta) \propto 1)$  we get that the conditional posterior has the form:

$$p(\alpha, \beta|y, n, x) \propto \prod_{i=1}^{k} p(y_i|\alpha, \beta, n_i, x_i)$$

First we compute, as a rough approximation of the parameters, the ML estimates.

```
# Data
x = c(-0.86, -0.30, -0.05, 0.73)
y = c(0,1,3,5)
n = rep(5,4)

# ML estimation
mlfit <- glm( cbind(y, n-y) ~ x, family = binomial)
mlfit$coefficients = round(mlfit$coefficients*1000)/1000
res1 = paste(paste("(",mlfit$coefficients[1], sep=""), paste(mlfit$coefficients[2],")", sep=""), sep="",</pre>
```

With the estimates  $(\hat{\alpha}, \hat{\beta}) = (0.847, 7.749)$ .

Now we will simulated values of  $\alpha$  and  $\beta$  from the posterior distribution. The approach is as follows.

• (i) Build a grid for all possible values of  $\alpha$  and  $\beta$ . Although this can be the whole real line, we use the ML estimates and some trial and error to restrict the space to  $\alpha \in [-5, 10]$  and  $\beta \in [-10, 40]$ .

```
alpha = seq(-5,10,.1)
beta = seq(-10,40,1)
```

• (ii) Compute the posterior density (equation \ref{post.1}) over the whole grid and normalize it to 1.

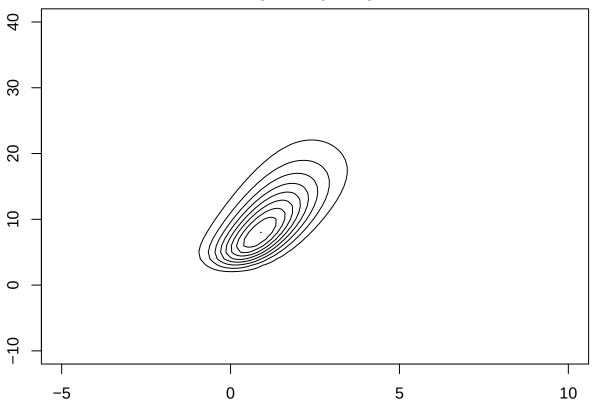
```
post.a = function(a,b,x,y,n) {
  post = 1
  for (i in 1:length(y)) {
    post = post * ( ((inv.logit(a+b*x[i]))^y[i])*((1-inv.logit(a+b*x[i]))^(n[i]-y[i])) )
    }
  post
}

post.dens = outer(alpha,beta,function(x1,x2) post.a(x1,x2,x,y,n))
post.dens = post.dens/sum(post.dens)
```

• (iii) Inspect the density using a contour plot (here we are looking for some indication that we are covering all the possible domain)

```
contours <- seq(min(post.dens), max(post.dens) , length=10)
par(mar = rep(2, 4))
contour(alpha, beta, post.dens,levels=contours, xlab=expression(alpha), ylab=expression(beta), xlim=c(max)</pre>
```

## Contour plot of joint posterior



• (iv) Compute the marginal posterior of  $\alpha$  by summing over all  $\beta$  for each value of  $\alpha$ . Notice so far that we had only compute the probability distribution of  $\alpha$  and  $\beta$ . Is not until this point where we would be able to do things like compute the mean, median and CI's of these parameters.

```
samps = 1000
alpha.dens = apply(post.dens ,1, sum)
```

- (v) For  $s = 1 \dots 1000$ :
  - a. Draw  $\alpha^s$  from its marginal posterior.

```
s.alpha = sample(alpha, samps, replace=TRUE, prob = alpha.dens)
```

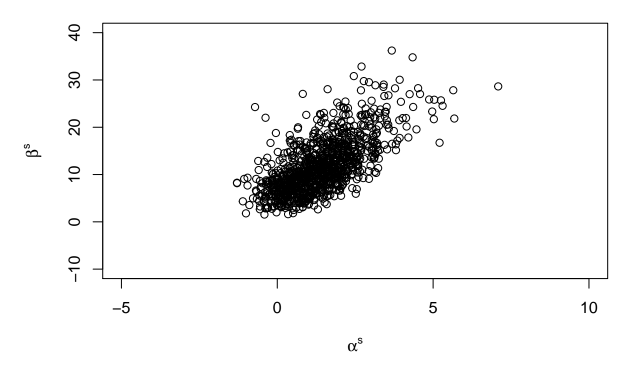
b. Draw  $\beta^s$  from  $p(\beta|\alpha, y)$  for each value of  $\alpha^s$ 

```
#Select the colum of the joint density corresponding to a specific value of alpha (p(beta/alpha))
cond.beta = function(x) {
  post.dens[which(alpha == s.alpha[x]),]
}
#Sample a value of beta according the the conditional probatility above
s.beta = sapply(1:samps,function(x) sample(beta,1,replace=TRUE,prob=cond.beta(x)))
```

c. For each sample values of  $\alpha$  and  $\beta$  add a uniform random jitter centered at zero with with equal to the grid spacing. This will make the simulation draws more continuous. Plot the sampled values.

```
s.beta = s.beta + runif(length(s.beta),-.5,.5)
s.alpha = s.alpha + runif(length(s.alpha),-.05,.05)
plot(s.alpha, s.beta, xlab=expression(alpha^s), ylab=expression(beta^s), xlim=c(min(alpha),max(alpha))
```

## Scatter Plot of Sample Draws of alpha and beta



The final result are two vectors  $(\{\alpha^s\}), (\{\beta^s\}),$  that represent probability distribution of each parameter.

4 - Description with my own words, of the fully Bayesian analysis of conjugate hierarchical models described in section 5.3 of BDA (pg 108 - 113)

The overall goal of hierarchical models is to describe a statistical (uncertain) phenomena where are multiple parameters involved and exists a dependence structure.

