## STA2201H Methods of Applied Statistics II

Monica Alexander

Week 8: Hierarchical models II

#### Overview

- ► Funnels of hell and reparameterization
- ► GLMs in a hierarchical context

# Reading

- Lesaffre and Lawson, 'Bayesian Biostatistics'. Lip cancer example is from here.
- ► GH Chapters 14-15
- BDA Chapters 15-16
- ► A nice overview of hierarchical funnels: https://arxiv.org/pdf/1312.0906.pdf

# Hierarchical models and funnels

#### Back to radon

$$y_{i}|\alpha_{j[i]}, \beta_{j[i]} \sim N\left(\alpha_{j[i]} + \beta_{j[i]}x_{i}, \sigma_{y}^{2}\right)$$
$$\alpha_{j} \sim N(\mu_{\alpha}, \sigma_{\alpha}^{2})$$
$$\beta_{j} \sim N(\mu_{\beta}, \sigma_{\beta}^{2})$$

- y<sub>i</sub> is log radon level
- $ightharpoonup \alpha_i$  is county-specific intercept
- $\triangleright \beta_j$  is county-specific slope
- ▶ x<sub>i</sub> is floor 1/ basement dummy

Let's fit this in Stan

#### Just a moment

$$y_{i}|\alpha_{j[i]}, \beta_{j[i]} \sim N\left(\alpha_{j[i]} + \beta_{j[i]}x_{i}, \sigma_{y}^{2}\right)$$
$$\alpha_{j} \sim N(\mu_{\alpha}, \sigma_{\alpha}^{2})$$
$$\beta_{j} \sim N(\mu_{\beta}, \sigma_{\beta}^{2})$$

- ▶ What's the interpretation of  $\sigma_{\beta}^2$ ?
- ▶ What happens to the  $\beta$ s when  $\sigma_{\beta}^2$  is small?

#### Fit in Stan

```
transformed parameters {
 vector[N] y_hat;
  for (i in 1:N)
    y_hat[i] = a1[county[i]] + a2[county[i]] * x[i];
model {
  mu_a1 \sim normal(0, 1);
  a1 ~ normal(mu_a1, sigma_a1);
  mu_a2 \sim normal(0, 1);
  a2 ~ normal(mu_a2, sigma_a2);
  y ~ normal(y_hat, sigma_y);
```

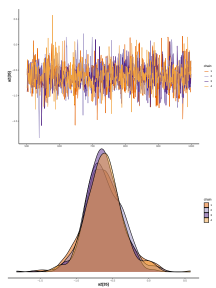
#### Fit in Stan

Fit to MN only. Here's some of the results:

```
## mean se_mean n_eff Rhat
## a1[1] 1.1658688 0.005594059 2011.76249 0.9997036
## a1[35] 1.0507016 0.005884807 1692.31830 0.9995272
## a2[1] -0.6108627 0.007584385 1726.54637 0.9995233
## a2[35] -0.6212198 0.006722520 1442.06768 1.0008974
## sigma_a1 0.3401203 0.001839355 665.76480 1.0046823
## sigma_a2 0.3251554 0.012747997 76.04905 1.0634959
```

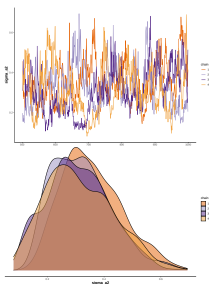
# Let's look at some diagnostics

Pick a county (number 35) and plot the trace and density

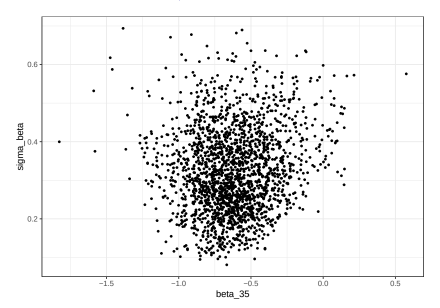


# What about the variance parameter

What's happening here?



