# Bayesian Hierarchical/ Multilevel and Latent-Variable (Random-Effects) Modeling

2: Fixed- and random-effects models in meta-analysis

#### **David Draper**

Department of Applied Mathematics and Statistics University of California, Santa Cruz

> draper@ams.ucsc.edu http://www.ams.ucsc.edu/~draper

National University of Ireland, Galway

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# Hierarchical Models for Combining Information

Formulating hierarchical models for quantitative outcomes from scientific context

Case Study: Meta-analysis of effects of aspirin on heart attacks. Table 5.1 (Draper et al., 1993a) gives the number of patients and mortality rate from all causes, for six randomized controlled experiments comparing the use of aspirin and placebo by patients following a heart attack.

Table 5.1. Aspirin meta-analysis data.

	Aspirin		<u>Placebo</u>	
	# of	Mortality	# of	Mortality
Study $(i)$	<b>Patients</b>	Rate (%)	<b>Patients</b>	Rate (%)
UK-1	615	7.97	624	10.74
CDPA	758	5.80	771	8.30
GAMS	317	8.52	309	10.36
UK-2	832	12.26	850	14.82
PARIS	810	10.49	406	12.81
AMIS	2267	10.85	2257	9.70
Total	5599	9.88	5217	10.73

Comparison				
	$y_i = \overline{\text{Diff}}$	$\sqrt{V_i} = SE$		
Study (i)	(%)	of Diff (%)	$Z_i^{\ddagger}$	$p_i^\S$
UK-1	2.77	1.65	1.68	.047
CDPA	2.50	1.31	1.91	.028
GAMS	1.84	2.34	0.79	.216
UK-2	2.56	1.67	1.54	.062
PARIS	2.31	1.98	1.17	.129
AMIS	-1.15	0.90	-1.27	.898
Total	0.86	0.59	1.47	.072

 $<sup>^{\</sup>ddagger}Z_i$  denotes the ratio of the difference in mortality rates over its standard error, assuming a binomial distribution.  $^\S p_i$  is the one-sided p value associated with  $Z_i$ , using the normal approximation.

#### Meta-Analysis

The first five trials are reasonably consistent in showing a (weighted average) **mortality decline** for aspirin patients of 2.3 percentage points, a **20% drop** from the (weighted average) placebo mortality of 11.5% (this difference is **highly clinically significant**).

However, the sixth and largest trial, AMIS, went the other way: an **increase** of 1.2 percentage points in aspirin mortality (a 12% rise from the placebo baseline of 9.7%).

Some relevant questions (Draper, 1995):

- $|\mathbf{Q}_1|$  Why did AMIS get such different results?
- $|\mathbf{Q}_2|$  What should be done next to **reduce the uncertainty** about  $Q_1$ ?
- $\mathbf{Q}_3$  If you were a doctor treating a patient like those eligible for the trials in Table 5.1, what therapy should you employ while answers to  $Q_1$  and  $Q_2$  are sought?

One possible **paraphrase** of  $Q_3$ :  $\mathbf{Q}_4$  How should the information from these six experiments be **combined** to produce a **more informative summary** than those obtained from each experiment by itself?

The discipline of | meta-analysis | is devoted to answering questions like  $Q_4$ .

One leading school of **frequentist meta-analysis** (e.g., Hedges and Olkin, 1985) looks for methods for combining the Z and p values in Table 5.1, an approach that often leads only to an overall p value.

#### **A** Gaussian HM

A more satisfying form of meta-analysis (which has both frequentist and Bayesian versions) builds a hierarchical model (HM) that indicates how to combine information from the mortality differences in the table.

A Gaussian meta-analysis model for the aspirin data, for example (Draper et al., 1993a), might look like

$$egin{array}{ll} \left(\mu,\sigma^2
ight) &\sim & p\left(\mu,\sigma^2
ight) & ext{(prior)} \\ \left( heta_i|\mu,\sigma^2
ight) &\stackrel{ ext{IID}}{\sim} & N\left(\mu,\sigma^2
ight) & ext{(underlying effects)} & ext{(1)} \\ \left(y_i|\theta_i\right) &\stackrel{ ext{indep}}{\sim} & N(\theta_i,V_i) & ext{(data)} \ . \end{array}$$

The bottom level of (1), the **data** level of the HM, says that—because of relevant differences in patient cohorts and treatment protocols—each study has its own **underlying treatment effect**  $\theta_i$ , and the observed mortality differences  $y_i$  are like random draws from a normal distribution with mean  $\theta_i$  and variance  $V_i$  (the normality is reasonable because of the **Central Limit Theorem**, given the large numbers of patients).

In meta-analyses of data like those in Table 5.1 the  $V_i$  are typically taken to be **known** (again because the patient sample sizes are so big),  $V_i = SE_i^2$ , where  $SE_i$  is the standard error of the mortality difference for study i in Table 5.1.

The middle level of the HM is where you would bring in the **study-level covariates**, if you have any, to try to explain why the studies differ in their underlying effects.

Here there are no study-level covariates, so the middle level of (1) is equivalent to a **Gaussian regression with no** predictor variables.

# A Gaussian HM (continued)

Why the "error" distribution should be **Gaussian** at this level of the HM is not clear—it's a **conventional** option, not a choice that's automatically scientifically reasonable (in fact I'll challenge it later).

 $\sigma^2$  in this model represents **study-level heterogeneity**.

The top level of (1) is where the **prior** distribution on the regression parameters from the middle level is specified.

Here, with only an intercept term in the regression model, a popular **conventional choice** is the normal/scaled-inverse- $\chi^2$  prior we looked at earlier in our first Gaussian case study.

Fixed effects and random effects. If  $\sigma^2$  were known somehow to be 0, all of the  $\theta_i$  would have to be equal deterministically to a common value  $\mu$ , yielding a simpler model:  $(y_i|\mu) \stackrel{\mathsf{indep}}{\sim} N(\mu, V_i)$ ,  $\mu \sim p(\mu)$ .

Meta-analysts call this a **fixed-effects** model, and refer to model (1) as a **random-effects** model.

When  $\sigma^2$  is not assumed to be 0, with this terminology the  $\theta_i$  are called **random effects** (this parallels the use of this term in the **random-effects Poisson regression** case study).

# Approximate Fitting of Gaussian Hierarchical Models: Maximum Likelihood and Empirical Bayes

Fitting HM (1). Some algebra based on model (1) yields that the conditional distributions of the study-level effects  $\theta_i$  given the data and the parameters  $(\mu, \sigma^2)$ , have a **simple** and revealing form (I'll show this later):

$$(\theta_i|y_i,\mu,\sigma^2) \stackrel{\mathsf{indep}}{\sim} N[\theta_i^*,V_i(1-B_i)],$$
 (2)

with 
$$\theta_i^* = (1 - B_i) y_i + B_i \mu$$
 and  $B_i = \frac{V_i}{V_i + \sigma^2}$ . (3)

In other words, the conditional mean of the effect for study i given  $y_i, \mu$ , and  $\sigma^2$  is a **weighted average** of the sample mean for that study,  $y_i$ , and the overall mean  $\mu$ .

The weights are given by the so-called **shrinkage factors**  $B_i$  (e.g., Draper et al., 1993a), which in turn depend on how the variability  $V_i$  within study i compares to the **between-study** variability  $\sigma^2$ : the more accurately  $y_i$  estimates  $\theta_i$ , the more weight the "local" estimate  $y_i$  gets in the weighted average.

The term **shrinkage** refers to the fact that, with this approach, unusually high or low individual studies are **drawn back** or "shrunken" toward the overall mean  $\mu$  when making the calculation  $(1 - B_i) y_i + B_i \mu$ .

Note that  $\theta_i^*$  uses data from all the studies to estimate the effect for study i—this is referred to as **borrowing strength** in the estimation process.

Closed-form expressions for  $p(\mu|y)$  and  $p(\theta_i|y)$  with  $y = (y_1, \dots, y_k), k = 6$  are not available even with a normal- $\chi^{-2}$  prior for  $(\mu, \sigma^2)$ ; MCMC is needed (see below).

# Maximum Likelihood and Empirical Bayes

In the meantime | maximum likelihood | calculations provide some idea of what to expect: the likelihood function based on model (1) is

$$l(\mu, \sigma^{2}|y) = c \left[ \prod_{i=1}^{k} \frac{1}{\sqrt{V_{i} + \sigma^{2}}} \right] \exp \left[ -\frac{1}{2} \sum_{i=1}^{k} \frac{(y_{i} - \mu)^{2}}{V_{i} + \sigma^{2}} \right].$$
 (4)

The maximum likelihood estimates (MLEs)  $(\hat{\mu}, \hat{\sigma}^2)$  then turn out to be the **iterative** solutions to the following equations:

$$\hat{\mu} = \frac{\sum_{i=1}^{k} \hat{W}_{i} y_{i}}{\sum_{i=1}^{k} \hat{W}_{i}} \quad \text{and} \quad \hat{\sigma}^{2} = \frac{\sum_{i=1}^{k} \hat{W}_{i}^{2} \left[ (y_{i} - \hat{\mu})^{2} - V_{i} \right]}{\sum_{i=1}^{k} \hat{W}_{i}^{2}}, \quad (5)$$

$$\text{where} \quad \hat{W}_{i} = \frac{1}{V_{i} + \hat{\sigma}^{2}}. \quad (6)$$

where 
$$\hat{W}_i = \frac{1}{V_i + \hat{\sigma}^2}$$
. (6)

Start with  $\hat{\sigma}^2 = 0$  and iterate (5–6) to convergence (if  $\hat{\sigma}^2$  converges to a negative value,  $\hat{\sigma}^2 = 0$  is the MLE); the MLEs of the  $\theta_i$  are then given by

$$\hat{\theta}_i = (1 - \hat{B}_i) \ y_i + \hat{B}_i \hat{\mu} \quad \text{where} \quad \hat{B}_i = \frac{V_i}{V_i + \hat{\sigma}^2}.$$
 (7)

These are called **empirical Bayes** (EB) estimates of the study-level effects, because it turns out that this analysis approximates a fully Bayesian solution by (in effect) using the data to **estimate** the prior specifications for  $\mu$  and  $\sigma^2$ .

**Large-sample** (mainly meaning large k) approximations to the (frequentist) distributions of the MLEs are given by

$$\widehat{\mu} \sim N\left(\mu, \left[\sum_{i=1}^{k} \frac{1}{V_i + \widehat{\sigma}^2}\right]^{-1}\right)$$
 and  $\widehat{\theta}_i \sim N\left[\theta_i, V_i(1 - \widehat{B}_i)\right]$ . (8)

# MLEB (continued)

NB The variances in (8) don't account fully for the uncertainty in  $\sigma^2$  and therefore underestimate the actual sampling variances for small k (adjustments are available; see, e.g., Morris (1983, 1988)).

MLEB estimation can be implemented simply in about 15 lines of R code (Table 5.2).