The analysis described here builds on what we just did in the example above. For this purpose we enumerate the steps followed, now in a more general notation.

- 1. Compute the joint posterior probability distribution of all the parameters conditional on the data and a prior distribution of those parameters (in the example above we use a non-informative prior).
- 2. Compute the marginal posterior of one parameter  $(p(\theta_2|y))$  by summing the joint posterior over all the possible values of the other parameter  $(\theta_1)$ .
- 3. Compute the marginal of the other parameter  $(p(\theta_1|y))$  using the following identity:

$$p(\theta_1|y) = \int p(\theta_1|\theta_2, y) p(\theta_2|y) d\theta_2 \approx p(\theta_1|\theta_2^s, y)$$

Where  $\theta_2^s$  is a random draw using  $p(\theta_2|y)$  (the justification for this last step is in section 2 of this document).

Now we will distinguish between two sets of parameters. The original parameters of interest, still represented by  $\theta$ . And the parameters of the prior distribution, or *hyperparameters* represented by  $\phi$ . This two sets of parameters fully describe the probabilistic process and have a joint distribution function:

$$p(\phi, \theta) = p(\phi)p(\theta|\phi)$$

And a joint posterior distribution:

$$p(\phi, \theta|y) \propto p(\theta, \phi)p(y|\theta, \phi)$$
$$= p(\phi)p(\theta|\phi)p(y|\theta)$$

Where in the last line we used the assumption that  $p(y|\theta,\phi)$  depends on  $\phi$  only through  $\theta$ . We call  $p(\phi)$  the hyperprior distribution,  $p(\theta|\phi)$  the population distribution, and the usual suspect  $p(y|\theta)$  is the likelihood, or sampling, distribution.

As always our goal is to make inference about  $\theta$  (and maybe  $\phi$  also?). As in the exercise before, our end result will be a set of matrices  $(\{\phi^s\}), (\{\theta^s\})$  with values for the parameters that follow the posterior joint distribution. To get this result we take the following steps:

- 1. Compute the conditional joint posterior  $p(\phi, \theta|y)$ . Choosing the right hyper-prior distribution is a whole topic on itself and will be addressed later.
- 2. Compute the marginal posterior of  $\theta$ , conditional on  $\phi$ ,  $p(\theta|\phi,y)$ . This conditional posterior of  $\theta$  can be obtain analytically in conjugate models for a given value of  $\phi$ , it is simply the posterior of the non-hierarchical Bayesian case.
- 3. Compute the posterior marginal of  $\phi$ . When step 2 has a closed form solution we can compute the marginal of  $\phi$  as:

$$p(\phi|y) = \frac{p(\theta, \phi|y)}{p(\theta|\phi, y)}$$

In the absence of a closed form solution for step 2. We can compute the marginal by integrating the joint over all the possible values of  $\theta$ .

- 4. Draw samples  $\phi^s$  from  $p(\phi|y)$ . Notice that  $\phi$  can have multiple components. If it has more than one element we follow the procedure described at the beginning of this section.
- 5. For each draw of  $\phi^s$ , draw a sample of  $\theta$  from  $p(\theta|\phi^s,y)$ . This allow us to fully characterize the parameter of interest (our goal, remember?)

#### Application

We have data from  $j=1\ldots J, J=71$  experiments where, in each experiment  $n_j$  rats were exposed to a drug and  $y_j$  of them, presented tumors afterwards (think also something similar to unemployment rate in different states with  $n_j$  people in the active labor force and  $y_j$  unemployed). The results from the experiments are assumed to follow a binomial distribution  $(y_j|\theta_j \sim Bin(n_j,\theta_j))$ , and the parameters  $\theta_j$  are assume to follow a beta prior distribution  $(\theta_j|\alpha,\beta \sim Beta(\alpha,\beta))$ , this last assumption is made to take advantage of conjugacy. With all this elements now we can follow the steps described above.

1. Compute the conditional joint posterior  $p(\phi = (\alpha, \beta), \theta = \{\theta_i\}_{i=1}^{J} | y)$ .

$$p(\theta, \alpha, \beta|y) \propto p(\alpha, \beta)p(\theta|\alpha, \beta)p(y|\theta, \alpha, \beta)$$

$$p(\theta, \alpha, \beta|y) \propto p(\alpha, \beta) \prod_{j=1}^{J} \frac{\Gamma(\alpha + \beta)}{\Gamma(\alpha)\Gamma(\beta)} \theta_j^{\alpha-1} (1 - \theta_j)^{\beta-1} \prod_{j=1}^{J} \theta_j^{y_j} (1 - \theta_j)^{n_j - y_j}$$

2. Compute the marginal posterior of  $\theta$ , conditional on  $\alpha$ ,  $\beta$ . For the beta-binomial case we have that given the hyper-parameters, each  $\theta_j$  has a posterior distribution  $Beta(\alpha + y_j, \beta + n_j - y_j)$ . Assuming exchangeability:

$$p(\theta|\alpha,\beta,y) = \prod_{j=1}^{J} \frac{\Gamma(\alpha+\beta+n_j)}{\Gamma(\alpha+y_j)\Gamma(\beta+n_j-y_j)} \theta_j^{\alpha+y_j-1} (1-\theta_j)^{\beta+n_j-y_j-1}$$

3. Compute the posterior marginal of  $(\alpha, \beta)$ . Given that we do have a closed form solution in step 2, we compute the ratio of (\ref{rat.joint.post}) and (\ref{rat.cond.post.theta}).

$$p(\alpha, \beta | y) \propto p(\alpha, \beta) \prod_{j=1}^{J} \frac{\Gamma(\alpha + \beta)}{\Gamma(\alpha)\Gamma(\beta)} \frac{\Gamma(\alpha + y_j)\Gamma(\beta + n_j - y_j)}{\Gamma(\alpha + \beta + n_j)}$$