# Scatterplot of $\beta_{35}$ and $\sigma_{\beta}$



#### Funnels of hell

- ► The density of these models looks like a 'funnel', with a region of high density but low volume below a region of low density and high volume
- ► This property of the posterior is a characteristic of the model and not a problem in itself, but makes sampling hard
- Especially a problem for Gibbs, but still a problem for HMC
- The narrower the space, the smaller the step size has to be
- Larger step sizes more likely to get rejected, so the sampler can get stuck

# We fit a centered parameterization

$$y_{i}|\alpha_{j[i]}, \beta_{j[i]} \sim N\left(\alpha_{j[i]} + \beta_{j[i]}x_{i}, \sigma_{y}^{2}\right)$$
$$\alpha_{j} \sim N(\mu_{\alpha}, \sigma_{\alpha}^{2})$$
$$\beta_{j} \sim N(\mu_{\beta}, \sigma_{\beta}^{2})$$

lacktriangle e.g.  $eta_j$  is directly dependent on hyperparameters  $\mu_eta$  and  $\sigma_eta^2$ 

## Non-centered parameterization

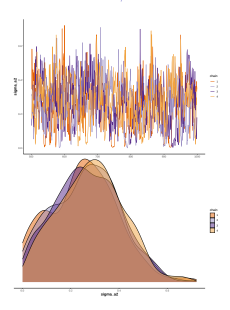
$$\begin{aligned} y_i | \alpha_{j[i]}, \beta_{j[i]} &\sim N \left( \alpha_{j[i]} + \beta_{j[i]} x_i, \sigma_y^2 \right) \\ &\cdots \\ \beta_j &= \mu_\beta + \eta_j \cdot \sigma_\beta \\ \eta_j &\sim N(0, 1) \\ &\cdots \end{aligned}$$

- $\beta_j$  is now a global mean + some offset, scaled by the  $\beta$ -specific standard deviation.
- ▶ In a non-centered parameterization we do not try to fit the group-level parameters directly, rather we fit a latent Gaussian variable from which we can recover the group-level parameters with a scaling and a translation
- The variables we are actually sampling are uncorrelated

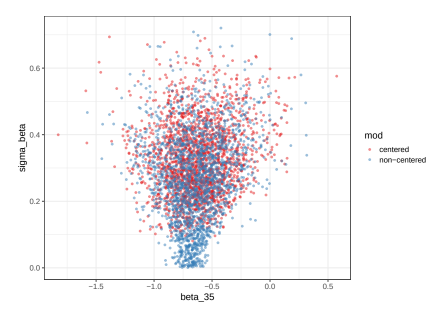
#### Non-centered Stan model

```
transformed parameters {
  vector[85] a1;
  vector[85] a2;
  vector[N] y_hat;
  a1 = mu_a1 + sigma_a1 * eta1;
  a2 = mu_a2 + sigma_a2 * eta2;
  for (i in 1:N)
    y_hat[i] = a1[county[i]] + a2[county[i]] * x[i];
model {
  mu_a1 \sim normal(0, 1);
  mu_a2 \sim normal(0, 1);
  eta1 \sim normal(0, 1);
  eta2 \sim normal(0, 1);
  y ~ normal(y_hat, sigma_y);
```

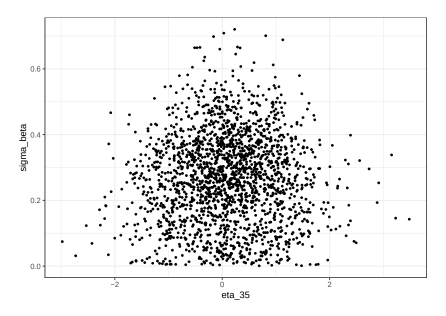
# Diagnostics for $\sigma_{\beta}$



# Better exploration of the funnel



# Less dependence between sampled parameters



# Summary

- Funnel structure of posterior density is a consequence of hierarchical model structure
- ▶ Not just the shape but the mass of density
- Mostly a problem when you don't have much data! More shrinkage = more mass in narrow bit of funnel

# GLMs in a hierarchical context

#### GLMs in a hierarchical context

- ► Last week we introduced hierarchical models in the setting where the data are assumed to be normally distributed
- But can easily extended to model other types of non-normal data hierarchically

Let's revisit the Poisson case:

$$y_i | \lambda_i \sim \text{Poisson}(\lambda_i)$$

where  $\lambda_i > 0$ , may depend on covariates, and vary by group membership j[i].