Table 5.2. R program to perform MLEB calculations.

```
mleb <- function( y, V, m ) {
    sigma2 <- 0.0
    for ( i in 1:m ) {
        W <- 1.0 / ( V + sigma2 )
        theta <- sum( W * y ) / sum( W )
        sigma2 <- sum( W^2 * ( ( y - theta )^2 - V ) ) / sum( W^2 )
        B <- V / ( V + sigma2 )
        effects <- ( 1 - B ) * y + B * theta
        se.theta <- 1.0 / sqrt( sum( 1.0 / ( V + sigma2 ) ) )
        se.effects <- sqrt( V * ( 1.0 - B ) )
        print( c( i, theta, se.theta, sigma2 ) )
        print( cbind( W, ( W / sum( W ) ), B, y, effects, se.effects ) )
    }
}</pre>
```

With the aspirin data it takes **18 iterations** (less than 0.1 second on a 400MHz UltraSPARC Unix machine) to get convergence to **4-digit accuracy**, leading to the summaries in Table 5.3 and the following estimates (standard errors in parentheses):

$$\hat{\mu} = 1.45 \ (0.809), \quad \hat{\sigma}^2 = 1.53.$$

Table 5.3. Maximum likelihood empirical Bayes meta-analysis of the aspirin data.

study(i)	$\widehat{W}_i$	normalized $\hat{W}_i$	$\widehat{B}_i$	$y_i$	$\widehat{ heta}_i$	$\widehat{SE}(\widehat{\theta}_i)$
1	0.235	0.154	0.640	2.77	1.92	0.990
2	0.308	0.202	0.529	2.50	1.94	0.899
3	0.143	0.0934	0.782	1.84	1.53	1.09
4	0.232	0.151	0.646	2.56	1.84	0.994
5	0.183	0.120	0.719	2.31	1.69	1.05
6	0.427	0.280	0.346	-1.15	-0.252	0.728

#### **Aspirin Meta-Analysis: Conclusions**

Note that (1) AMIS gets **much less weight** (normalized  $\hat{W}_i$ ) than would have been expected given its small  $V_i$ ; (2) the **shrinkage factors**  $(\hat{B}_i)$  are considerable, with AMIS shrunk almost all the way into positive territory (see Figure 5.1); (3) there is **considerable study-level heterogeneity**  $(\hat{\sigma} \doteq 1.24)$  percentage points of mortality); and (4) the standard errors of the effects are by and large smaller than the  $\sqrt{V_i}$  (from the **borrowing of strength**) but are still considerable.

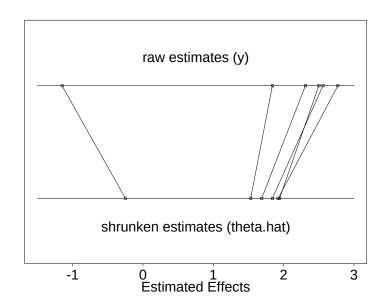


Figure 5.1. Shrinkage plot for the aspirin MLEB meta-analysis.

The 95% interval estimate of  $\mu$ , the overall underlying effect of aspirin on mortality, from this approach comes out

$$\hat{\mu} \pm 1.96 \cdot \widehat{SE}(\hat{\mu}) \doteq (-0.140, 3.03),$$

which if interpreted Bayesianly gives

$$P( ext{aspirin reduces mortality}| ext{data}) \doteq 1 - \Phi(rac{0-1.45}{0.809}) = extbf{0.96}$$
 ,

where  $\Phi$  is the **standard normal CDF**.

Thus although the interval includes 0, so that in a frequentist sense the effect is not statistically significant, in fact from a Bayesian point of view the evidence is running strongly in favor of aspirin's usefulness.

#### **MCMC Details**

In many cases (as with this example) empirical Bayes methods have the advantage of yielding **closed-form solutions**, but I view them at best as approximations to fully Bayesian analyses—which can in any case be carried out with MCMC—so I won't have any more to say about EB methods here (see Carlin and Louis, 1996, for more on this topic).

MCMC details. First let's derive that likelihood function I mentioned on page 7: the model, once again, is

$$\begin{pmatrix} (\mu, \sigma^2) & \sim & p(\mu, \sigma^2) & \text{(prior)} \\ (\theta_i | \mu, \sigma^2) & \overset{\text{IID}}{\sim} & N(\mu, \sigma^2) & \text{(underlying effects)} & (9) \\ (y_i | \theta_i) & \overset{\text{indep}}{\sim} & N(\theta_i, V_i) & \text{(data)} . \end{cases}$$

The **parameters** we're interested in here are  $(\mu, \sigma^2)$ ; **Bayes's** Theorem gives (as usual)

$$p(\mu, \sigma^2 | y) = c p(\mu, \sigma^2) p(y | \mu, \sigma^2),$$
 (10)

so let's look at the **sampling distribution** for a single  $y_i$ :

$$p(y_{i}|\mu,\sigma^{2}) = \int_{-\infty}^{\infty} p(y_{i},\theta_{i}|\mu,\sigma^{2}) d\theta_{i}$$

$$= \int_{-\infty}^{\infty} p(y_{i}|\theta_{i},\mu,\sigma^{2}) p(\theta_{i}|\mu,\sigma^{2}) d\theta_{i} \qquad (11)$$

$$= \int_{-\infty}^{\infty} p(y_{i}|\theta_{i}) p(\theta_{i}|\mu,\sigma^{2}) d\theta_{i}$$

(what we're doing here is **integrating out** the **random effect**  $\theta_i$ ).

Now  $p(y_i|\theta_i)$  is **normal** in this model, and  $p(\theta_i|\mu,\sigma^2)$  is **also normal**; you could put in the **normal densities** and **grind away** at the **algebra** and **integration**, but there's a **better way**: the last line of (11) is a **mixture representation**, and a **normal mixture of normals is normal**, so I know that  $p(y_i|\mu,\sigma^2)$  is **normal**, and the only questions are, what are its **mean** and **variance**?

#### Adam and Eve

These questions can be answered with **little difficulty** via the **Double Expectation Theorem**, which has **two parts** that are so **central** to **Bayesian calculations** that **Carl Morris** refers to them as **Adam** and **Eve**: for any two random variables X and Y,

$$E(Y) = E_X [E(Y|X)]$$
 (Adam)  
 $V(Y) = E_X [V(Y|X)] + V_X [E(Y|X)]$  (Eve), (12)

in which  $E_X$  and  $V_X$  refer to **expectation** and **variance** with respect to the **distribution** of X.

If there's **additional conditioning** going on, you just need to remember to **include it** in all the **relevant places**: for any three random variables X, Y and Z,

$$E(Y|Z) = E_{(X|Z)} [E(Y|X,Z)] V(Y|Z) = E_{(X|Z)} [V(Y|X,Z)] + V_{(X|Z)} [E(Y|X,Z)],$$
and **so on**. (13)

The application here is in two parts (Adam and Eve):

$$E(y_i|\mu,\sigma^2) = E_{(\theta_i|\mu,\sigma^2)} \left[ E(y_i|\mu,\sigma^2,\theta_i) \right]$$

$$= E_{(\theta_i|\mu,\sigma^2)} \left[ E(y_i|\theta_i) \right]$$

$$= E_{(\theta_i|\mu,\sigma^2)} \left[ \theta_i \right]$$

$$= \mu, \text{ and}$$

$$V(y_{i}|\mu,\sigma^{2}) = E_{(\theta_{i}|\mu,\sigma^{2})} \left[ V(y_{i}|\mu,\sigma^{2},\theta_{i}) \right] + V_{(\theta_{i}|\mu,\sigma^{2})} \left[ E(y_{i}|\mu,\sigma^{2},\theta_{i}) \right]$$

$$= E_{(\theta_{i}|\mu,\sigma^{2})} \left[ V(y_{i}|\theta_{i}) \right] + V_{(\theta_{i}|\mu,\sigma^{2})} \left[ E(y_{i}|\theta_{i}) \right]$$

$$= E_{(\theta_{i}|\mu,\sigma^{2})} \left[ V_{i} \right] + V_{(\theta_{i}|\mu,\sigma^{2})} \left[ \theta_{i} \right]$$

$$= V_{i} + \sigma^{2}. \tag{14}$$

#### **Direct Use of Gibbs Sampling**

So (a)  $(y_i|\mu,\sigma^2) \sim N(\mu,V_i+\sigma^2)$ , (b) by **inspection** of the **form** of the **model**, the  $y_i$  are **independent** given  $(\mu,\sigma^2)$ , so

$$l(\mu, \sigma^{2}|y) = c p(y|\mu, \sigma^{2}) = c \prod_{i=1}^{k} p(y_{i}|\mu, \sigma^{2})$$

$$= c \left[ \prod_{i=1}^{k} \frac{1}{\sqrt{V_{i} + \sigma^{2}}} \right] \exp \left[ -\frac{1}{2} \sum_{i=1}^{k} \frac{(y_{i} - \mu)^{2}}{V_{i} + \sigma^{2}} \right],$$
as **desired**. (15)

MCMC: how best to sample from the posterior?

All MCMC (with a parameter space of fixed dimension) is one special case or another of the Metropolis-Hastings algorithm, but (as usual) we have a number of possibilities: generic (e.g., random-walk) Metropolis? Metropolis mixed with Gibbs steps? All Gibbs? With or without auxiliary (e.g., latent) variables? ...

First let's try **direct Gibbs**, for which we would need the **full conditionals**:

$$p(\mu|\sigma^2, y) = c p(\mu, \sigma^2, y)$$

$$= c p(\mu, \sigma^2) p(y|\mu, \sigma^2).$$
(16)

By virtue of **integrating out** the **random effects** above, we have  $p(y|\mu,\sigma^2)$  as a **product** of **independent univariate Gaussians**; what shall we take for the **prior**  $p(\mu,\sigma^2)$ , given that there's no **conjugate choice**?

Even with somewhat informative priors on a vector of parameters, for simplicity people often assume independence of the components — in this case,  $p(\mu, \sigma^2) = p(\mu) \, p(\sigma^2)$  — on the ground that whatever correlation the parameters should have in the posterior will be learned via the likelihood function; let's make this simplifying assumption; then

$$p(\mu|\sigma^2, y) = c p(\mu) p(\sigma^2) p(y|\mu, \sigma^2) = c p(\mu) p(y|\mu, \sigma^2).$$
 (17)

#### **Direct Gibbs; Latent Gibbs**

Now the **product of two Gaussians is Gaussian**, so if we take the **prior** for  $\mu$  to be **Gaussian** we'll have a **Gaussian** full conditional for  $\mu$  that'll be easy to sample from; what about  $\sigma^2$ ?

$$p(\sigma^{2}|\mu,y) = c p(\mu,\sigma^{2},y)$$

$$= c p(\mu,\sigma^{2}) p(y|\mu,\sigma^{2})$$

$$= c p(\mu) p(\sigma^{2}) p(y|\mu,\sigma^{2})$$

$$= c p(\sigma^{2}) p(y|\mu,\sigma^{2}).$$
(18)

Here we run into **trouble**: when considered as a **function** of  $\sigma^2$  for fixed  $\mu$  and y,  $p(y|\mu,\sigma^2)$  is **not recognizable** as a member of a **standard parametic family** (because the  $y_i$  (given  $\mu$  and  $\sigma^2$ ) are **independent** but **not identically distributed**); we could choose, e.g., a  $\chi^{-2}$  prior on  $\sigma^2$  and use **rejection sampling** to sample from the resulting **non-standard full conditional**, but that would not be especially **pleasant**.