And here is the code for this function:

```
rat.marg.post.phi = function(alpha, beta) {
  post = 1
  for (i in 1:length(y)) {
    post = post * ( ( gamma(alpha + beta) )/( gamma(alpha)*gamma(beta) ) ) * ( ( gamma(alpha + y[i] )
    }
    # The hyper prior is defined below
    rat.hyper.prior(alpha,beta) * post
}
```

**Note:** we assume a hyper-prior of the form  $p(\alpha, \beta) \propto (\alpha + \beta)^{-5/2}$ . So far I do not understand very well the whole discussion about proper and improper priors. **Need to review Ch10 of DeGroot's book** 

4. Draw samples  $(\alpha^s, \beta^s)$  from  $p(\alpha, \beta|y)$ . Before drawing the samples, the authors applied the following transformation to the parameter space:  $(\alpha, \beta) \to (log(\frac{\alpha}{\beta}), log(\alpha + \beta))$ . I still don't know why they do this and why is that the transformation only applies to the hyper-prior only. This step requires a important number of sub-steps:

4.1. Compute the (prior) probability distribution of the transformed variables. Here is where we multiply by the Jacobian of the inverse transformation. The transformation method states that if  $u \sim p_u(u)$  and there is a 1:1 function over u, v = f(u), then  $v \sim p_u(u)|J|$  where the J is the Jacobian of the function  $f^{-1}(v)$  with respect to v. For this case we have  $v_1 = f_1(\alpha, \beta) = \log(\alpha/\beta)$  and  $v_2 = f_2(\alpha, \beta) = \log(\alpha+\beta)$ , with inverse  $f_1^{-1}(v_1, v_2) = \exp(v_1 + v_2)/(1 + \exp(v_1))$  and  $f_2^{-1}(v_1, v_2) = \exp(v_2)/(1 + \exp(v_1))$ . Hence the Jacobian is:

$$J = \begin{bmatrix} \frac{\partial f_1^{-1}}{\partial v_1} & \frac{\partial f_1^{-1}}{\partial v_2} \\ \frac{\partial f_2^{-1}}{\partial v_1} & \frac{\partial f_2^{-1}}{\partial v_2} \end{bmatrix} = -\alpha\beta$$

Hence, using the transformation method we have that  $p(\log(\frac{\alpha}{\beta}), \log(\alpha + \beta)) \propto \alpha\beta(\alpha + \beta)^{-5/2}$ . And here is the code for this function:

```
c(0,
                   0,
                       0,
                           0, 0,
                                    0, 0,
                                            Ο,
                           1,
                                    1,
                                        1,
                                2,
                       2.
                            2,
                                    2,
                                        2,
                                            2,
                                                2,
                            2,
                                7,
                                    7,
                                        3,
                                            3,
                                                2,
                       4,
                            4,
                               4,
                                    4,
                                        4, 10,
                                                4,
               5, 11, 12,
                           5,
                               5,
                                    6,
                                       5,
                                            6,
              16, 15, 15,
                           9,
                               4)
          = c(20, 20, 20, 20, 20, 20, 20, 19, 19, 19, 19,
n
              18, 18, 17, 20, 20, 20, 20, 19, 19, 18, 18,
              27, 25, 24, 23, 20, 20, 20, 20, 20, 20, 10,
              49, 19, 46, 17, 49, 47, 20, 20, 13, 48, 50,
              20, 20, 20, 20, 20, 20, 20, 48, 19, 19, 19,
              22, 46, 49, 20, 20, 23, 19, 22, 20, 20, 20,
              52, 46, 47, 24, 14)
rat.hyper.prior = function(alpha,beta) {
  alpha*beta*(alpha + beta)^(-5/2)
}
```

4.2. Identify the relevant domain for the transformed variables and its counterpart with the original variables. To do so we start computing the parameters  $\alpha$  and  $\beta$  that would match the sample mean and standard deviation of all 70 experiments with a beta prior distribution:

$$\hat{\mu} = 0.1381 = \frac{\hat{\alpha_0}}{\hat{\alpha_0} + \hat{\beta_0}}$$

$$\hat{\sigma^2} = 0.0109 = \frac{\hat{\alpha_0}\hat{\beta_0}}{(\hat{\alpha_0} + \hat{\beta_0})^2(\hat{\alpha_0} + \hat{\beta_0} + 1)}$$

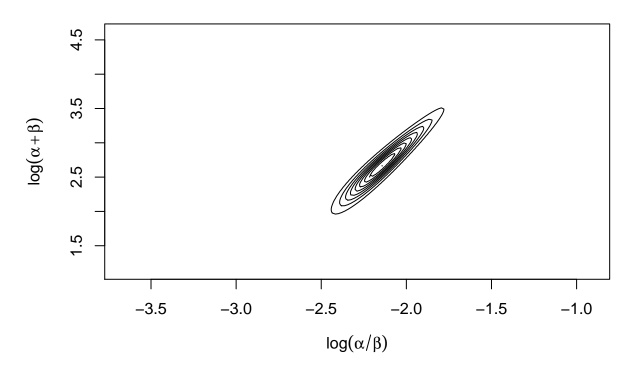
Solving for  $(\hat{\alpha_0}, \hat{\beta_0})$ :

We get:  $(\hat{\alpha_0}, \hat{\beta_0}) = (1.4, 8.6)$ .

