## STA 360/602L: Module 5.2

HIERARCHICAL NORMAL MODELS WITH CONSTANT VARIANCE: TWO GROUPS (ILLUSTRATION)

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No pre-recorded video for this module. To be done during discussion session.



## FULL CONDITIONALS RECAP

$$\mu|Y,\delta,\sigma^2 \sim \mathcal{N}(\mu_n,\gamma_n^2), \quad ext{where}$$
  $\gamma_n^2 = rac{1}{rac{1}{\gamma_0^2} + rac{n_m + n_f}{\sigma^2}}$   $\mu_n = \gamma_n^2 \left[rac{\mu_0}{\gamma_0^2} + rac{\sum\limits_{i=1}^{n_m} (y_{i,male} - \delta) + \sum\limits_{i=1}^{n_f} (y_{i,female} + \delta)}{\sigma^2}
ight].$ 

## Full conditionals

$$\delta|Y,\mu,\sigma^2 \sim \mathcal{N}(\delta_n, au_n^2), \quad ext{where}$$
  $au_n^2 = rac{1}{rac{1}{ au_0^2} + rac{n_m + n_f}{\sigma^2}}$   $\delta_n = au_n^2 \left[ rac{\delta_0}{ au_0^2} + rac{\sum\limits_{i=1}^{n_m} (y_{i,male} - \mu) + (-1)\sum\limits_{i=1}^{n_f} (y_{i,female} - \mu)}{\sigma^2} 
ight].$ 

## Full conditionals

$$\sigma^2|Y,\mu,\delta\sim \mathcal{IG}(rac{
u_n}{2},rac{
u_n\sigma_n^2}{2}), ~~ ext{where}$$
  $onumber 
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■ The data we will use in the R package rethinking.

```
#install.packages(c("coda", "devtools", "loo", "dagitty"))
#library(devtools)
#devtools::install_github("rmcelreath/rethinking", ref="Experimental")
#library(rethinking)
data(Howell1)
Howell1[1:15,]
```

```
height
               weight age male
##
## 1 151.765 47.82561 63.0
## 2 139.700 36.48581 63.0
## 3 136.525 31.86484 65.0
## 4 156.845 53.04191 41.0
## 5 145.415 41.27687 51.0
## 6 163.830 62.99259 35.0
## 7 149.225 38.24348 32.0
## 8 168.910 55.47997 27.0
## 9 147.955 34.86988 19.0
## 10 165.100 54.48774 54.0
## 11 154.305 49.89512 47.0
## 12 151.130 41.22017 66.0
## 13 144.780 36.03221 73.0
## 14 149.900 47.70000 20.0
## 15 150.495 33.84930 65.3
```



• For now, focus on data for individuals under age 15.

```
htm <- Howell1$height/100
bmi <- Howell1$weight/(htm^2)</pre>
y_male <- bmi[Howell1$age<15 & Howell1$male==1]</pre>
y_female <- bmi[Howell1$age<15 & Howell1$male==0]</pre>
n_m <- length(y_male)</pre>
n_f <- length(y_female)</pre>
n f
## [1] 84
n_m
## [1] 77
summary(y_male)
     Min. 1st Qu. Median Mean 3rd Qu.
                                             Max.
##
##
     12.07 13.87 14.63 14.84 15.53
                                              18.22
summary(y_female)
     Min. 1st Qu. Median Mean 3rd Qu.
                                               Max.
     9.815 13.559 14.305 14.585 15.712 18.741
```