So instead let's use a **trick** that's generally helpful in **random-effects** models: treat the **(latent)** random effects as **auxiliary** variables to be sampled along with  $(\mu, \sigma^2)$ .

In other words, letting  $\theta = (\theta_1, \dots, \theta_k)$ , we're going to sample from the **augmented posterior**  $p(\mu, \sigma^2, \theta|y)$ ; the hope is that this will have **completely tractable full conditionals**; let's see.

$$p(\mu|\sigma^2, \theta, y) = c p(\mu, \sigma^2, \theta, y)$$

$$= c p(\mu, \sigma^2) p(\theta|\mu, \sigma^2) p(y|\theta, \mu, \sigma^2)$$
(19)

Notice how naturally this **factorization** matches the **hierarchical character** of (9), which starts at the **top** with a model for  $(\mu, \sigma^2)$ , and then builds a **model** for  $(\theta|\mu, \sigma^2)$ , and then at the **bottom** there's a model for  $p(y|\theta,\mu,\sigma^2)$ , which — by virtue of the **hierarchical** structure — can be **simplified** to  $p(y|\theta)$ .

# **Latent Gibbs (continued)**

Since (a) we're **assuming** that  $p(\mu, \sigma^2) = p(\mu) p(\sigma^2)$  and (b)  $p(y|\theta)$  **doesn't involve**  $\mu$ , the **full conditional** for  $\mu$  becomes

$$p(\mu|\sigma^2, \theta, y) = c p(\mu) p(\theta|\mu, \sigma^2); \tag{20}$$

with a **Gaussian** prior on  $\mu$  this will be **Gaussian**; how about  $\sigma^2$ ?

$$p(\sigma^{2}|\mu,\theta,y) = c p(\mu,\sigma^{2},\theta,y)$$

$$= c p(\mu,\sigma^{2}) p(\theta|\mu,\sigma^{2}) p(y|\theta,\mu,\sigma^{2})$$

$$= c p(\mu) p(\sigma^{2}) p(\theta|\mu,\sigma^{2}) p(y|\theta)$$

$$= c p(\sigma^{2}) p(\theta|\mu,\sigma^{2}).$$
(21)

Here's another **trick**: instead of **slogging** through the **details**, try to **recognize** situations in which you already know the **conjugate updating**, and just use what you **already know**.

For example, in this calculation  $(\theta|\mu,\sigma^2)$  is **Gaussian** with **known**  $\mu$  and **unknown**  $\sigma^2$ , and we know the **conjugate prior** for  $\sigma^2$  in that model —  $\chi^{-2}$  — so with that **prior choice** the **full conditional** for  $\sigma^2$  will also be  $\chi^{-2}$ ; how about  $\theta$ ?

$$p(\theta|\mu, \sigma^{2}, y) = c p(\mu, \sigma^{2}, \theta, y)$$

$$= c p(\mu, \sigma^{2}) p(\theta|\mu, \sigma^{2}) p(y|\theta, \mu, \sigma^{2})$$

$$= c p(\theta|\mu, \sigma^{2}) p(y|\theta).$$
(22)

Here  $p(\theta|\mu, \sigma^2)$  and  $p(y|\theta)$  are both **Gaussian**, so the **full** conditional for  $\theta$  — the **product** — will also be **Gaussian**.

Thus using the **latent Gibbs** approach in this **random-effects** model, all of the **full conditionals** have familiar forms; this approach will **work smoothly**; we just need to work out the **details**.

(I recommend this as a **basic Gibbs strategy**: in the first step make a **sketchy pass** through the **full conditionals** without working out all of the details, to ensure that everything **works fine**, and then go back and **fill in the details**.)

#### **Details**

(1) Full conditional for  $\mu$ :

$$p(\mu|\sigma^2, \theta, y) = c p(\mu) p(\theta|\mu, \sigma^2). \tag{23}$$

In this **calculation** (a)  $\sigma^2$  is known and (b) the **latent vector**  $\theta = (\theta_1, \dots, \theta_k)$  acts like the **data vector**  $y = (y_1, \dots, y_n)$  in the model  $\mu \sim N(\mu_0, \sigma_{\mu_0}^2), (y_i|\mu) \stackrel{\text{IID}}{\sim} N(\mu, \sigma^2)$   $(i = 1, \dots, n)$ , so we already know the **answer**:  $(\mu|\sigma^2, \theta, y) \sim N(\mu_k, \sigma_k^2)$ , where

$$\mu_k = \frac{k_0 \mu_0 + k \overline{\theta}}{k_0 + k} \quad \text{and} \quad \sigma_k^2 = \frac{\sigma^2}{k_0 + k}, \tag{24}$$

and in which the **prior sample size** is  $k_0 = \frac{\sigma^2}{\sigma_{\mu_0}^2}$  and

$$\bar{\theta} = \frac{1}{k} \sum_{i=1}^{k} \theta_i.$$

(2) Full conditional for  $\sigma^2$ :

$$p(\sigma^2|\mu,\theta,y) = c p(\sigma^2) p(\theta|\mu,\sigma^2). \tag{25}$$

In **parallel** with the situation with  $\mu$ , in this **calculation** (a)  $\mu$  is known and (b) the **latent vector**  $\theta = (\theta_1, \dots, \theta_k)$  acts like the **data vector**  $y = (y_1, \dots, y_n)$  in the model  $\sigma^2 \sim \chi^{-2}(\nu_0, \sigma_{\sigma^0}^2), (y_i | \sigma^2) \stackrel{\text{IID}}{\sim} N(\mu, \sigma^2) \ (i = 1, \dots, n)$ , so we already know the **answer**:  $(\sigma^2 | \mu, \theta, y) \sim \chi^{-2}(\nu_k, \sigma_k^2)$ , where

$$\nu_k = \nu_0 + k \quad \text{and} \quad \sigma_k^2 = \frac{\nu_0 \, \sigma_{\sigma 0}^2 + k \, v}{\nu_0 + k},$$
in which  $v = \frac{1}{k} \sum_{i=1}^k (\theta_i - \mu)^2$ .

# **Details** (continued)

(3) Full conditional for  $\theta$ :

$$p(\theta|\mu,\sigma^2,y) = c p(\theta|\mu,\sigma^2) p(y|\theta)$$
$$= c \prod_{i=1}^k p(\theta_i|\mu,\sigma^2) p(y_i|\theta_i). \tag{27}$$

Now  $(\theta_i|\mu,\sigma^2) \sim N(\mu,\sigma^2)$  and  $(y_i|\theta_i) \sim N(\theta_i,V_i)$  (with  $V_i$  known), so this is just our **old friend** 

{Gaussian likelihood (for  $y_i$ ) with unknown mean  $\theta_i$  and known variance  $V_i$  + Gaussian prior for  $\theta_i$  with hyper-parameters  $\mu$  and  $\sigma^2$ };

the **(un-normalized) product**  $p(\theta_i|\mu,\sigma^2) \, p(y_i|\theta_i)$  is just the **posterior** for  $\theta_i$ , and the **answer** is therefore the **same** as it was in the **full conditional** for  $\mu$ :

$$(\theta_i|\mu,\sigma^2,y)\sim N(\theta_i^*,\sigma_i^2)$$
, with

$$\theta_{i}^{*} = \frac{\frac{1}{\sigma^{2}}\mu + \frac{1}{V_{i}}y_{i}}{\frac{1}{\sigma^{2}} + \frac{1}{V_{i}}} = \frac{V_{i}\mu + \sigma^{2}y_{i}}{V_{i} + \sigma^{2}} = B_{i}\mu + (1 - B_{i})y_{i} \quad \text{and}$$

$$\sigma_{i}^{2} = \frac{1}{\frac{1}{\sigma^{2}} + \frac{1}{V_{i}}} = \frac{V_{i}\sigma^{2}}{V_{i} + \sigma^{2}} = V_{i}(1 - B_{i}), \tag{28}$$

in which  $B_i = \frac{V_i}{V_i + \sigma^2}$  is the **shrinkage factor** for study i (this is the **demonstration** of equations (2) and (3) earlier).

Thus  $(\theta|\mu, \sigma^2, y) \sim N_k(\theta^*, \Sigma)$  with  $\theta^* = (\theta_1^*, \dots, \theta_k^*)$  and  $\Sigma = \text{diag}(\sigma_i^2)$ , and **one scan** of the **Gibbs sampler** can be described as follows:

- (a) draw  $\mu$  from  $p(\mu|\sigma^2, \theta, y)$ , obtaining  $\mu_*$ ;
- (b) draw  $\sigma^2$  from  $p(\sigma^2|\mu_*,\theta,y)$ , obtaining  $\sigma^2_*$ ; and
- (c) draw  $\theta$  from  $p(\theta|\mu_*, \sigma_*^2, y)$ , either univariately on the  $\theta_i$  (one by one) or multivariately on  $\theta$  all at once.

#### R Code

```
meta.analysis.gibbs <- function( mu.0, sigma2.mu.0, nu.0, sigma2.sigma.0,
  mu.initial, sigma2.initial, theta.initial, y, V, M, B) {
  k <- length( y )
  mu \leftarrow rep(0, M + B + 1)
  sigma2 < - rep(0, M + B + 1)
  theta \leftarrow matrix(0, M + B + 1, k)
  mu[ 1 ] <- mu.initial</pre>
  sigma2[ 1 ] <- sigma2.initial</pre>
  theta[ 1, ] <- theta.initial
  for ( m in 2:( M + B + 1 ) ) {
    mu[ m ] <- mu.full.conditional( mu.0, sigma2.mu.0, sigma2[ m - 1 ],</pre>
      theta[ m - 1, ], y )
    sigma2[ m ] <- sigma2.full.conditional( nu.0, sigma2.sigma.0,</pre>
      mu[ m ], theta[ m - 1, ], y )
    theta[ m, ] <- theta.full.conditional( mu[ m ], sigma2[ m ], y, V )
    if ( m %% 1000 == 0 ) print( m )
  }
  return(cbind(mu, sigma2, theta))
}
mu.full.conditional <- function( mu.0, sigma2.mu.0, sigma2.current,
  theta.current, y ) {
  k <- length(y)
  k.0 <- sigma2.current / sigma2.mu.0
  theta.bar <- mean( theta.current )</pre>
```