We center the grid (approximately) around that initial estimate and expand the grid to cover up to a factor of 4 of each parameter. The result is plotted in the following figure:

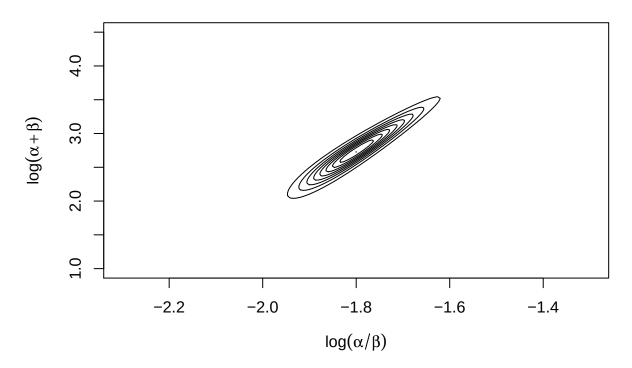
```
post.dens = post.dens/sum(post.dens)
contours <- seq(min(post.dens), max(post.dens) , length=10)
contour(v1, v2, post.dens,levels=contours, xlab=expression(log(alpha/beta)), ylab=expression(log(alpha+beta))</pre>
```

## Contour plot of joint posterior



4.3. Recalculate the range of the grid such that includes all the density

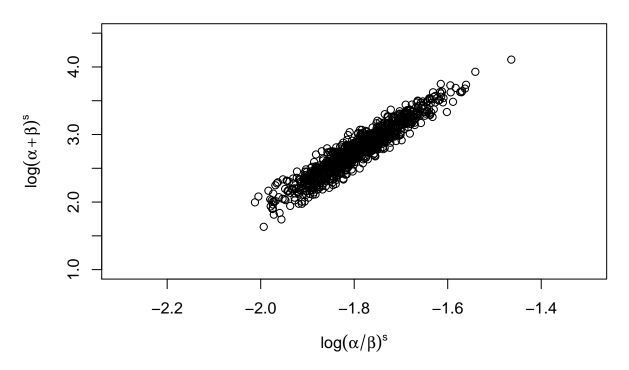
## Contour plot of joint posterior



4.4. Draw samples  $(\alpha^s, \beta^s)$  from  $p(\alpha, \beta|y)$  (finally!). Here we repeat the procedure used in section 3.(v) of this document.

```
samps
            = 1000
            = apply(post.dens ,1, sum)
v1.dens
            = sample(v1, samps, replace=TRUE, prob = v1.dens)
s.v1
#Select the colum of the joint density corresponding to a specific value of v1 (p(v2/v1))
cond.v2
            = function(x) {
  post.dens[which(v1 == s.v1[x]),]
#Sample a value of v2 according the the conditional probatility above
            = sapply(1:samps,function(x) sample(v2,1,replace=TRUE,prob=cond.v2(x)))
s.v2
#Add a uniform random jitter centered at zero with with equal to the grid spacing. This will make the s
            = v1[2]-v1[1]
grid.v1
            = v2[2]-v2[1]
grid.v2
            = s.v2 + runif(length(s.v2),-grid.v2/2,grid.v2/2)
s.v2
            = s.v1 + runif(length(s.v1),-grid.v1/2,grid.v1/2)
plot(s.v1, s.v2, xlab=expression(log(alpha/beta)^s), ylab=expression(log(alpha+beta)^s), xlim=c(min(v1)
```

## Scatter Plot of Sample Draws of log(alpha/beta) and log(alpha+beta



Note: this two figures do not match exactly their counterparts in the textbook (5.3a and 5.3b in BDA3), but so far I have not been able to detect my mistake.

4.5 By applying the inverse of the transformation we recover the marginal distribution of the original hyper-parameters.

```
s.beta = \exp(s.v2)/(\exp(s.v1)+1)
s.alpha = \exp(s.v2+s.v1)/(\exp(s.v1)+1)
```

5. For each draw of  $\phi^s$ , draw a sample of  $\theta$  from  $p(\theta|\phi^s,y)$ 

```
s.beta = \exp(s.v2)/(\exp(s.v1)+1)

s.alpha = \exp(s.v2+s.v1)/(\exp(s.v1)+1)

theta.dist = \operatorname{sapply}(1:71, \operatorname{function}(x) \operatorname{rbeta}(1000, s.alpha+y[x], s.beta + n[x] - y[x]))

theta.dist = \operatorname{apply}(\operatorname{theta.dist}, 2, \operatorname{sort})

\operatorname{plot}(0:450/1000, 0:450/1000, \operatorname{type="l"}, \operatorname{xlab="Observed rate"}, \operatorname{ylab="95\% CI and median of posterior"})

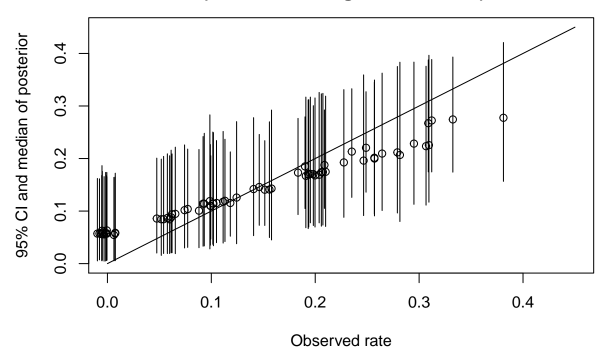
\operatorname{jitter.x} = \operatorname{y/n} + \operatorname{runif}(\operatorname{length}(y), -0.01, 0.01)

\operatorname{points}(\operatorname{jitter.x}, \operatorname{theta.dist}[500,])

\operatorname{segments}(\operatorname{jitter.x}, \operatorname{theta.dist}[25,], \operatorname{jitter.x}, \operatorname{theta.dist}[975,])

\operatorname{title}(\operatorname{main="Posterior Distribution of Tumor Rates for all 71 Experiments \n (Remember the goal? This is)
```

## Posterior Distribution of Tumor Rates for all 71 Experiments (Remember the goal? This is it!)



#### 5 - Replicating section 5.5: Experiments in eight schools (normal model) [Without Stan]

```
#Data:
school.id <- LETTERS[1:8]
effect <- c(28,8,-3,7,-1,1,18,12)
se.effect <- c(15,10,16,11,9,11,10,18)

pool.est <- sum(effect*se.effect^-2)/sum(se.effect^-2)
pool.var <- sum(se.effect^-2)^-1
pool.ci <- c(-1.96,1.96)*pool.var^.5 + pool.est
```

The pooled estimated effect and variance are 7.69 and 16.58, with a 95% CI of [-0.3, 15.67].

