BAYESIAN HIERARCHICAL MODELS

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1. Bayesian Thinking

2. Hierarchical Models

3. Monte Carlo Study

4. Application

Bayesian Thinking

Bayes' Theorem

$$p(\theta \mid \text{data}) = \frac{p(\text{data} \mid \theta)p(\theta)}{p(\text{data})} \propto p(\text{data} \mid \theta)p(\theta)$$

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$$posterior = \frac{likelihood \times prior}{evidence} \propto likelihood \times prior$$

Setting: $\{y_i : i = 1, ..., n\}$ with $y_i \stackrel{\text{iid}}{\sim} \mathcal{N}(\mu, \sigma^2)$ and σ^2 known

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         Why?
```

Let $\mu \sim \mathcal{N}\left(\mu_0, \sigma_0^2\right)$

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$$p(\mu \mid y) \propto p(y \mid \mu)p(\mu)$$
$$\propto \exp\left(\frac{-n}{2\sigma^2}(\mu - \bar{y})^2\right) \exp\left(\frac{-1}{2\sigma_0^2}(\mu - \mu_0)^2\right)$$

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$$\propto \exp\left(-\frac{1}{2\sigma_u^2}(\mu - \bar{\mu})^2\right)$$

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$$\propto \exp\left(-\frac{1}{2\sigma_\mu^2}(\mu - \bar{\mu})^2\right)$$

$$\implies \mu \mid y \sim \mathcal{N} \left(\bar{\mu}, \sigma_\mu^2\right)$$

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, with
$$\sigma_{\mu}^{2} = \left(\frac{1}{\sigma^{2}/n} + \frac{1}{\sigma_{0}^{2}}\right)^{-1}$$

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$$= \alpha \bar{\mathbf{y}} + (1 - \alpha) \mu_{0}$$

A normal model with known variance and no features, really?



Object of interest: $\theta \mid data$

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Quantity of interest: $\mathbb{E}[h(\theta) \mid data] =: \mathbb{E}_{\theta}[h]$

```
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Quantity of interest: \mathbb{E}\left[h(\theta) \mid \text{data}\right] =: \mathbb{E}_{\theta}[h]

Estimation: Let \theta^{(1)}, ..., \theta^{(n)} \stackrel{\text{iid}}{\sim} p(\theta \mid \text{data}), then
\frac{1}{\sqrt{n}} \left(\sum_{i} h(\theta^{(i)}) - \mathbb{E}_{\theta}[h]\right) \stackrel{\text{d}}{\longrightarrow} \mathcal{N}\left(0, \omega\right)
```

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(i.) of unkown form

(ii.) very complex
```

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Problems: p(\theta \mid \text{data}) might be

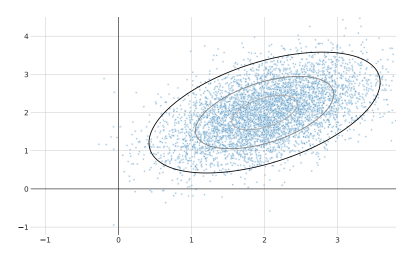
(i.) of unkown form

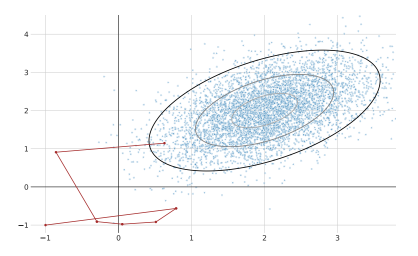
(ii.) very complex

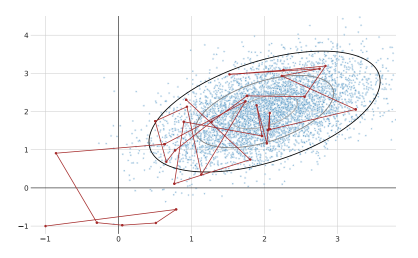
(iii.) only known up to an integration const.
```

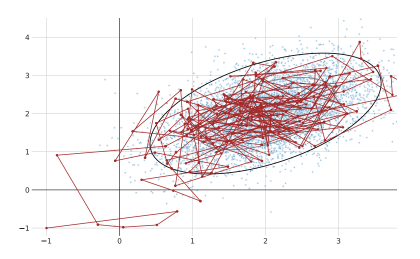
Algorithm Metropolis-Hastings (1953, 1970)