# R Code (continued)

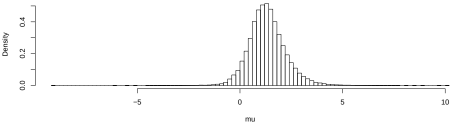
```
mu.k \leftarrow (k.0 * mu.0 + k * theta.bar) / (k.0 + k)
  sigma2.k <- sigma2.current / ( k.0 + k )</pre>
  mu.star <- rnorm( n = 1, mean = mu.k, sd = sqrt( sigma2.k ) )</pre>
  return( mu.star )
}
sigma2.full.conditional <- function( nu.0, sigma2.sigma.0,</pre>
  mu.current, theta.current, y ) {
  k <- length( y )
  nu.k \leftarrow nu.0 + k
  v <- mean( ( theta.current - mu.current )^2 )</pre>
  sigma2.k \leftarrow (nu.0 * sigma2.sigma.0 + k * v) / (nu.0 + k)
  sigma2.star <- rsichi2( 1, nu.k, sigma2.k )</pre>
  return( sigma2.star )
}
rsichi2 <- function( n, nu, sigma2 ) {
  sigma2.star <- 1 / rgamma( n, shape = nu / 2,</pre>
    rate = nu * sigma2 / 2)
  return( sigma2.star )
}
theta.full.conditional <- function( mu.current, sigma2.current, y, V ) {
  k <- length( y )
  theta.star <- ( V * mu.current + sigma2.current * y ) /
    ( V + sigma2.current )
```

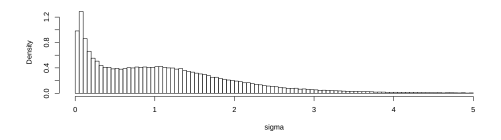
# R Code (continued)

```
sigma2.star <- V * sigma2.current / ( V + sigma2.current )</pre>
  theta.sim <- rnorm( n = k, mean = theta.star,
    sd = sqrt( sigma2.star ) )
  return( theta.sim )
}
mu.0 < -0.0
sigma2.mu.0 <- 100^2
nu.0 <- 0.001
sigma2.sigma.0 <- 1.53
mu.initial <- 1.45
sigma2.initial <- 1.53
theta.initial \leftarrow c( 1.92, 1.94, 1.53, 1.84, 1.69, -0.252 )
y \leftarrow c(2.77, 2.50, 1.84, 2.56, 2.32, -1.15)
V \leftarrow c(1.65, 1.31, 2.34, 1.67, 1.98, 0.90)^2
M <- 100000
B <- 1000
mcmc.data.set <- meta.analysis.gibbs( mu.0, sigma2.mu.0, nu.0,
  sigma2.sigma.0, mu.initial, sigma2.initial, theta.initial,
  y, V, M, B)
% took 47 seconds
mcmc.data.set <- cbind( mcmc.data.set[ , 1:2 ],</pre>
  sqrt( mcmc.data.set[ , 2 ] ), mcmc.data.set[ , 3:8 ] )
```

# R Code (continued)

```
apply( mcmc.data.set[ 1001:101001, ], 2, mean )
        mu
               sigma2
1.33013835 2.24106295 1.12196766 1.68639681 1.67526967 1.38514567 1.62389213
1.51615795 0.09356775
apply( mcmc.data.set[ 1001:101001, ], 2, sd )
             sigma2
       mu
0.9042468 4.4707971 0.9910910 1.1576621 1.0311309 1.2381000 1.1391841
1.1917662 0.9944885
mu.star <- mcmc.data.set[ 1001:101001, 1 ]</pre>
sum( mu.star > 0 ) / length( mu.star )
[1] 0.9484605
sigma.star <- mcmc.data.set[ 1001:101001, 3 ]</pre>
par(mfrow = c(2, 1))
hist( mu.star, nclass = 100, main = '', probability = T,
  xlab = 'mu' )
hist( sigma.star[ sigma.star < 5 ], nclass = 100, main = '',</pre>
  probability = T, xlab = 'sigma' )
```





#### WinBUGS Analysis of Aspirin Data

Aspirin meta-analysis revisited. I create three files for WinBUGS: a model file, a data file, and an initial values file (I'm using the most recent release, 1.4.1, of WinBUGS).

The (first) model file for the aspirin data:

```
mu ~ dnorm( 0.0, 1.0E-6 )
tau.theta ~ dgamma( 1.0E-3, 1.0E-3 )

for ( i in 1:k ) {
   theta[ i ] ~ dnorm( mu, tau.theta )
   y[ i ] ~ dnorm( theta[ i ], tau.y[ i ] )
}

sigma.theta <- 1.0 / sqrt( tau.theta )
}</pre>
```

#### WinBUGS Analysis of Aspirin Data

Here  $\mu$  plays the role of  $\theta$  in model (10) above to avoid using the name theta twice for two different purposes in the WinBUGS program.

In specifying a normal distribution WinBUGS works not with a standard deviation (SD) or a variance but with a precision—the reciprocal of the variance—so that the  $N(\mu,\sigma^2)$  distribution is specified by dnorm( mu, tau ) with  $\tau=\frac{1}{\sigma^2}$ .

Then the **SD** has to be computed as a derived quantity  $(\sigma = \frac{1}{\sqrt{\tau}})$  which is written above as sigma.theta <- 1.0 / sqrt( tau.theta )

If—before the aspirin experiments were performed—I'm relatively **ignorant** about the quantities  $\theta$  ( $\mu$ ) and  $\sigma$  in model (10), or equivalently  $\mu$  and  $\tau = \frac{1}{\sigma^2}$ , I can build a **diffuse** or flat prior for both quantities that expresses this relative ignorance.

Since  $\mu$  lives on  $(-\infty,\infty)$  a convenient choice for a flat prior for it is a **normal** distribution with mean (say) 0 and very small precision: mu  $\sim$  dnorm( 0.0, 1.0E-6 )

For tau.theta, which lives on  $(0,\infty)$ , I want something that's flat throughout (almost) all of that range; a convenient choice (to get an **initial idea** of where the posterior distribution for sigma.theta is **concentrated**) is a **gamma** distribution with small positive values of both of its parameters.

This is the  $\Gamma(\epsilon,\epsilon)$  distribution for some **small**  $\epsilon>0$  like 0.001: tau.theta  $\sim$  dgamma( 1.0E-3, 1.0E-3 )

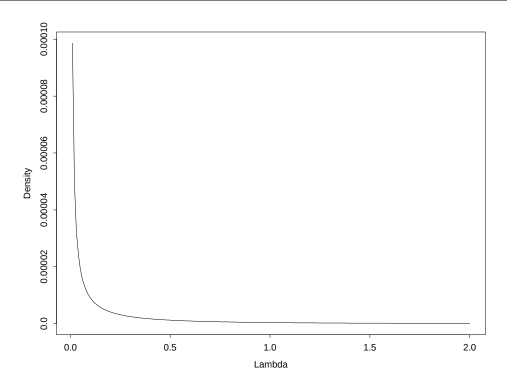


Figure 3.1. The  $\Gamma(0.001, 0.001)$  distribution.

The data file in the aspirin meta-analysis is

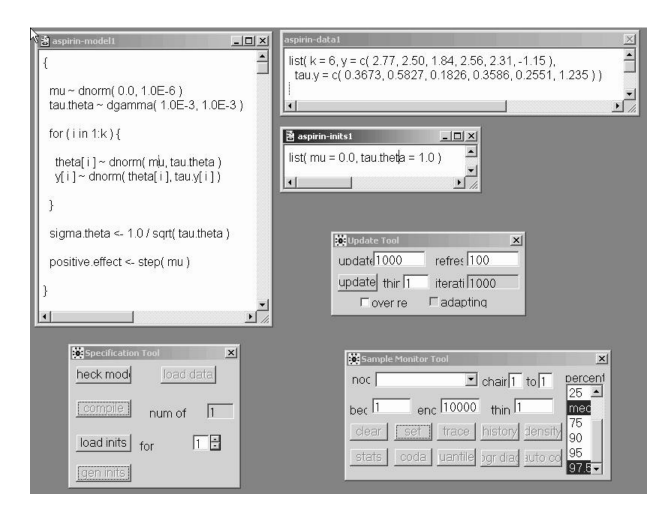
```
list( k = 6, y = c(2.77, 2.50, 1.84, 2.56, 2.31, -1.15), tau.<math>y = c(0.3673, 0.5827, 0.1826, 0.3586, 0.2551, 1.235)
```

Here, e.g., tau.y[1] =  $\frac{1}{1.65^2} \doteq 0.3673$ , where 1.65 is the standard error of the difference y[1] for experiment 1 in Table 2.1 on p. 20.

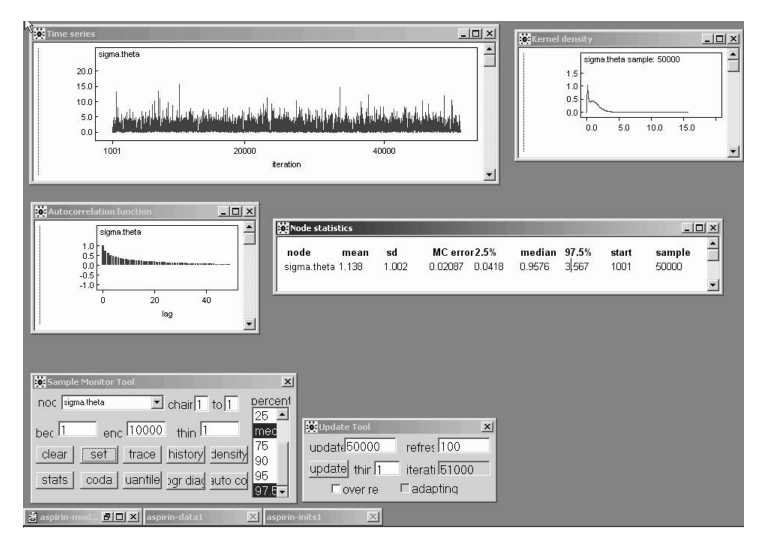
Finally, the initial values file in the aspirin meta-analysis is

list( 
$$mu = 0.0$$
,  $tau.theta = 1.0$ )

In a simple example like this there's no harm in starting the Markov chain off in a **generic** location: since  $\mu$  and  $\tau_{\theta}$  live on  $(-\infty,\infty)$  and  $(0,\infty)$ , convenient generic choices for their starting values are 0 and 1, respectively (more care may be required in models with **more complex** random-effects structure).



- I (1) get a Specification Tool from the Model menu, (2) click on the **model** window and click check model, (3) click on the **data** window and click load data and compile, (4) click on the **initial values** window and click load inits, and (5) click gen inits (because the random effects  $\theta_i$  were uninitialized in the inits file); I'm now ready to do some MCMC sampling.
- I (6) get an Update Tool from the Model menu, and click update to perform a **burn-in** of 1,000 iterations (the default), which takes **Os** at 1.6 Pentium GHz; (7) I then get a Sample Monitoring Tool from the Inference menu, and type sigma.theta and click set.



(8) I type 50000 in the updates window in the Update Tool and click update to get a **monitoring** run of **50,000** iterations (this took **15s**).

Then (9) selecting sigma.theta in the node window, all 10 buttons from clear through autoC are active, and I click on history (to get a Time Series window), density (to get a Kernel density window), autoC to get an Autocorrelation function window, and stats (to get a Node statistics window), yielding the screen above.