## Lip cancer

- Let's look at an example of estimating the risk of lip cancer by region in the former German Democratic Republic (Lesaffre and Lawson, chapter 9)
- ▶ In 1989, 2,342 deaths were recorded from lip cancer among males in 195 regions of GDR.
- For each region i we observed the number of deaths  $y_i$
- We also know the expected count e<sub>i</sub>, based on
  - age-specific mortality rates for whole country
  - age distribution of each region
- We also know the percentage of the population working outside
- ▶ Goal: estimate **relative risk** for each region  $\theta_i$

## Lip cancer

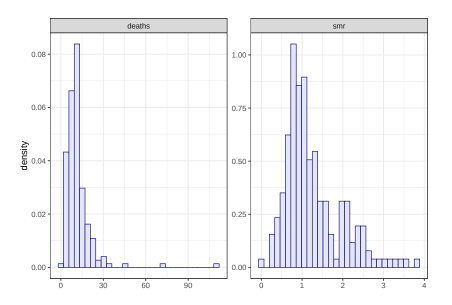
- ▶ Define the standardized mortality rate  $SMR_i = y_i/e_i$
- This is an estimate of the underlying relative risk  $\theta_i$ , which captures the relative difference in risk of dying for region i as compared to the reference population.
- Problem of small populations, some of the SMRs are based on very low counts, so are very uncertain
  - ▶ 15% based on counts < 5
  - ▶ 62% based on counts < 10

#### Set-up

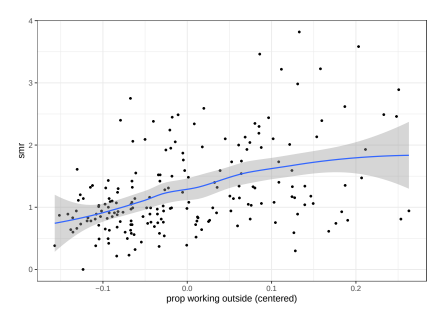
$$y_i \sim \text{Poisson } (\theta_i \cdot e_i)$$
  
 $\theta_i = ??$ 

From last week, in a broad sense, what are our three options for  $\theta_i$ ?

### What the data look like



### What the data look like



## Lip cancer

- 1. Model  $\theta_i$ s separately
- 2. Find one  $\theta$  for all regions
- 3. Use hierarchical model to exchange information about  $\theta_i$ s across regions

$$y_i \sim \text{Poisson } (\theta_i \cdot e_i)$$
 $\log \theta_i = \alpha_i + \beta(x_i^c)$ 
 $\alpha_i \sim N(\mu, \sigma_\mu^2)$ 

Where  $x_i^c$  is the (centered) percent of male population engaged in agriculture/forestry and fisheries in region i

## Lip cancer

#### Full model

$$y_i \sim \operatorname{Poisson} (\theta_i \cdot e_i)$$
 $\log \theta_i = \alpha_i + \beta(x_i^c)$ 
 $\alpha_i \sim N(\mu, \sigma_{\mu}^2)$ 
 $\mu, \beta \sim N(0, 1)$ 
 $\sigma_{\mu} \sim N_{+}(0, 1)$ 

# Fitting in Stan

- ▶ Relatively straightforward extension of normal models
- ► Be careful of types

## Fitting in Stan

```
data {
 int<lower=1> N;
 vector[N] x;
 vector[N] offset;
 int<lower=0> deaths[N];
 int<lower=0> region[N];
parameters {
 vector[N] alpha;
 real mu;
 real beta;
  real<lower=0> siama_mu;
model {
 vector[N] log_lambda;
 for (i in 1:N){
    loa_lambda[i] = alpha[i] + beta*x[i] + offset[i]:
 alpha ~ normal(mu, sigma_mu);
 mu \sim normal(0, 1);
 beta \sim normal(0,1);
  siama_mu ~ normal(0, 1);
  deaths ~ poisson_log(log_lambda);
```

#### Stan interlude

Note in the previous slide I wrote

```
deaths ~ poisson_log(log_lambda)
```

for the likelihood.