Posterior simulation under ther hierarchical model Using the identity:

$$p(\theta, \mu, \tau | y) = p(\tau | y)p(\mu | \tau, y)p(\theta | \mu, \tau, y)$$

And the results from BDA in equation 5.17, 5.20 and 5.21 we code the joint posterior:

```
# Eqn 5.17 of BDA3
post.theta.j <- function(mu,tau,j) (effect[j]/(se.effect[j]^2) + mu/(tau^2)) / (1/(se.effect[j]^2) +
post.v.theta.j <- function(tau,j) 1/(1/(se.effect[j]^2) + 1/(tau^2))
# Eqn 5.20 of BDA3
post.mu.hat <- function(tau) sum(effect*1/(se.effect^2 +tau^2))/sum(1/(se.effect^2 +tau^2))</pre>
```

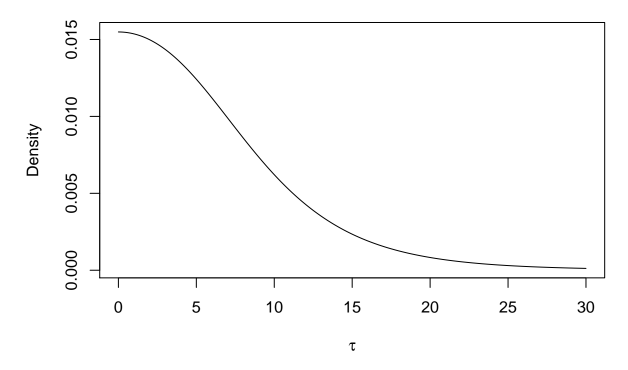
Define a hyper-prior and draw 200 samples from each distribution (for all 8 schools).

```
samps <- 200

hyper.prior <- function(tau) 1
tau.grid <- seq(0.001,30, length=samps)
pdf.tau <- sapply(tau.grid,function(x) marginal.tau(x))
pdf.tau <- pdf.tau/sum(pdf.tau)

plot(tau.grid,pdf.tau, type="l", main="Figure 5.5 from BDA3", xlab=expression(tau), ylab="Density")</pre>
```

Figure 5.5 from BDA3



The sampling method in BDA3 suggest to apply the inverse method from the posterior of  $\tau$ . I don't do this for two reasons: (i) I'm not sure the posterior has a closed for solution for its inverse, and (ii) given that I already have the density, I can directly draw from that distribution sampling using the sample command (which leads me to think that this command applies the inverse method, but **need to check with Susan**).

The following figures replicate the figures in pg 122 in BDA. Before doing the plots we need to 'average over  $\mu$ '

$$\begin{split} E(\theta_{j}|\tau,y) &= E_{\mu} \left[ E(\theta_{j}|\tau,y,\mu) | \tau,y \right] \\ &= E_{\mu} \left[ \frac{\frac{1}{\sigma_{j}^{2}} y_{j} + \frac{1}{\tau^{2}} \mu}{\frac{1}{\sigma_{j}^{2}} + \frac{1}{\tau^{2}}} | \tau,y \right] = \frac{\frac{1}{\sigma_{j}^{2}} y_{j} + \frac{1}{\tau^{2}} \hat{\mu}}{\frac{1}{\sigma_{j}^{2}} + \frac{1}{\tau^{2}}} \\ Var(\theta_{j}|\tau,y) &= E_{\mu} \left[ Var(\theta_{j}|\tau,y,\mu) | \tau,y \right] + Var_{\mu} \left[ E(\theta_{j}|\tau,y,\mu) | \tau,y \right] \\ &= \frac{1}{\frac{1}{\sigma^{2}} + \frac{1}{\tau^{2}}} + V_{\mu} \left( \frac{\frac{1}{\tau^{2}}}{\frac{1}{\sigma^{2}} + \frac{1}{\tau^{2}}} \right) \end{split}$$

Where  $V_{\mu}$  and  $\hat{\mu}$  correspond to the expressions defined in Eq 5.20 of BDA3. Below is the code and plot of both equations.

```
post.theta.j.no.mu <- function(tau,j) post.theta.j(post.mu.hat(tau),tau,j)
post.se.theta.j.no.mu <- function(tau,j) sqrt( (post.v.theta.j(tau,j)) * (1+post.v.mu(tau)*tau^(-2)) )

plot( tau.grid,sapply(tau.grid, function(x) post.theta.j.no.mu(x,1)), type="l", ylim=c(-5,30), xlab="",
lines(tau.grid,sapply(tau.grid, function(x) post.theta.j.no.mu(x,2)))
lines(tau.grid,sapply(tau.grid, function(x) post.theta.j.no.mu(x,3)))
lines(tau.grid,sapply(tau.grid, function(x) post.theta.j.no.mu(x,4)))
lines(tau.grid,sapply(tau.grid, function(x) post.theta.j.no.mu(x,5)))
lines(tau.grid,sapply(tau.grid, function(x) post.theta.j.no.mu(x,6)))
lines(tau.grid,sapply(tau.grid, function(x) post.theta.j.no.mu(x,7)))
lines(tau.grid,sapply(tau.grid, function(x) post.theta.j.no.mu(x,8)))
title(main="Figure 5.6 from BDA3", xlab=expression(tau), ylab="Estimated treatment effect")</pre>
```