- We will set the hyper-parameters as:
  - $\mu_0 = 15, \gamma_0 = 5$ ,
  - $\delta_0 = 0, \tau_0 = 3$ ,
  - $\nu_0 = 1, \sigma_0 = 5.$
- Do these values seem reasonable?

```
#priors
mu0 <- 15; gamma02 <- 5^2
delta0 <- 0; tau02 <- 3^2
nu0 <- 1; sigma02 <- 5^2

#starting values
mu <- (mean(y_male) + mean(y_female))/2
delta <- (mean(y_male) - mean(y_female))/2
#no need for starting values for sigma_squared, we can sample it first

MU <- DELTA <- SIGMA2 <- NULL</pre>
```

```
#set seed
set.seed(1234)
#set number of iterations and hurn-in
n iter <- 10000; burn in <- 0.2*n iter
##Gibbs sampler
for (s in 1:(n iter+burn in)) {
#update sigma2
sigma2 < -1/rgamma(1,(nu0 + n_m + n_f)/2,
                    (nu0*sigma02 + sum((y_male-mu-delta)^2) + sum((y_female-mu+delta)^2))/2
#update mu
gamma2n <- 1/(1/gamma02 + (n_m + n_f)/sigma2)
mun <- gamma2n*(mu0/gamma02 + sum(y_male-delta)/sigma2 + sum(y_female+delta)/sigma2)</pre>
mu <- rnorm(1,mun,sqrt(gamma2n))</pre>
#update delta
tau2n <- 1/(1/tau02 + (n_m+n_f)/sigma2)
deltan <- tau2n*(delta0/tau02 + sum(y_male-mu)/sigma2 - sum(y_female-mu)/sigma2)</pre>
delta <- rnorm(1,deltan,sqrt(tau2n))</pre>
#save parameter values
MU <- c(MU,mu); DELTA <- c(DELTA,delta); SIGMA2 <- c(SIGMA2,sigma2)
```

## Posterior summaries

```
#library(coda)
MU.mcmc <- mcmc(MU,start=1)</pre>
summary(MU.mcmc)
##
## Iterations = 1:12000
## Thinning interval = 1
## Number of chains = 1
## Sample size per chain = 12000
##
## 1. Empirical mean and standard deviation for each variable,
##
      plus standard error of the mean:
##
                                        Naive SE Time-series SE
##
             Mean
                              SD
        14.712517
                        0.118765
                                        0.001084
                                                       0.001089
##
##
## 2. Ouantiles for each variable:
##
  2.5%
           25%
                 50%
                       75% 97.5%
## 14.48 14.63 14.71 14.79 14.95
(mean(y_male) + mean(y_female))/2 #compare to data
## [1] 14.7127
```



### Posterior summaries

```
DELTA.mcmc <- mcmc(DELTA,start=1)</pre>
summary(DELTA.mcmc)
##
## Iterations = 1:12000
## Thinning interval = 1
## Number of chains = 1
## Sample size per chain = 12000
##
## 1. Empirical mean and standard deviation for each variable,
     plus standard error of the mean:
##
##
                                       Naive SE Time-series SE
##
                              SD
             Mean
##
         0.127657
                        0.119522
                                       0.001091
                                                      0.001091
##
  2. Quantiles for each variable:
##
##
       2.5%
                 25%
                          50%
                                   75%
                                          97.5%
## -0.10691 0.04791 0.12743 0.20796 0.36407
summary((2*DELTA)) #rescale as difference in group means
##
      Min. 1st Qu. Median
                                  Mean 3rd Qu.
                                                    Max.
## -0.63464 0.09582 0.25487 0.25531 0.41592 1.23660
mean(y_male) - mean(y_female) #compare to data
```

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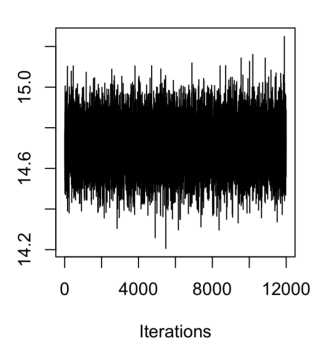
## Posterior summaries

```
SIGMA2.mcmc <- mcmc(SIGMA2,start=1)</pre>
summary(SIGMA2.mcmc)
##
## Iterations = 1:12000
## Thinning interval = 1
## Number of chains = 1
## Sample size per chain = 12000
##
## 1. Empirical mean and standard deviation for each variable,
      plus standard error of the mean:
##
##
##
                                        Naive SE Time-series SE
                              SD
             Mean
                                        0.002352
         2.287927
                        0.257689
##
                                                       0.002352
##
## 2. Quantiles for each variable:
##
  2.5%
           25%
                 50%
                       75% 97.5%
##
## 1.833 2.107 2.272 2.455 2.841
```