The output of an MCMC sampler, when considered as a **time series**, often exhibits **positive autocorrelation**; in fact it often looks like a realization of an **autoregressive**  $AR_p$  model of order p = 1 ( $\theta_t = \alpha + \beta \theta_{t-1} + e_t$ ) with **positive first-order autocorrelation**  $\rho$ .

This does not affect the **validity** of Monte Carlo inferences about the unknowns (e.g., the mean of any **stationary stochastic process** is a **consistent** estimator of the underlying process mean), but it does affect the **efficiency** of these inferences: for example, the Monte Carlo variance of the sample mean  $\bar{\theta}$  based on M draws from an  $AR_1$  time series is

$$V(\bar{\theta}) = \frac{\sigma_{\theta}^2}{M} \left( \frac{1+\rho}{1-\rho} \right), \tag{29}$$

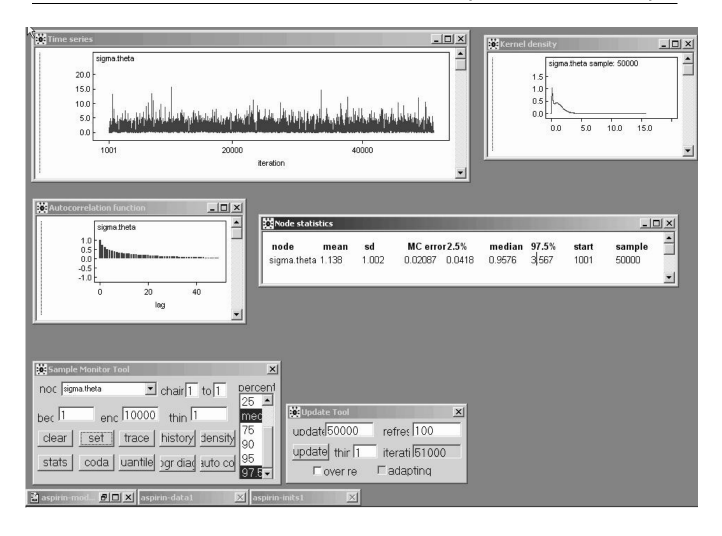
and the sample size inflation factor  $\frac{1+\rho}{1-\rho} \to \infty$  as  $\rho \to +1$ .

An MCMC sampler which produces output for any given unknown  $\theta$  with  $\rho$  near 0 (if  $\rho=0$  the output is white noise, i.e., equivalent to IID draws from the posterior) is said to be **mixing well** in that unknown.

The time series trace for  $\sigma_{\theta}$  above is only mixing **moderately** well: the autocorrelation function has the familiar ski-slope shape of an  $AR_1$  series with  $\rho \doteq 0.7$  (the height of the bar at lag 1).

The marginal posterior distribution for  $\sigma_{\theta}$  (from the Kernel density window) looks heavily skewed to the right, which makes sense for a scale parameter.

The **posterior mean** and **SD** of  $\sigma_{\theta}$  (using the  $\Gamma(\epsilon, \epsilon)$  prior for  $\tau_{\theta}$ ) are estimated to be 1.14 and 1.00, respectively; the **Monte Carlo standard error** of the posterior mean estimate is 0.021 (so that with 50,000 monitoring iterations I don't yet have **3 significant figures** of accuracy for the posterior mean); the **posterior median** is estimated to be 0.96; and a **95% central interval** for  $\sigma_{\theta}$  with this prior is estimated to run from 0.042 to 3.57.

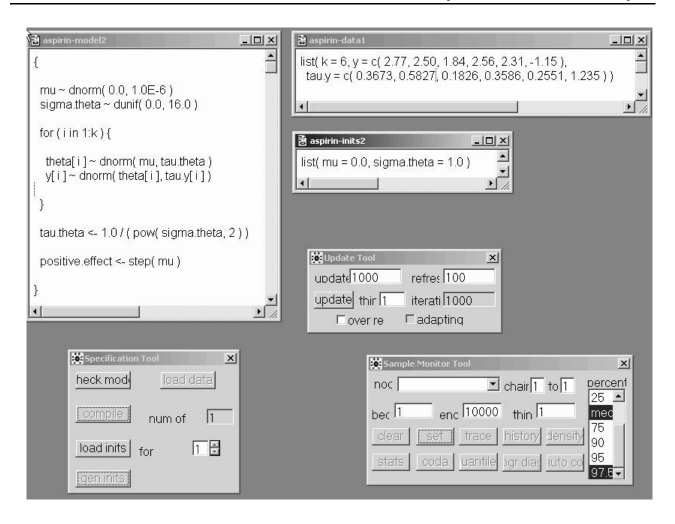


The main thing to notice, however, is that the **range of plausible values** for sigma.theta in its posterior is approximately from **0 to 16**.

It has recently been shown that the **simplest diffuse prior** on  $\sigma_{\theta}$  that has **good calibration properties** (i.e., such that 95% **nominal** Bayesian interval estimates for all of the parameters in model (10) do in fact have **actual coverage close to 95%**) is

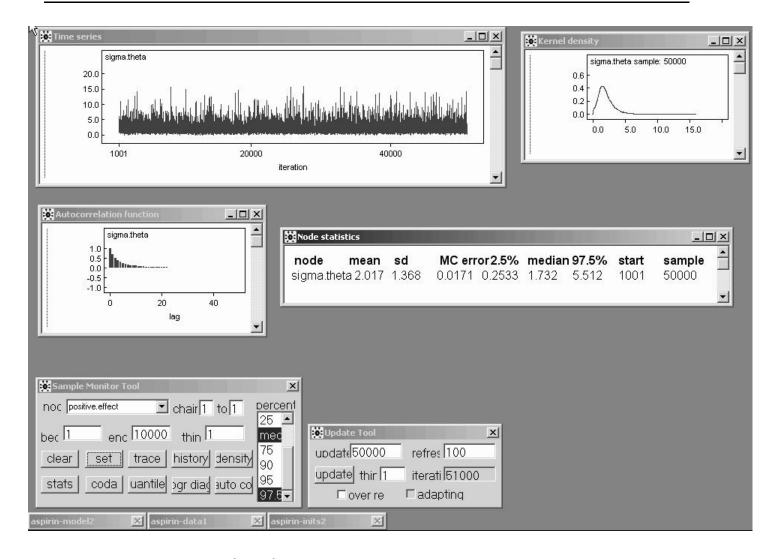
$$\sigma_{\theta} \sim U(0, c),$$
 (30)

where c is chosen to be (roughly) the **smallest value that** doesn't truncate the likelihood function for  $\sigma_{\theta}$ ; here it's evident that  $c \doteq 16$  will work well.



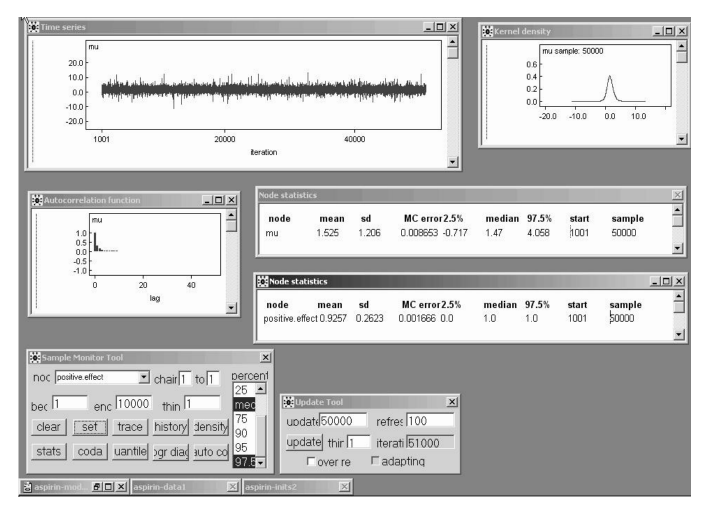
So I estimate a **second model** placing a Uniform(0,c) prior on  $\sigma_{\theta}$  (this model also requires a **new initial values file** that initializes sigma.theta instead of tau.theta).

This time in the Sample Monitor Tool I set all of the interesting quantities: mu, sigma.theta, theta, and positive.effect, and I use the same MCMC strategy as before (a burn-in of 1,000 followed by a monitoring run of 50,000).



With the Uniform(0,c) prior on  $\sigma_{\theta}$  the posterior mean of  $\sigma_{\theta}$  is now **sharply higher** than before (**2.02** versus the **1.14** value I got with the initial  $\Gamma(\epsilon,\epsilon)$  prior (this sort of **discrepancy** will only arise when the number of studies k is **small**; when it does arise I **trust** the results from the Uniform(0,c) prior).

Note that the posterior mean of  $\sigma_{\theta}$  is also **quite a bit bigger** than the value (1.24) obtained from **MLEB** back on page 25—this is a reflection of the **tendency of MLEB to understate the between-study heterogeneity** in model (10) with small k.

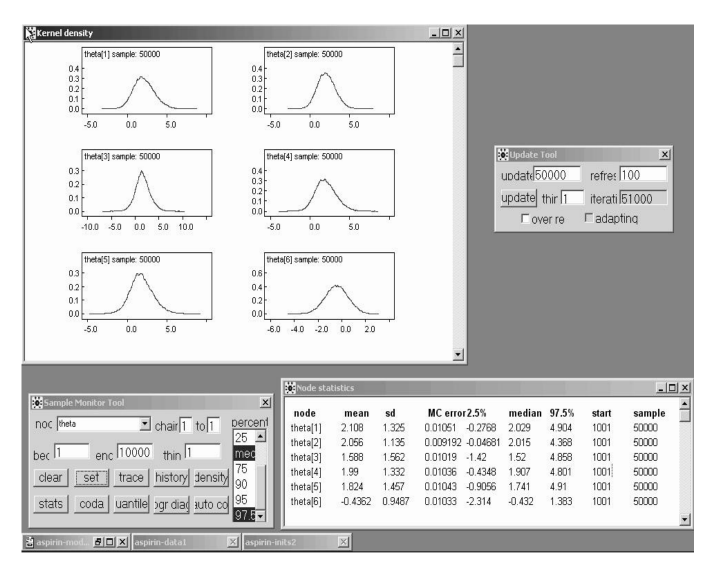


On pp. 25–26 above we saw that the MLEB estimate of  $\mu$  was **1.45** with an approximate standard error of **0.809**, and an approximate 95% confidence interval for  $\mu$  ran from -0.14 to +3.03.

The corresponding **Bayesian** results are: posterior **mean** 1.52, posterior **SD** 1.21, 95% interval (-0.72, 4.06).

As is often true, the simple MLEB approximations leading to these estimates have **underestimated the actual uncertainty** about  $\mu$ : the Bayesian 95% interval with the Uniform prior is **50% wider**.

It's easy to monitor the **posterior probability that aspirin** is beneficial, with the built-in step function applied to mu:  $P(\mu > 0 | \text{data, diffuse prior information}) \doteq \textbf{0.93}$ , i.e., posterior betting odds of about **12.5 to 1 that aspirin** reduces mortality.



The marginal density plots of the  $\theta_i$  values show **interesting** departures from normality, and the Bayesian estimates (a) exhibit rather less shrinkage and (b) have 27–43% larger uncertainty estimates.