There are a million (well, at least two) other options, see here: https://mc-stan.org/docs/2\_18/functions-reference/poisson.html

- target += poisson\_lpmf( deaths | lambda)
  i.e. "Increment target log probability density with..."
- deaths ~ poisson(lambda) this is shorthand for the above, to make people used to BUGS/JAGS happier.

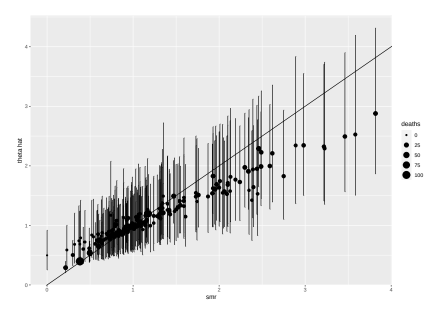
In the above example, I used poisson\_log so I didn't have to exponentiate my expression, but you do you.

Also note that I used an 'offset' in the same way that we did in glm, but you could equally write in terms of the expected deaths e.g. lambda[i] = theta[i]\*expected[i], etc etc

# Interpretation of coefficient on proportion working outside?

```
## mean se_mean n_eff Rhat
## mu 0.08663494 0.0005613354 4499.129 1.0006931
## beta 1.97939284 0.0059926765 3089.442 0.9996098
## sigma_mu 0.38681959 0.0006737879 2207.329 1.0009135
```

# Observed SMR v estimated $\theta_i$



## Overdispersion

- Recall that in many applications, counts are likely to be overdispersed
- Actually not bad in lip cancer case (how do you tell?)
- Going back k weeks, what were our two main options for dealing with overdispersion?

# Overdispersion with a quasi-Poisson set-up

In the lip cancer case we had

$$y_i \sim \text{Poisson } (exp(\alpha_i + \beta x_i^c) \cdot e_i)$$
 ...

For the overdispersed case we would have

$$y_i \sim \text{Poisson } (exp(\alpha_i + \beta x_i^c + \varepsilon_i) \cdot e_i)$$
...
 $\varepsilon_i \sim N(0, \sigma_{\varepsilon}^2)$ 
...

How to compare?

# Logistic regression

# Logistic regression in a hierarchical context

Can easily extend these hierarchical models to model binary outcomes and the probability of an event happening by various groups of interest, e.g. in the simplest form we would have

$$y_i|\pi_i\sim \operatorname{Bern}\left(\pi_i\right),\operatorname{OR}$$
  $y_i|\pi_i\sim \operatorname{Bin}\left(n_i,\pi_i\right),\ ext{if total number of trials is }n_i$  and then

$$ext{logit}(\pi_i) = \log\left(\frac{\pi_i}{1 - \pi_i}\right)$$

$$= ext{ some function of covariates, e.g.}$$
 $ext{logit}(\pi_i) = \beta_0 + \beta_1 x_i$ 

and then could put some hierarchical structure on the  $\beta$ 's for instance.

# Fertility intentions

- Using data from the 2015-2017 National Survey of Family Growth
- Gathers information on family life, marriage and divorce, pregnancy, infertility, use of contraception, and men's and women's health in the US.
- Some microdata are public: https://www.cdc.gov/nchs/nsfg/index.htm

# Fertility intentions

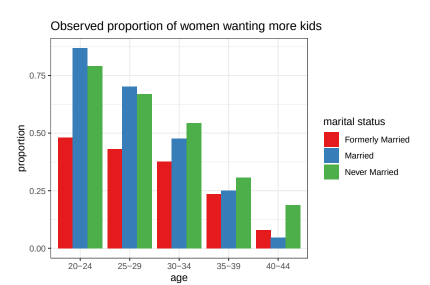
Interested in looking at fertility intentions, i.e. do women intend to have more children (and if so, how many)