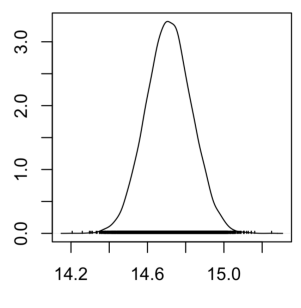


plot(MU.mcmc)

#### Trace of var1

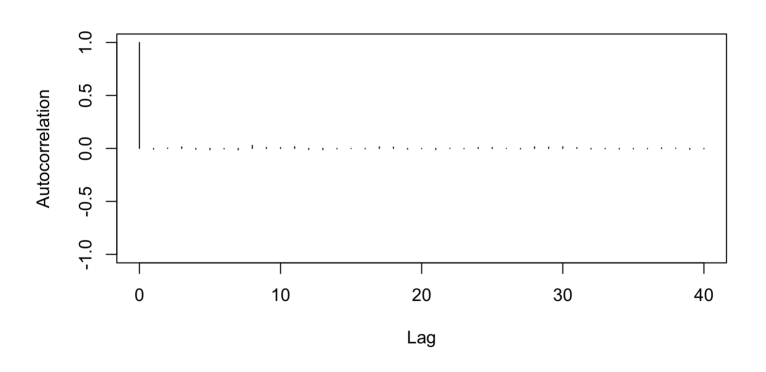


#### **Density of var1**



N = 12000 Bandwidth = 0.01924

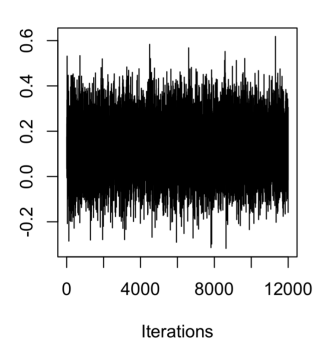
autocorr.plot(MU.mcmc)



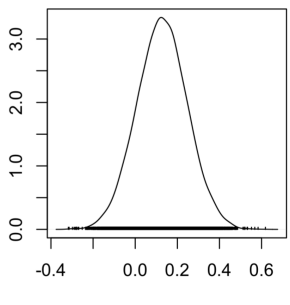


plot(DELTA.mcmc)

#### Trace of var1

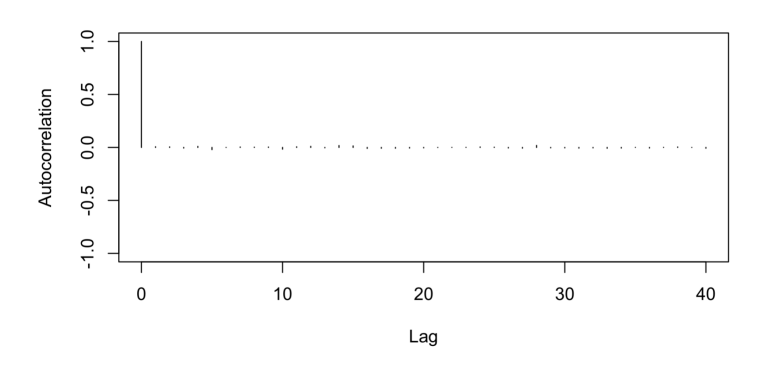


#### **Density of var1**



N = 12000 Bandwidth = 0.01935

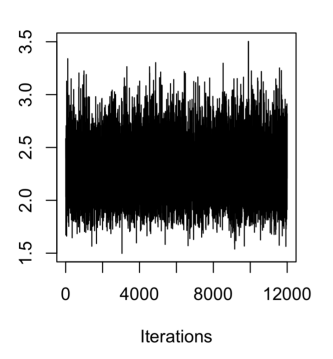
autocorr.plot(DELTA.mcmc)