```
Input: (\pi, q, T) = (\text{target, proposal, no. of samples})
 1: initialize x_0 in supp q
 2: for t = 0, ..., T do
           candidate: y \sim q(\cdot \mid x_t)
 4: acceptance prob.: \mathcal{A} \leftarrow \min \left\{ \frac{\pi(y)}{\pi(x_t)} \frac{q(y \mid x_t)}{q(x_t \mid y)}, 1 \right\}
         update: x_{t+1} \leftarrow \begin{cases} y & \text{,with prob. } A \\ x_t & \text{,with remaining prob.} \end{cases}
 6: return \{x_t : t = 1, ..., T\}
```









Hierarchical Models

Structure of HM - Setup

Hierarchical Data:

Structure of HM - Setup

Hierarchical Data:

Individual Level: (y_i, x_i) for i = 1, ..., n

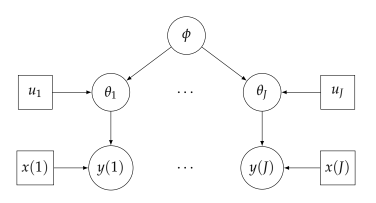
Structure of HM - Setup

Hierarchical Data: Individual Level: (y_i, x_i) for i = 1, ..., nGroup Level: u_i for j = 1, ..., J

Structure of HM - Setup

```
Hierarchical Data:
    Individual Level: (y_i, x_i) for i = 1, ..., n
    Group Level: u_i for j = 1, ..., J
Example:
    Test Outcome: y_i
    Parental Income: x_i
    Num. of Teachers: u_i
```

Structure of HM



The Prior Revisited

```
Before: Model: p(\text{data} \mid \theta)
```

Prior: $p(\theta)$

The Prior Revisited

```
Before:
     Model: p(\text{data} \mid \theta)
     Prior: p(\theta)
Now:
     Model: p(\text{data} \mid \theta, \phi) = p(\text{data} \mid \theta)
     Prior: p(\theta \mid \phi)
     Hyperprior: p(\phi)
```

The Posterior Revisited

Posterior:

$$p(\theta, \phi \mid \text{data}) \propto p(\text{data} \mid \theta)p(\theta, \phi)$$

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 $\propto p(\text{data} \mid \theta)p(\theta \mid \phi)p(\phi)$

The Posterior Revisited

Posterior:

$$p(\theta, \phi \mid \text{data}) \propto p(\text{data} \mid \theta)p(\theta, \phi)$$

$$\propto p(\text{data} \mid \theta)p(\theta \mid \phi)p(\phi)$$

$$p(\phi \mid \text{data}) \propto \int p(\theta, \phi \mid \text{data})d\theta$$

Setup: Individual *i* in group *j*

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Monte Carlo Study

Stan

What is Stan? C++ package (fast run times)

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Use cases: Bayesian/Maximum Likelihood
estimation of statistical models

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Use cases: Bayesian/Maximum Likelihood
estimation of statistical models
Interfaces: Pystan, Rstan, Stan.jl,...

$$y_i = \alpha + \beta_{j[i]} x_i + \epsilon_i$$
,
 $\epsilon \sim \mathcal{N}(0, 1)$

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,
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 $\beta_j = \gamma_0 + \gamma_1 u_j + \eta_j$,
 $\eta \sim \mathcal{N}(0, 1)$

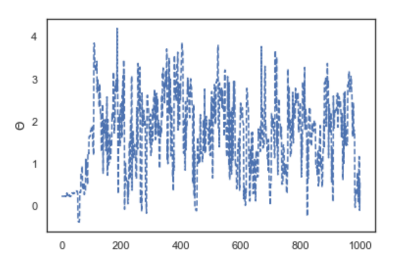
```
\begin{array}{ll} y_i = \alpha + \beta_{j[i]} x_i + \epsilon_i \,, & \text{data } \{ \\ \text{vector [N]} \ \ y; \\ \epsilon \sim \mathcal{N} \ (0,1) & \text{vector [N]} \ \ x; \\ \text{vector [N]} \ \ u; \\ \text{vector [N]} \ \ u; \\ \text{vector [N]} \ \ u; \\ \text{int < lower = 0 > } \ \ J; \\ \text{int < lower = 0 > } \ \ N; \\ \text{int < lower = 1, upper = J > } \\ \text{group [N]}; \end{array}
```