Table 3.1. MLEB and Bayesian (posterior mean) estimates of the  $\theta_i$ .

	Maximum Likelihood		Bayesian Posterior	
study(i)	$\widehat{ heta}_i$	$\widehat{SE}(\widehat{ heta}_i)$	mean	SD
1	1.92	0.990	2.11	1.33
2	1.94	0.899	2.06	1.14
3	1.53	1.09	1.59	1.56
4	1.84	0.994	1.99	1.33
5	1.69	1.05	1.82	1.46
6	-0.252	0.728	-0.44	0.95

#### **Hierarchical Model Expansion**

Looking at the **shrinkage plot** on p. 26 or the **raw data values** themselves, it's evident that a **Gaussian** model for the  $\theta_i$  may not be appropriate: study 6 is so different than the other 5 that a **heavier-tailed distribution** may be a better choice.

This suggests **expanding** the HM (10), by embedding it in a **richer model class** of which it's a **special case** (this is the main Bayesian approach in practice to **dealing with model inadequacies**).

A **natural choice** would be a t model for the  $\theta_i$  with **unknown degrees of freedom**  $\nu$ :

$$(\theta, \sigma^2, \nu) \sim p(\theta, \sigma^2, \nu)$$
 (prior)   
 $(\theta_i | \theta, \sigma^2, \nu) \stackrel{\text{IID}}{\sim} t(\theta, \sigma^2, \nu)$  (underlying effects) (31)   
 $(y_i | \theta_i) \stackrel{\text{indep}}{\sim} N(\theta_i, V_i)$  (data).

Here  $\eta \sim t \left( \theta, \sigma^2, \nu \right)$  just means that  $\left( \frac{\eta - \theta}{\sigma} \right)$  follows a **standard** t **distribution** with  $\nu$  degrees of freedom. This is **amazingly easy** to implement in WinBUGS (it is considerably more difficult to carry out an **analogous ML analysis**).

The new model file is

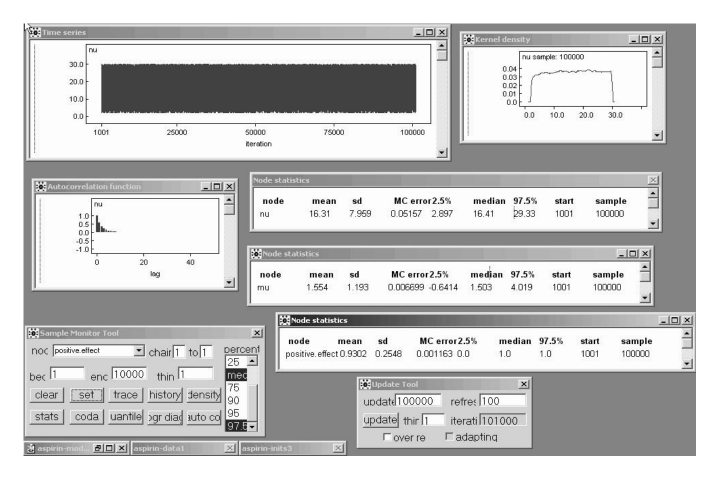
```
mu ~ dnorm( 0.0, 1.0E-6 )
sigma.theta ~ dunif( 0.0, 16.0 )
nu ~ dunif( 3.0, 30.0 )

for ( i in 1:k ) {
   theta[ i ] ~ dt( mu, tau.theta, nu )
   y[ i ] ~ dnorm( theta[ i ], tau.y[ i ] )
}

tau.theta <- 1.0 / pow( sigma.theta, 2 )
}</pre>
```

{

#### Model Expansion (continued)



To express comparative prior ignorance about  $\nu$  I use a **uniform** prior on the interval from 2.0 to 30.0 (below  $\nu=2$  the t distribution has **infinite variance**, and above about 30 it starts to be **indistinguishable** in practice from the Gaussian).

A burn-in of 1,000 and a monitoring run of 100,000 iterations takes about twice as long as with 50,000 iterations in the Gaussian model (i.e., about the same speed per iteration) and yields the posterior summaries above.

It's clear that there's little information in the likelihood function about  $\nu$ : the prior and posterior for this parameter virtually coincide.

The results for  $\mu$  and the  $\theta_i$  are **almost unchanged**; this would not necessarily be the case if study 6 had been **more extreme**.

#### **Educational Meta-Analysis**

#### **Incorporating Study-Level Covariates**

Case Study: Meta-analysis of the effect of teacher expectancy on student IQ (Bryk and Raudenbush, 1992). Do teachers' expectations influence students' intellectual development, as measured by IQ scores?

Table 5.4. Results from 19 experiments estimating the effects of teacher expectancy on pupil IQ.

	Weeks of	Estimated	Standard
	Prior	Effect	Error of
Study (i)	Contact $(x_i)$	Size $(y_i)$	$y_i = \sqrt{V_i}$
1. Rosenthal et al. (1974)	2	0.03	0.125
2. Conn et al. (1968)	3	0.12	0.147
3. Jose & Cody (1971)	3	-0.14	0.167
4. Pellegrini & Hicks (1972)	0	1.18	0.373
5. Pellegrini & Hicks (1972)	0	0.26	0.369
6. Evans & Rosenthal (1969)	3	-0.06	0.103
7. Fielder et al. (1971)	3	-0.02	0.103
8. Claiborn (1969)	3	-0.32	0.220
9. Kester & Letchworth (1972)	0	0.27	0.164
10. Maxwell (1970)	1	0.80	0.251
11. Carter (1970)	0	0.54	0.302
12. Flowers (1966)	0	0.18	0.223
13. Keshock (1970)	1	-0.02	0.289
14. Henrickson (1970)	2	0.23	0.290
15. Fine (1972)	3	-0.18	0.159
16. Greiger (1970)	3	-0.06	0.167
17. Rosenthal & Jacobson (1968)	1	0.30	0.139
18. Fleming & Anttonen (1971)	2	0.07	0.094
19. Ginsburg (1970)	3	-0.07	0.174

#### **Teacher Expectancy**

Raudenbush (1984) found k = 19 experiments, published between 1966 and 1974, estimating the effect of teacher expectancy on student IQ (Table 5.4).

In each case the experimental group was made up of children for whom teachers were (deceptively) encouraged to have high expectations (e.g., experimenters gave treatment teachers lists of students, actually chosen at random, who allegedly displayed dramatic potential for intellectual growth), and the controls were students about whom no particular expectations were encouraged.

The estimated **effect sizes**  $y_i = \frac{\bar{T}_i - \bar{C}_i}{\text{SD}_{i:pooled}}$  (column 3 in Table 5.4) ranged from -.32 to +1.18; why?

One good reason: the studies differed in **how well the experimental teachers knew their students** at the time they were given the deceptive information—this time period  $x_i$  (column 2 in Table 5.4) ranged from 0 to 3 weeks.

Figure 5.2 plots  $y_i$  against  $x_i$ —you can see that **the studies** with bigger  $x_i$  had smaller IQ effects on average.

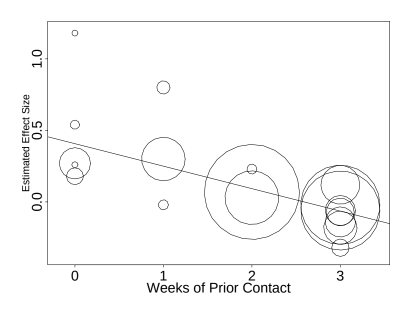


Figure 5.2. Scatterplot of estimated effect size against weeks of prior contact in the IQ meta-analysis. Radii of circles are proportional to  $w_i = V_i^{-1}$  (see column 4 in Table 5.4); fitted line is from weighted regression of  $y_i$  on  $x_i$  with weights  $w_i$ .

#### **Conditional Exchangeability**

Evidently model (1) will not do here — it says that your predictive uncertainty about all the studies is **exchangeable** (similar, i.e., according to (1) the underlying study-level effects  $\theta_i$  are like IID draws from a normal distribution), whereas Figure 5.2 **clearly shows** that the  $x_i$  are useful in predicting the  $y_i$ .

This is another way to say that your uncertainty about the studies is **not unconditionally exchangeable** but

conditionally exchangeable given x (Draper et al., 1993b).

In fact Figure 5.2 suggests that the  $y_i$  (and therefore the  $\theta_i$ ) are related **linearly** to the  $x_i$ .

Bryk and Raudenbush, working in the **frequentist** paradigm, fit the following HM to these data:

$$(\theta_i | \alpha, \beta, \sigma_{\theta}^2) \overset{\text{indep}}{\sim} N(\alpha + \beta x_i, \sigma_{\theta}^2)$$
 (underlying effects)   
  $(y_i | \theta_i) \overset{\text{indep}}{\sim} N(\theta_i, V_i)$  (data). (32)

According to this model the estimated effect sizes  $y_i$  are like **draws from a Gaussian** with mean  $\theta_i$  and variance  $V_i$ , the squared standard errors from column 4 of Table 5.4—here as in model (1) the  $V_i$  are taken to be known—and the  $\theta_i$  themselves are like **draws from a Gaussian** with mean  $\alpha + \beta x_i$  and variance  $\sigma_{\theta}^2$ .

The top level of this HM in effect assumes, e.g., that the 5 studies with x = 0 are sampled **representatively** from {all possible studies with x = 0}, and similarly for the other values of x.

This (and the Gaussian choice on the top level) are conventional assumptions, not automatically scientifically reasonable—for example, if you know of some way in which (say) two of the studies with x=3 differ from each other that's relevant to the outcome of interest, then you should include this in the model as a study-level covariate along with x.

#### An MLEB Drawback

Bryk and Raudenbush used MLEB methods, based on the **EM algorithm**, to fit this model.

As in Section 5.2, this estimation method combines the two levels of model (9) to construct a **single likelihood** for the  $y_i$ , and then **maximizes** this likelihood as usual in the ML approach.

They obtained  $(\hat{\alpha}, \hat{\beta}) = (.407 \pm .087, -.157 \pm .036)$  and  $\hat{\sigma_{\theta}}^2 = 0$ , naively indicating that **all of the study-level** variability has been "explained" by the covariate x.