Figure 5.6 from BDA3

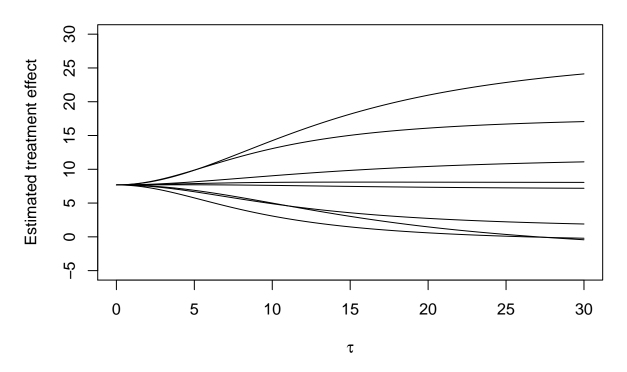


Figure 5.7 from BDA3

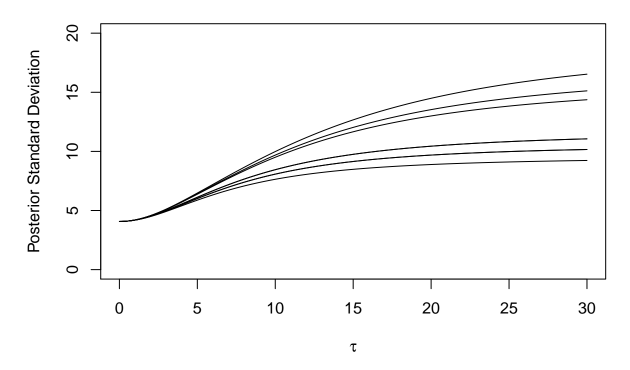


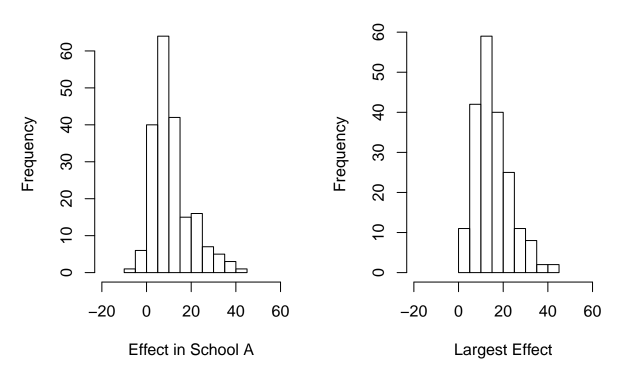
Table 5.3 from BDA3:

School					
	2.5%	25%	median	75%	97.5%
A	-0.959	5.126	8.799	14.701	31.639
В	-4.707	3.321	7.599	11.384	20.985
$\mathbf{C}$	-15.041	2.191	6.602	10.44	17.38
D	-9.037	2.942	6.87	11.321	20.173
$\mathbf{E}$	-8.086	2.217	6.21	9.562	18.036
$\mathbf{F}$	-6.893	2.292	6.263	10.17	19.596
G	-1.023	5.646	9.295	14.107	29.198
Н	-7.684	4.477	7.525	12.355	24.877

Here we reproduce figure 5.8 (with the same problems as above)

```
par(mfrow=c(1,2))
domain          <- c(-20,60)
hist(s.theta[,1], breaks=10, xlab="Effect in School A", main="", xlim=domain)
hist(apply(s.theta,1,max), breaks=10, xlab="Largest Effect", main="", xlim=domain)</pre>
```

Figure 5.8 from BDA3



This last figure ("largest effect") is a good example of one the main advantage of a fully Bayesian hierarchical model: once we have correctly simulated the posterior, we can test all kinds of complicated hypothesis.

#### 6 - Replicating section 5.5: Experiments in eight schools (normal model) [With Stan]

Appendix C in BDA3

## 7 - Replicating example of section 11.2: Metropolis sampling from bivariate normal [Witout Stan]

Here we follow the steps in p278 to simulate draws from a bivariate normal  $N(0, I_2)$ .

1. Define a starting point  $\theta^0$ , for which  $p(\theta^0|y) > 0$ :

```
set.seed(142857)
library(mvtnorm)
theta.0 <- c(-2.5, 2.5)</pre>
```

Our starting point is (-2.5, 2.5), the upper left most square dot from figure 11.1a.

2. For  $t = 1, 2, \dots$ :

(a) Sample a proposal  $\theta^*$  at time t from  $J_t(\theta^*|\theta^{t-1})$ . For this example the jumping distribution is  $N(\theta^*|\theta^{t-1}, 0.2^2I_2)$ :

(b) Calculate the ratio of densities,

$$r = \frac{p(\theta^*|y)}{p(\theta^{t-1}|y)}$$

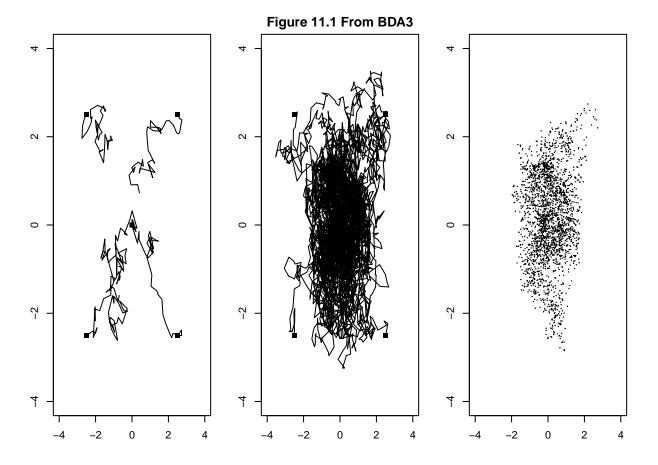
(c) Set

$$\theta^t = \begin{cases} \theta^* & \text{with probability } min(r,1) \\ \theta^{t-1} & \text{otherwise.} \end{cases}$$