```
\begin{array}{ll} y_i = \alpha + \beta_{j[i]} x_i + \epsilon_i \,, & \text{parameter } \{ \\ \epsilon \sim \mathcal{N} \left( 0, 1 \right) & \text{real alpha} \,; \\ \beta_j = \gamma_0 + \gamma_1 u_j + \eta_j \,, & \text{real gamma\_1} \,; \\ \eta \sim \mathcal{N} \left( 0, 1 \right) & \text{vector[J] eta\_b} \,; \\ \eta \sim \mathcal{N} \left( 0, 1 \right) & \text{real < lower=0} > \text{sigma\_b} \,; \\ \end{array}
```

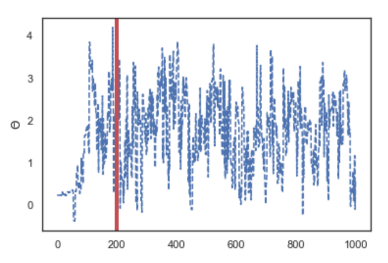
```
model
                                      for (i in 1:N) {
y_i = \alpha + \beta_{i[i]} x_i + \epsilon_i,
                                         beta[i] = gamma_0 +
                                                 u[i] * gamma_1 +
\epsilon \sim \mathcal{N}\left(0,1\right)
                                                     eta[group[i]]
\beta_i = \gamma_0 + \gamma_1 u_i + \eta_i,
                                         y_hat[i] = alpha +
                                                x[i] * beta[i];
\eta \sim \mathcal{N}\left(0,1\right)
                                         ~ normal(
                                                y_hat , sigma_y );
```