- predictor of childbearing
- gaps tell you something about unmet need, due to social, economic conditions, cultural changes

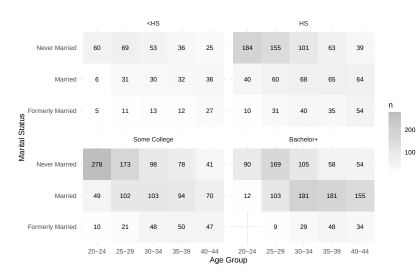
We are interested in the probability that a women wants more children.

- Individual covariates: age, education, marital status
- Possible extensions: state/region, parity (current number of children)

# Some clear patterns



# Sample size differences



#### Non-nested hierarchical model

- ➤ So far we have considered the simplest hierarchical structure of individuals *i* in groups *j*
- In the polls example, individuals have different group memberships (based on age and education) that we may want to pool across.

#### E.g. a reasonable model set-up would be

- to have hierarchies for age and education
- an extended model that includes geography may also have county nested within state, for example

## Just one hirerachical model

A hierarchical model with just age would be

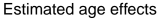
$$egin{aligned} y_i | \pi_i &\sim \mathsf{Bern}\left(\pi_i
ight) \ \mathsf{Pr}\left(y_i = 1
ight) = \pi_i = \mathsf{logit}^{-1}\left(lpha_{j[i]}
ight) \ lpha_j &\sim \mathsf{N}\left(\mu_lpha, \sigma_lpha^2
ight), \ \mathsf{for} \ j = 1, \dots, 5 \end{aligned}$$

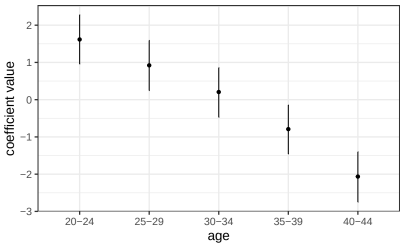
### Add in other effects

```
\begin{split} y_i | \pi_i &\sim \mathsf{Bern}\left(\pi_i\right) \\ \pi_i &= \mathsf{logit}^{-1}\left(\beta_0 + \beta_1\mathsf{formerly married}_i + \beta_2\mathsf{married}_i + \alpha_{j[i]}^{\mathsf{age}} + \alpha_{k[i]}^{\mathsf{edu}}\right) \\ \alpha_j^{\mathsf{age}} &\sim \mathsf{N}\left(0, \sigma_{\mathsf{age}}^2\right), \ \mathsf{for} \ j = 1, \dots, 5 \\ \alpha_k^{\mathsf{edu}} &\sim \mathsf{N}\left(0, \sigma_{\mathsf{edu}}^2\right), \ \mathsf{for} \ k = 1, \dots, 4 \end{split}
```

- ▶ Indexes: individual i, age j, education k
- Notice that the effects for age, educ, are centered around zero and now there is a 'global' intercept  $\beta_0$ 
  - $\blacktriangleright$  any non-zero means for the  $\alpha s$  could be folded into the global intercept
- The baseline category is never married