(d) Execute:

```
theta
                 <- function(sims, theta.0) {
  s.theta
                 <- matrix(NA, sims, 2)
  s.theta[1,]
                 <- t(as.matrix(theta.0))
  for (t in 2:sims) {
    s.theta.star<- theta.star(s.theta[t-1,])</pre>
           <- r.dens(s.theta[t-1,], s.theta.star)</pre>
    s.theta[t,] <- theta.t(s.theta[t-1,], s.theta.star, r)</pre>
  }
return(s.theta)
par(mfrow=c(1,3))
par(mar = rep(2,4))
# 11.1a
s.theta.1
                 <- theta(50, theta.0)
s.theta.2
               \leftarrow theta(50,c(2.5,2.5))
s.theta.3
              \leftarrow theta(50,c(2.5,-2.5))
```

```
\leftarrow theta(50,c(-2.5,-2.5))
s.theta.4
s.theta.5
                \leftarrow theta(50,c(0,0))
plot(s.theta.1, xlim=c(-4,4), ylim=c(-4,4), type="l", sub = "50 simulations")
points(theta.0[1], theta.0[2], pch=15)
lines(s.theta.2)
points(2.5, 2.5, pch=15)
lines(s.theta.3)
points(2.5, -2.5, pch=15)
lines(s.theta.4)
points(-2.5, -2.5, pch=15)
lines(s.theta.5)
points(0, 0, pch=15)
# 11.1b
s.theta.1
                <- theta(1000, theta.0)
s.theta.2
                \leftarrow theta(1000, c(2.5,2.5))
                \leftarrow theta(1000, c(2.5,-2.5))
s.theta.3
                \leftarrow theta(1000, c(-2.5,-2.5))
s.theta.4
s.theta.5
                \leftarrow theta(1000, c(0,0))
plot(s.theta.1, xlim=c(-4,4), ylim=c(-4,4), type="l", sub = "1000 simulations", main="Figure 11.1 From
points(theta.0[1], theta.0[2], pch=15)
lines(s.theta.2)
points(2.5, 2.5, pch=15)
lines(s.theta.3)
points(2.5, -2.5, pch=15)
lines(s.theta.4)
points(-2.5, -2.5, pch=15)
lines(s.theta.5)
points(0, 0, pch=15)
                 <- s.theta.1[501:1000,] + matrix(runif(1000)/20, nrow = 500, ncol = 2)
s.theta.1
                 <- s.theta.2[501:1000,] + matrix(runif(1000)/20, nrow = 500, ncol = 2)
s.theta.2
s.theta.3
                <- s.theta.3[501:1000,] + matrix(runif(1000)/20, nrow = 500, ncol = 2)
                \leftarrow s.theta.4[501:1000,] + matrix(runif(1000)/20, nrow = 500, ncol = 2)
s.theta.4
s.theta.5
                <- s.theta.5[501:1000,] + matrix(runif(1000)/20, nrow = 500, ncol = 2)
# 11.1c
plot(s.theta.1, xlim=c(-4,4), ylim=c(-4,4), type="p", pch = 20, cex = .1, sub = "Second half of 1000 sim")
points(s.theta.2, pch = 20, cex =.1)
points(s.theta.3, pch = 20, cex =.1)
points(s.theta.4, pch = 20, cex =.1)
points(s.theta.5, pch = 20, cex =.1)
```



8 - Replicating example of section 11.6: Gibbs sampling for a hierarchical normal model (Coagulations experiment data) [Witout Stan]

#### The Model

Data  $y_{ij}$ ,  $i=1,...,n_j$ , j=1,...,J are iid within each of J groups with  $y_{ij} \sim N(\theta_j, \sigma^2)$ . The prior distribution of  $\theta_j$  is assume to be normal with hyperparameters  $\mu, \tau$  ( $\theta_j \sim N(\mu, \tau^2)$ ),  $\sigma$  is assumed to be unkonwn (confusing!) and the hyperprior is assumed to be uniform over  $(\mu, log(\sigma), \tau)$ , which implies  $p(\mu, log(\sigma), log(\tau)) \propto \tau$ . The joint posterior density for all the parameters is:

$$p(\theta, \mu, log(\sigma), log(\tau)|y) \propto \tau \prod_{j=1}^{J} N(\theta_j | \mu, \tau^2) \prod_{j=1}^{J} \prod_{i=1}^{n_j} N(y_{ij} | \theta_j, \sigma^2)$$

### $Starting\ points$

Select 10  $\theta_j$  randomly from the  $y_{ij}$  sample. And take  $\mu$  to be the average starting values of  $\theta_j$ .

```
set.seed(142857)
theta.0 <- sapply(1:4,function(x) sample(y[id==LETTERS[x]],10, replace=TRUE))
mu.0 <- apply(theta.0, 1,mean)</pre>
```

[Here I follow the exact reverse order that is proposed in the book, going from step 4 to 1 instead of 1 to 4. It is not clear to me how to do it otherwise].