```
\begin{aligned} y_i &= \alpha + \beta_{j[i]} x_i + \epsilon_i \,, \\ &\epsilon \sim \mathcal{N} \left( 0, 1 \right) & \text{gamma}_0 \sim \text{normal} \left( 1 \,, \, 1 \right) \,; \\ \beta_j &= \gamma_0 + \gamma_1 u_j + \eta_j \,, \\ \eta \sim \mathcal{N} \left( 0, 1 \right) & \text{sigma}_y \sim \text{cauchy} \left( 0 \,, \, 5 \right) \,; \\ \text{sigma}_b \sim \text{cauchy} \left( 0 \,, \, 5 \right) \,; \end{aligned}
```

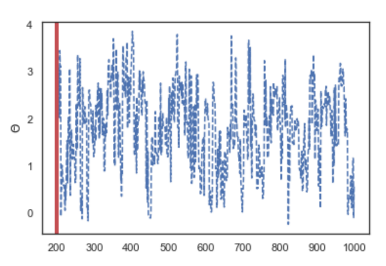
How can we be sure that we sample from the right distribution?

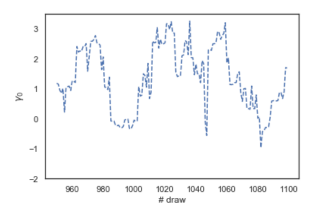


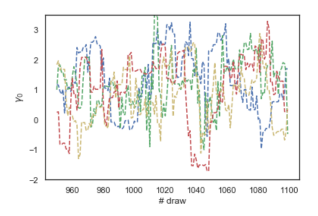
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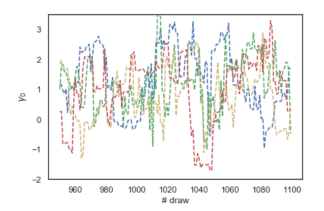


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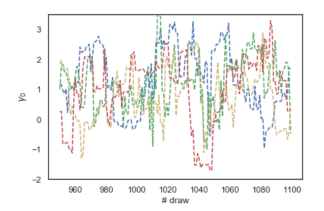






Variance of a single chain:

$$s_m^2 = \frac{1}{N-1} \sum_{n=1}^{N} (\theta_m^{(n)} - \overline{\theta}_m)^2$$



Average within chain variance:

$$W = \frac{1}{M} \sum_{m=1}^{M} s_m^2$$

Brooks and Gelman convergence criterium

Average Variance between chains:

$$B/N = \frac{1}{M-1} \sum_{m=1}^{M} (\overline{\theta}_m - \overline{\theta})^2$$

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Total Variance:

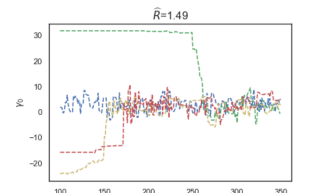
$$\widehat{\operatorname{Var}}^+(\theta \mid y) = \frac{N-1}{N}W + \frac{1}{N}B$$

Scale Reducing Factor:

$$\widehat{R} = \sqrt{\frac{\widehat{\text{Var}}^+(\theta \mid y)}{W}}$$

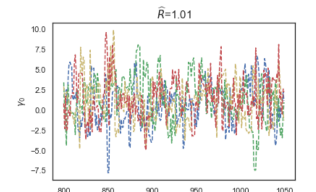
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Good Prior: $\gamma_0 \sim \mathcal{N}(1,1)$, $\gamma_1 \sim \mathcal{N}(1,1)$

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Bad Prior: $\gamma_0 \sim \mathcal{N}\left(2,1\right)$, $\gamma_1 \sim \mathcal{N}\left(2,1\right)$

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In all models: σ_v , $\sigma_b \sim \text{half-Cauchy}(0,5)$

Posterior Distribution - good prior

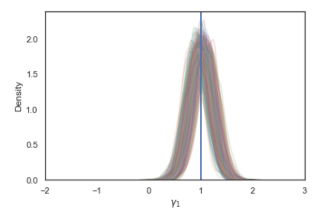


Figure: Posterior Draws of γ_1 with N=200, J=10 and 300 simulations

What happense if we decrease the number of levels *[*?

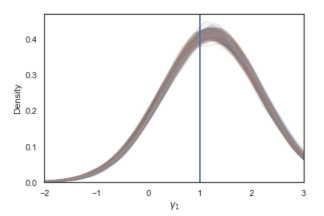


Figure: Posterior Draws of γ_1 with N=50, J=5 and 300 simulations

Posterior Distribution - bad prior

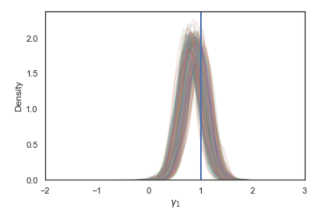


Figure: Posterior Draws of γ_1 with N=200, J=10 and 300 simulations

Posterior Distribution - weak, bad prior

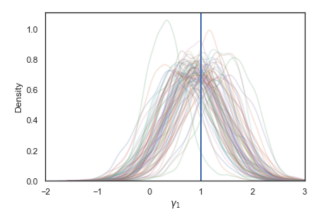