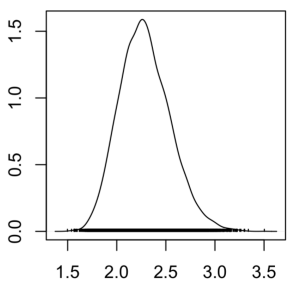


plot(SIGMA2.mcmc)



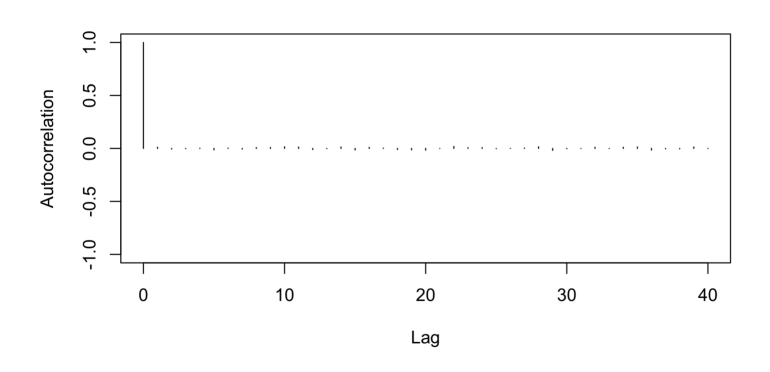


#### **Density of var1**



N = 12000 Bandwidth = 0.04174

autocorr.plot(SIGMA2.mcmc)





- Posterior probability that boys have larger average BMI than girls is 0.86!
- Posterior medians and 95% credible intervals for the group means are actually quite similar to the unpooled gender specific intervals from classical inference (do a t-test to confirm).

```
#mean for boys
quantile((MU+DELTA),probs=c(0.025,0.5,0.975))
      2.5%
##
                 50%
                        97.5%
## 14.50255 14.84146 15.17925
#mean for girls
quantile((MU-DELTA),probs=c(0.025,0.5,0.975))
       2.5%
                 50%
                        97.5%
## 14.26848 14.58276 14.90761
#posterior probability boys have larger BMI than girls
mean(DELTA > 0)
## [1] 0.8571667
```



ullet Let's look at a different sub-population. For older individuals >75, we only have 8 male and 4 female.

```
y_male <- bmi[Howell1$age > 75 & Howell1$male==1]
y_female <- bmi[Howell1$age > 75 & Howell1$male==0]
n_m <- length(y_male)
n_f <- length(y_female)
n_m

## [1] 8

## [1] 4</pre>
```



■ A 95% confidence interval for the difference between genders in BMI (estimated as 0.24) is (-4.20,4.68).

```
mean(y_male) - mean(y_female)

## [1] 0.2408966

t.test(y_male,y_female)

##

## Welch Two Sample t-test

##

## data: y_male and y_female

## t = 0.13801, df = 5.1869, p-value = 0.8954

## alternative hypothesis: true difference in means is not equal to 0

## 95 percent confidence interval:

## -4.197948 4.679741

## sample estimates:

## mean of x mean of y

## 18.06751 17.82662
```



Let's apply the Bayesian model with these priors:

• 
$$\mu_0 = 18, \gamma_0 = 5$$
,

• 
$$\delta_0 = 0, \tau_0 = 3$$
,

• 
$$\nu_0 = 1, \sigma_0 = 5.$$

- The R code for running the sampler is suppressed here. Basically, just rerun the same Gibbs sampler from before on this new data.
- Using the results from the model, the posterior mean is 0.25 with 95% CI (-3.45, 3.88).

```
mean((DELTA*2))

## [1] 0.2493733

quantile((DELTA*2),probs=c(0.025,0.5,0.975))

## 2.5% 50% 97.5%

## -3.4466931 0.2758598 3.8762543
```



- The width of this interval is smaller than that of the 95% confidence interval from before.
- In a way, precision has been improved by borrowing of information across the groups. Of course the prior is important here given the sample sizes.



# WHAT'S NEXT?

MOVE ON TO THE READINGS FOR THE NEXT MODULE!