However, from a **Bayesian** point of view, this model is **missing a third layer**:

$$(\alpha, \beta, \sigma_{\theta}^{2}) \sim p(\alpha, \beta, \sigma_{\theta}^{2})$$

$$(\theta_{i} | \alpha, \beta, \sigma_{\theta}^{2}) \stackrel{\text{indep}}{\sim} N(\alpha + \beta(x_{i} - \bar{x}), \sigma_{\theta}^{2})$$

$$(y_{i} | \theta_{i}) \stackrel{\text{indep}}{\sim} N(\theta_{i}, V_{i}).$$

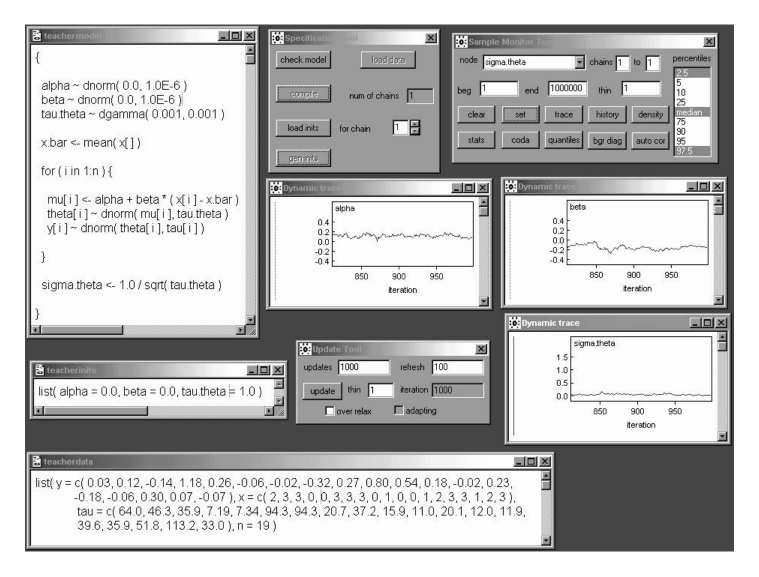
$$(33)$$

(it will help **convergence** of the sampling-based MCMC methods to make  $\alpha$  and  $\beta$  uncorrelated by **centering** the  $x_i$  at 0 rather than at  $\bar{x}$ ).

As will subsequently become clear, the trouble with MLEB is that in Bayesian language it assumes in effect that the posterior for  $\sigma_{\theta}^2$  is point-mass on the MLE. This is bad (e.g., Morris, 1983) for two reasons:

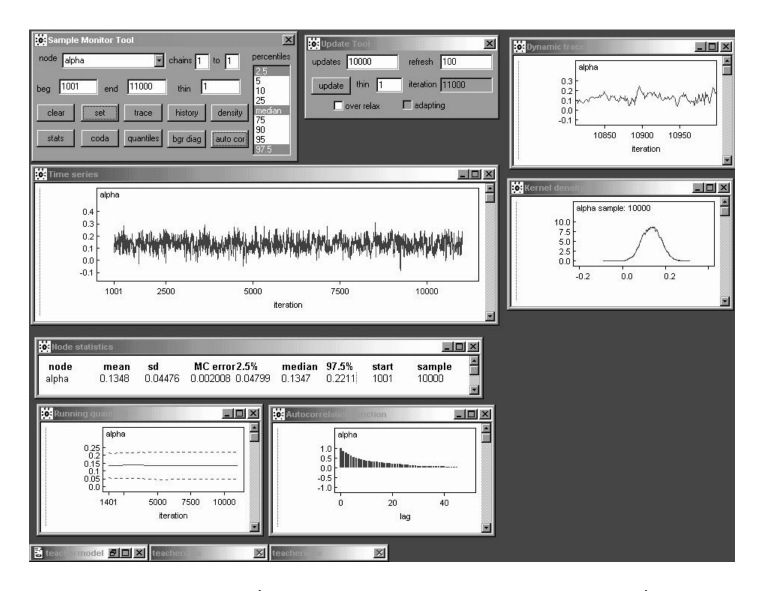
- ullet If the posterior for  $\sigma_{\theta}^2$  is highly **skewed**, the mode will be a **poor summary**; and
- Whatever point-summary you use, pretending the posterior SD for  $\sigma^2$  is zero **fails to propagate uncertainty** about  $\sigma_{\theta}^2$  through to uncertainty about  $\alpha, \beta$ , and the  $\theta_i$ .

The best way to carry out a fully Bayesian analysis of model (10) is with **MCMC** methods.



For  $p(\alpha, \beta, \sigma_{\theta}^2)$  in model (10) I've chosen the **usual** WinBUGS **diffuse prior**  $p(\alpha)p(\beta)p(\sigma_{\theta}^2)$ : since  $\alpha$  and  $\beta$  live on the whole real line I've taken marginal Gaussian priors for them with mean 0 and precision  $10^{-6}$ , and since  $\tau_{\theta} = \frac{1}{\sigma^2}$  is positive I use a  $\Gamma(0.001, 0.001)$  prior for it.

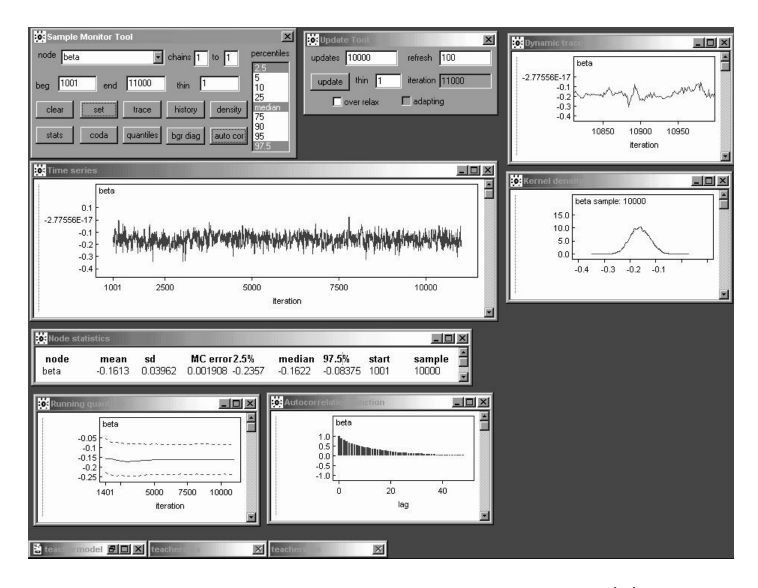
Model (10) treats the variances  $V_i$  of the  $y_i$  as **known** (and equal to the squares of column 4 in Table 5.4); I've **converted these into precisions** in the data file (e.g.,  $\tau_1 = \frac{1}{0.125^2} = 64.0$ ).



A burn-in of 1,000 (certainly longer than necessary) from default initial values  $(\alpha, \beta, \tau_{\theta}) = (0.0, 0.0, 1.0)$  and a monitoring run of 10,000 yield the following preliminary MCMC results.

Because this is a **random-effects model** we don't expect anything like IID mixing: the output for  $\alpha$  behaves like an  $AR_1$  time series with  $\hat{\rho}_1 \doteq 0.86$ .

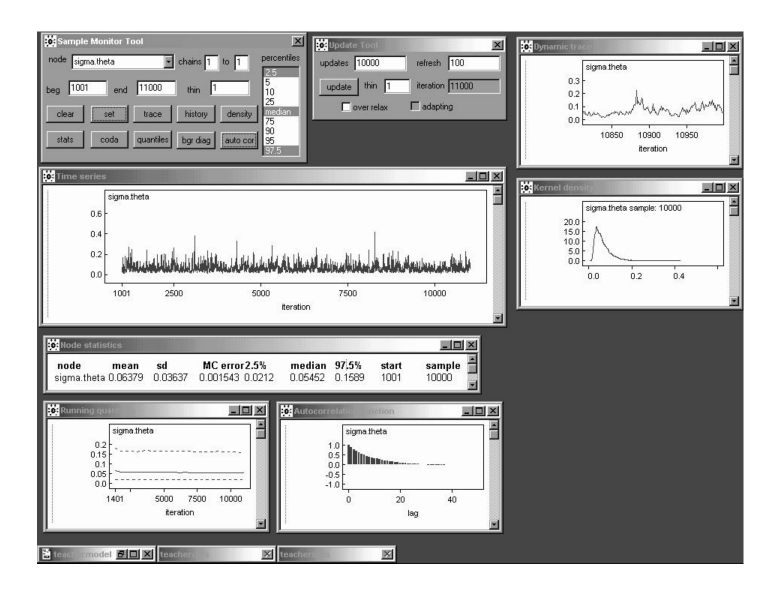
The posterior mean for  $\alpha$ , 0.135 (with an MCSE of 0.002), shows that  $\alpha$  in model (10) and  $\alpha$  in model (9) are **not comparable** because of the **recentering** of the predictor x in model (10): the MLE of  $\alpha$  in (9) was  $0.41 \pm 0.09$ .



But  $\beta$  means the same thing in both models (9) and (10): its posterior mean in (10) is  $-0.161 \pm 0.002$ , which is not far from the MLE -0.157.

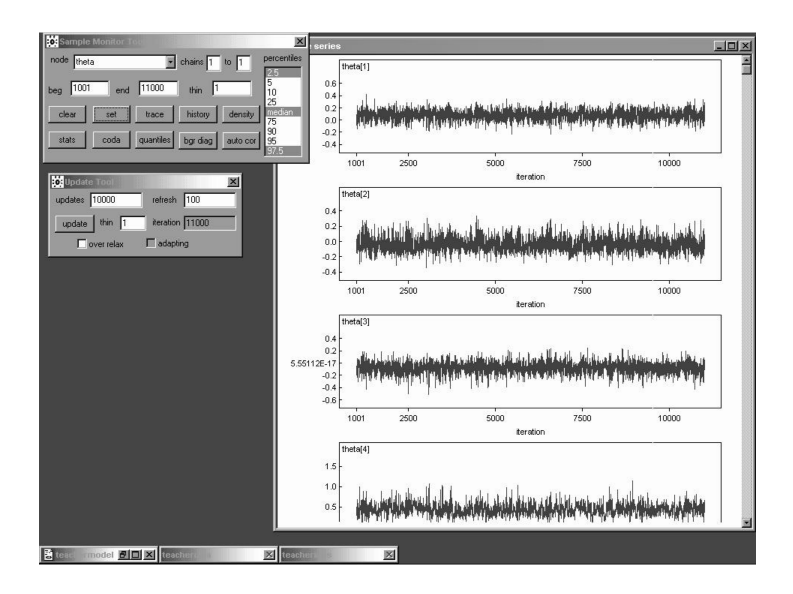
Note, however, that the posterior SD for  $\beta$ , 0.0396, is **10%** larger than the standard error of the maximum likelihood estimate of  $\beta$  (0.036).

This is a reflection of the **underpropagation of uncertainty** about  $\sigma_{\theta}$  in maximum likelihood mentioned on page 15.



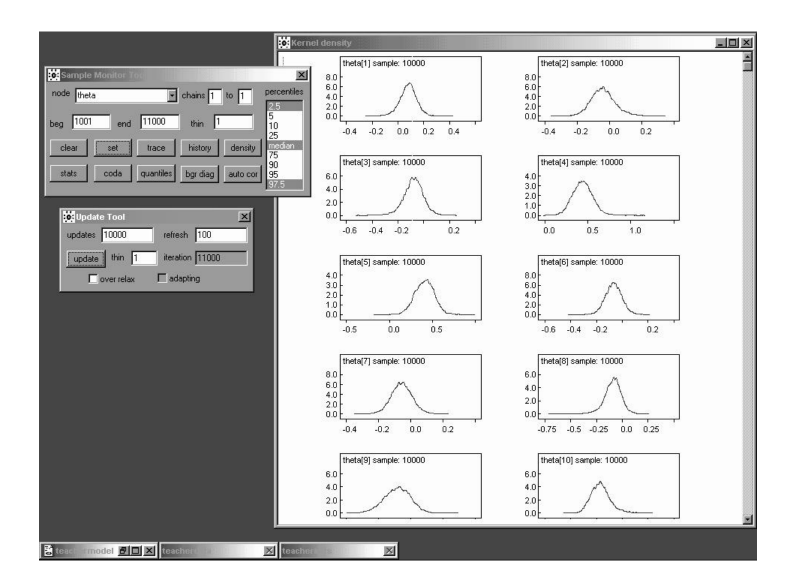
In these preliminary results  $\sigma_{\theta}$  has posterior mean  $0.064 \pm 0.002$  and SD 0.036, providing **clear evidence** that the MLE  $\hat{\sigma}_{\theta} = 0$  is a **poor summary**.