Figure: Posterior Draws of γ_1 with N=200, J=10 and 150 simulations

Not a good idea: Uniform Prior

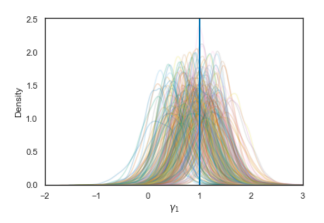


Figure: Posterior Draws of γ_1 with N=200, J=10 and 300 simulations

Increase sample size dramatically

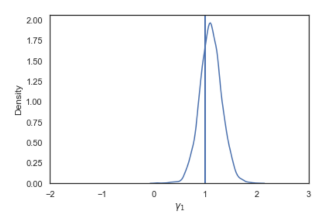


Figure: Single posterior draw for model with wrong prior and N=500, I=50

Application



The Data

Description: General Certificate of Secondary Education (GCSE) exam scores of 1,905 students from 73 schools in England on a science subject

Variables of interest: school identifier, student identifier, gender, total score on written paper and total score of course work.

ML and Bayesian Approach Application

Comparison

The model: Varying intercept and slope model with a single predictor

$$y_i = \alpha_{j[i]} + \beta_{j[i]} x_i + \epsilon_i, \tag{1}$$

$$\alpha_j = \mu_\alpha + \mu_j, \tag{2}$$

$$\beta_j = \mu_\beta + v_j, \tag{3}$$

$$y_i = \mu_{\alpha} + \mu_{\beta} x_i + u_{j[i]} + v_{j[i]} x_i + \epsilon_i$$
 (4)

Comparison

Maximum Likelihood (ML) Estimation

Package: lmer

The Imer Package: combines of ML estimation of model parameters and empirical Bayes (EB) predictions of the varying intercepts and/or slopes resulting in the Best Linear Unbiased Predictions (BLUPs) of the model parameters.

Why *lmer*?? allows for comparison between parameter estimates

Comparison

Bayesian Estimation:

Package: stan

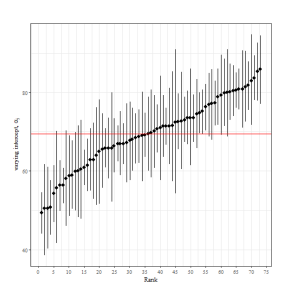
Priors: weekly informative normally distributed priors for hyperparameters

Results

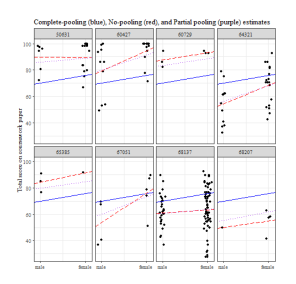
Dependent variab	le: Course	test score	
	ML	Bayes	
A. Random effects			(i.) point estimates
Intercept	- (10.146)	- (10.249)	almost the same
Female		- (7.000)	(ii.) Bayes standard
	(6.924)	(7.099)	deviations for
B. Fixed effects			random effects n
Intercept	69.425	69.413	
	(1.352)	(1.287)	be higher becaus
Female	7.128	7.132	ML does not take
	(1.131)	(1.165)	into account area
			into account gro
N			level variance
Students	1725	1725	
Schools	73	73	3'

Bayesian Approach: Further Analysis
Application

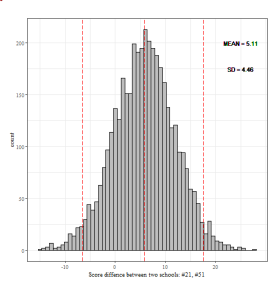
Posterior distribution ranking:



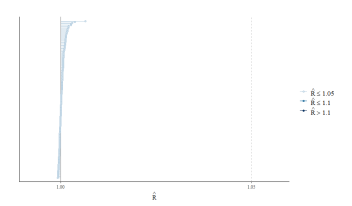
School specific regression lines and pooling:



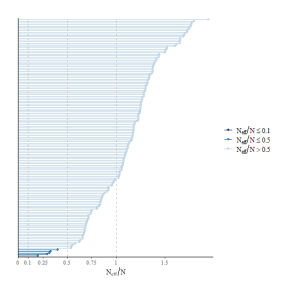
Making comparisons between individual schools:



Convergence



Convergence



https://github.com/timmens/

bayesian-hierarchical-models

http://mfviz.com/hierarchical-models/