4. Draw from conditional posterior of  $\tau^2$  using:

$$\hat{\tau}^2 = \frac{1}{J-1} \sum_{j=1}^{J} (\theta_j - \mu)^2$$

And the fact that:

$$\tau^2 | \theta, \mu, \sigma, y \sim Inv - \chi^2 (J - 1, \hat{\tau}^2)$$

We can draws samples for  $\tau^2$ 

```
s.tau.post <- function(theta) {
  tau.2 <- tau.hat.2(theta)
  tau.cond <- (J - 1) * (tau.2)/rchisq(1,J-1)
  return(tau.cond)
}</pre>
```

3. Draw from conditional posterior of  $\sigma^2$  Using:

$$\hat{\sigma}^2 = \frac{1}{n} \sum_{i=1}^{J} \sum_{i=1}^{n_j} (y_{ij} - \theta_j)^2$$

```
f.sigma.hat.2 <- function(theta) {
  sigma.hat.2 <- sapply(1:4, function(x) (y[id==LETTERS[x]] - theta[x])^2)
  sigma.hat.2 <- (1/n) * sum(unlist(sigma.hat.2))
  return(sigma.hat.2)
}</pre>
```

And the fact that:

$$\sigma^2 | \theta, \mu, \tau, y \sim Inv - \chi^2(n, \hat{\sigma}^2)$$

We can draws samples for  $\sigma^2$ 

```
s.sigma.post <- function(theta) {
  sigma2.hat <- f.sigma.hat.2(theta)
  sigma2.post <- (n) * (sigma2.hat)/rchisq(1,n)
  return(sigma2.post)
}</pre>
```

2. Draw from conditional posterior of  $\mu$  Using:

$$\hat{\mu} = \frac{1}{J} \sum_{j=1}^{J} \theta_j$$

```
mu.hat <- function(theta) {
  mean(theta)
}</pre>
```

And the fact that:

$$\mu | \theta, \sigma, \tau, y \sim N(\hat{\mu}, \tau^2/J)$$

We can draw values for  $\mu$ 

```
s.mu <- function(theta,tau2) {
  mu.hat <- mu.hat(theta)
  rnorm(1,mu.hat,sqrt(tau2/J))
}</pre>
```

1. Finally, we can draw values for  $\theta$  Using the fact that:

$$\theta_j | \mu, \sigma, \tau, y \sim N(\hat{\theta_j}, V_{\theta_j})$$

With:

$$\hat{\theta_j} = \frac{\frac{1}{\tau^2}\mu + \frac{n_j}{\sigma^2}\bar{y}\bar{j}}{\frac{1}{\tau^2} + \frac{n_j}{\sigma^2}}$$

$$V_{\theta_j} = \frac{1}{\frac{1}{\tau^2} + \frac{n_j}{\sigma^2}}$$

```
theta.hat.j <- function(j,mu,sigma2,tau2) {</pre>
              <- mean(y[id==LETTERS[j]])</pre>
  y.bar.j
 n.j
                <- length(y[id==LETTERS[j]])
  ( (1/tau2) * mu + (n.j/sigma2) * y.bar.j ) / ( (1/tau2) + (n.j/sigma2) )
V.theta.hat.j <- function(j,mu,sigma2,tau2) {</pre>
                <- length(y[id==LETTERS[j]])
  (1)/((1/tau2) + (n.j/sigma2))
}
s.theta
                <- function(mu,sigma2,tau2) {
  theta
                <- NULL
  for (j in 1:J) {
   t.hat
              <- theta.hat.j(j,mu,sigma2,tau2)</pre>
    v.t.hat
               <- V.theta.hat.j(j,mu,sigma2,tau2)
    theta[j] <- rnorm(1,t.hat,sqrt(v.t.hat))</pre>
  }
  return(theta)
}
```

```
<- function(chain) {
mcmc.gibbs
                <- as.list(NULL)
  res1
  sims
                <- 200
  param
                <- 7
                <- matrix(NA, ncol = param, nrow = sims )
  s.param
  colnames(s.param) <- c("theta1", "theta2", "theta3", "theta4", "mu", "sigma2", "tau2")
  s.param[1,1:4]<- theta.0[chain,]
  s.param[1,7] <- s.tau.post(theta.0[chain,])</pre>
  s.param[1,6] <- s.sigma.post(theta.0[chain,])</pre>
  s.param[1,5] \leftarrow s.mu(theta.0[chain,],s.param[1,7])
  for (s in 2:sims) {
    s.param[s,1:4] <- s.theta(s.param[s-1,5],s.param[s-1,6],s.param[s-1,7])
    s.param[s,7] \leftarrow s.tau.post(s.param[s,1:4])
    s.param[s,6] <- s.sigma.post(s.param[s,1:4])
    s.param[s,5] \leftarrow s.mu(s.param[s,1:4],s.param[s,7])
  }
return(s.param)
#Warm-up
}
                  <- lapply(1:10,function(x) mcmc.gibbs(x))
s.param
                  <- s.param[[1]][101:200, ]
s.param.1
#Transform the variance in to sd.
s.param.1[,6:7]
                  <- sqrt(s.param.1[,6:7])
t(apply(s.param.1,2, function(x) quantile(x, c(.025,.25,.5,.75,.975))))
##
            2.5%
                    25%
                           50%
                                  75% 97.5%
## theta1 58.539 60.668 61.290 62.136 63.551
## theta2 64.097 65.187 65.966 66.521 67.694
## theta3 65.779 67.124 67.719 68.310 69.505
## theta4 59.654 60.640 61.285 61.767 62.598
          58.590 62.809 63.768 65.404 69.947
## sigma2 1.851 2.169 2.395 2.694 3.328
## tau2
           1.825 2.777 3.833 6.389 17.368
                  <- function(arg1) {
r.hat
                  <- dim(arg1)[1]
  n
  m
                  <- dim(arg1)[2]
  В
                  (n/(m-1)) * sum( (apply(arg1,2,mean) - mean(arg1))^2)
                  (1/m) * (1/(n-1)) * sum((arg1 - apply(arg1,2,mean))^2)
  sqrt((n-1)/(n) + B/(W*n))
}
r.hat(sapply(1:10,function(x) s.param[[x]][,4] ))
```

## [1] 1.032

Simulate 200 draws for each of the 10 chains

**Note:** Problems with  $\sigma$  and  $\tau$ , need to review.