Note, however, that the likelihood for  $\sigma_{\theta}$  may be appreciable in the vicinity of  $\mathbf{0}$  in this case, meaning that some sensitivity analysis with diffuse priors other than  $\Gamma(0.001, 0.001)$ —such as U(0,c) for c around 0.5—would be in order.

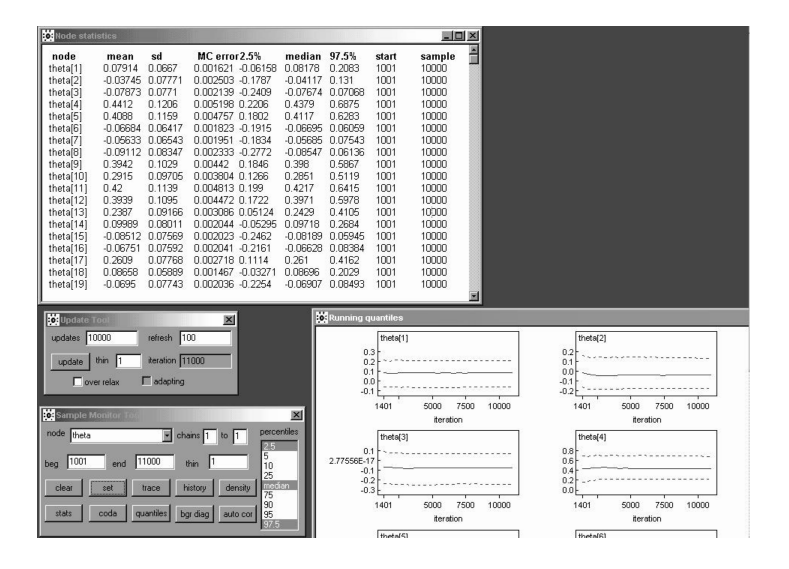


When you specify node theta in the Sample Monitor Tool and then look at the results, you see that WinBUGS presents **parallel findings with a single click** for all elements of the vector  $\theta$ .

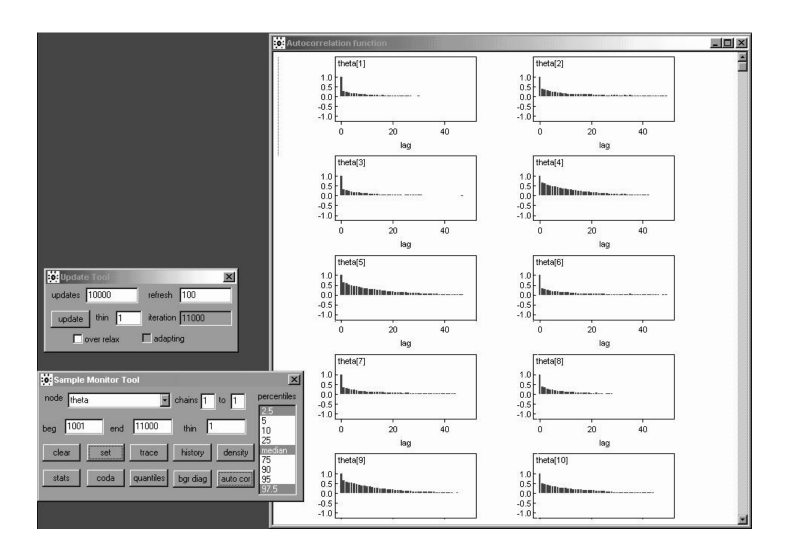
Some of the  $\theta_i$  are evidently mixing better than others.



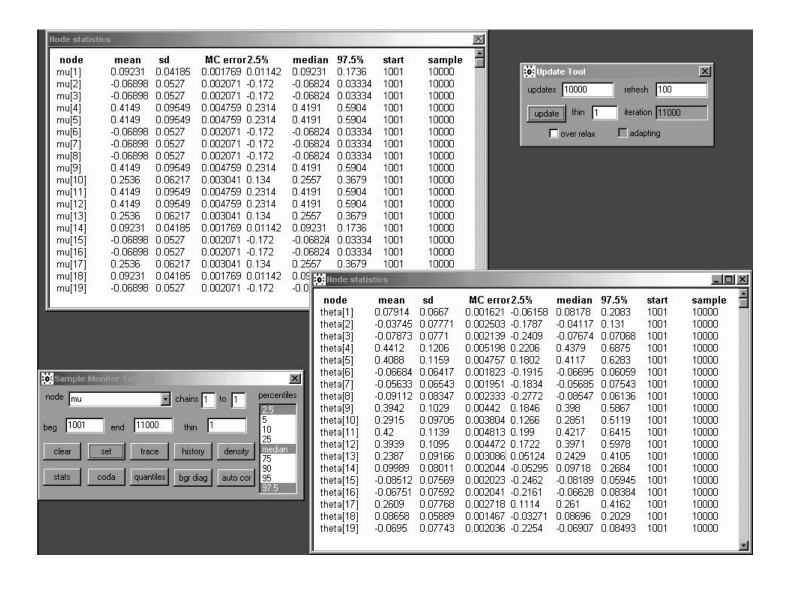
The marginal density traces of the  $\theta_i$  look rather like t distributions with fairly low degrees of freedom (fairly heavy tails).



Many of the  $\theta_i$  have **posterior probability** concentrated near **0**, but not all;  $\theta_4, \theta_5, \theta_9, \theta_{11}$ , and  $\theta_{12}$  are particularly large (looking back on page 12, what's **special** about the corresponding studies?).



Some of the  $\theta_i$  are **not far from white noise**; others are **mixing quite slowly**.



It's also useful to monitor the  $\mu_i = \alpha + \beta(x_i - \bar{x})$ , because they represent an **important part of the** shrinkage story with model (10).

## **Shrinkage Estimation**

In a manner parallel to the situation with the simpler model (1), the posterior means of the **underlying study effects**  $\theta_i$  should be at least approximately related to the **raw effect sizes**  $y_i$  and the  $\mu_i$  via the **shrinkage equation** 

$$E(\theta_i|y) \doteq (1 - \hat{B}_i)y_i + \hat{B}_i E(\mu_i|y); \qquad (34)$$

here  $\hat{B}_i = \frac{V_i}{V_i + \hat{\sigma}_{\theta}^2}$  and  $\hat{\sigma}_{\theta}^2$  is the posterior mean of  $\sigma_{\theta}^2$ .

#### This is easy to check in R:

```
> mu <- c( 0.09231, -0.06898, -0.06898, 0.4149, 0.4149, -0.06898, -0.06898, -0.06898, 0.4149, 0.2536, 0.4149, 0.4149, 0.2536, 0.09231, -0.06898, -0.06898, 0.2536, 0.09231, -0.06898 )
> y <- c( 0.03, 0.12, -0.14, 1.18, 0.26, -0.06, -0.02, -0.32, 0.27, 0.80, 0.54, 0.18, -0.02, 0.23, -0.18, -0.06, 0.30, 0.07, -0.07 )
> theta <- c( 0.08144, -0.03455, -0.07456, 0.4377, 0.4076, -0.0628, -0.05262, -0.08468, 0.3934, 0.289, 0.4196, 0.3938, 0.2393, 0.1014, -0.08049, -0.06335, 0.2608, 0.08756, -0.06477 )
> V <- 1 / tau
> B.hat <- V / ( V + 0.064^2 )
> theta.approx <- ( 1 - B.hat ) * y + B.hat * mu</pre>
```

# The Shrinkage Story (continued)

> cbind( y, theta, mu, sigma.2, B.hat, theta.approx )

```
B.hat theta.approx
              theta
                                    V
                          mu
 [1,]
      0.03 0.08144 0.09231 0.015625 0.7923026
                                                  0.07936838
 [2,]
      0.12 -0.03455 -0.06898 0.021609 0.8406536
                                                 -0.03886671
 [3,] -0.14 -0.07456 -0.06898 0.027889 0.8719400
                                                 -0.07807482
 [4,]
      1.18
           0.43770 0.41490 0.139129 0.9714016
                                                 0.43678060
 [5,]
     0.26
            0.40760 0.41490 0.136161 0.9707965
                                                  0.41037637
 [6,] -0.06 -0.06280 -0.06898 0.010609 0.7214553
                                                 -0.06647867
 [7,] -0.02 -0.05262 -0.06898 0.010609 0.7214553
                                                 -0.05533688
 [8,] -0.32 -0.08468 -0.06898 0.048400 0.9219750
                                                 -0.08856583
 [9,]
     0.27 0.39340 0.41490 0.026896 0.8678369
                                                  0.39574956
[10,]
     0.80 0.28900 0.25360 0.063001 0.9389541
                                                  0.28695551
[11,] 0.54 0.41960 0.41490 0.091204 0.9570199
                                                  0.42027681
[12,]
     0.18 0.39380 0.41490 0.049729 0.9239015
                                                  0.39702447
[13,] -0.02 0.23930 0.25360 0.083521 0.9532511
                                                  0.24080950
[14,]
     0.23 0.10140 0.09231 0.084100 0.9535580
                                                  0.09870460
                                                 -0.08445939
[15,] -0.18 -0.08049 -0.06898 0.025281 0.8605712
[16,] -0.06 -0.06335 -0.06898 0.027889 0.8719400
                                                 -0.06783002
[17,] 0.30 0.26080 0.25360 0.019321 0.8250843
                                                  0.26171609
            0.08756 0.09231 0.008836 0.6832663
[18,] 0.07
                                                  0.08524367
[19,] -0.07 -0.06477 -0.06898 0.030276 0.8808332
                                                 -0.06910155
```

You can see that equation (11) is indeed a **good** approximation to what's going on: the posterior means of the  $\theta_i$  (column 3 of this table, counting the leftmost column of study indices) all fall between the  $y_i$  (column 2) and the posterior means of the  $\mu_i$  (column 4), with the closeness to  $y_i$  or  $E(\mu_i|y)$  expressed through the **shrinkage factor**  $\widehat{B}_i$ .

Since  $\hat{\sigma}_{\theta}^2$  is small (i.e., most—but not quite all—of the between-study variation has been explained by the covariate x), the raw  $y_i$  values are shrunken **almost all of the way** toward the regression line  $\alpha + \beta(x_i - \bar{x})$ .