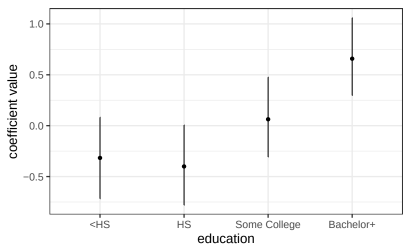
# Some results



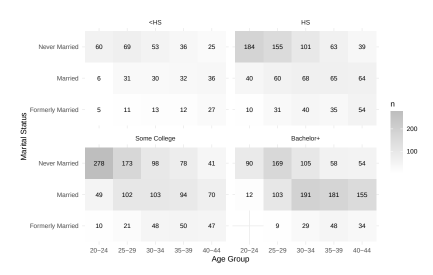


## Some results

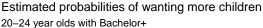


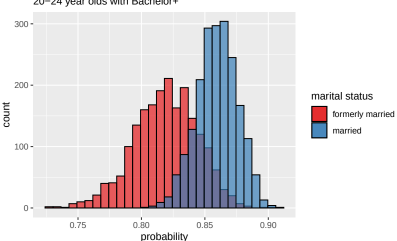


## Remember we had no observations from one cell



## Estimates from model



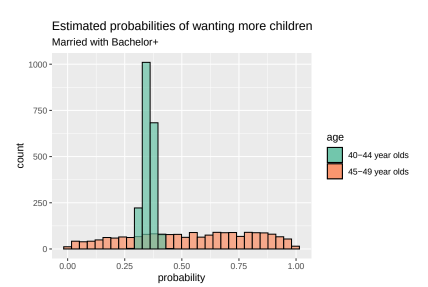


# What about getting estimates for the 45-49 year olds?

How do we generate these?

$$\begin{split} y_i | \pi_i &\sim \mathsf{Bern}\left(\pi_i\right) \\ \pi_i &= \mathsf{logit}^{-1}\left(\beta_0 + \beta_1\mathsf{formerly married}_i + \beta_2\mathsf{married}_i + \alpha_{j[i]}^{\mathsf{age}} + \alpha_{k[i]}^{\mathsf{edu}}\right) \\ \alpha_j^{\mathsf{age}} &\sim \mathsf{N}\left(0, \sigma_{\mathsf{age}}^2\right), \; \mathsf{for} \; j = 1, \dots, 5 \\ \alpha_k^{\mathsf{edu}} &\sim \mathsf{N}\left(0, \sigma_{\mathsf{edu}}^2\right), \; \mathsf{for} \; k = 1, \dots, 4 \end{split}$$

# Results, compared to 40-44 year olds



## Issues

We draw a new  $\tilde{\alpha}_{j}^{\mathrm{age}}$  based on

$$ilde{lpha}_{j}^{\mathsf{age}} \sim \mathsf{N}\left(\mathsf{0}, \sigma_{\mathsf{age}}^{\mathsf{2}}\right)$$

This assumes the age effects are conditionally independent of each other and centered at zero.

But we can do better!

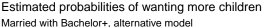
# An alternative specification on age

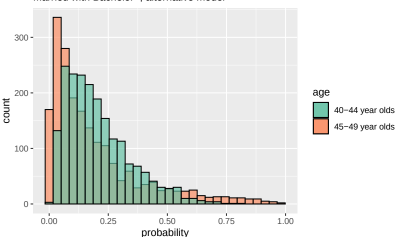
Consider the model as before but with one change:

$$\begin{split} y_i | \pi_i \sim \mathsf{Bern}\left(\pi_i\right) \\ \pi_i &= \mathsf{logit}^{-1}\left(\beta_0 + \beta_1 \mathsf{formerly married}_i + \beta_2 \mathsf{married}_i + \alpha_{j[i]}^{\mathsf{age}} + \alpha_{k[i]}^{\mathsf{edu}}\right) \\ \alpha_j^{\mathsf{age}} &\sim \mathsf{N}\left(\alpha_{j-1}^{\mathsf{age}}, \sigma_{\mathsf{age}}^2\right), \ \mathsf{for} \ j = 2, \dots, 5 \\ \alpha_k^{\mathsf{edu}} &\sim \mathsf{N}\left(0, \sigma_{\mathsf{edu}}^2\right), \ \mathsf{for} \ k = 1, \dots, 4 \end{split}$$

- ► This is assuming that the age effect in group j is similar to that in the previous age group (plus some noise)
- We are placing a random walk on the age effects; assuming some structure over age
- Note: need to put a prior on the first age group here

# Now look at the estimated probabilities for 45-49 year olds





# Summary

- Can model non-normal data in hierarchical setting
- Can have different hierarchies going on at the same time
- In the fertility intentions case, hierarchical models makes sense here because cell counts get very small: want to stabilize estimates.
- ► We can think about structured priors on the hierarchical effects (e.g. over age)
  - this idea extends to other dimensions, e.g. particularly thinking about geographic space