Class 6: Hierarchical generalised linear models

Andrew Parnell andrew.parnell@mu.ie

PRESS RECORD

Learning outcomes:

- Inderstand the modelling implications of moving from linear to hierarchical generalised linear models (HGLMs)
- \triangleright Know some of the different versions of Hierarchical GLMs
- \triangleright Be able to fit HGLMS in JAGS
- \triangleright Be able to expand and summarise fitted models

From LMs to HGLMs

- \blacktriangleright Reminder: a hierarchical model has prior distributions on the parameters which depend on further parameters
- \triangleright A generalised linear model is one in which the probability distribution is not normal, and a link function serves to match the mean of the distribution to the covariates
- \triangleright Within this framework, we can borrow the ideas from the previous class to create hierarchical GLMs
- ▶ We will go through four examples: binomial-logit, Poisson, robust regression, and ordinal regression

Example 1: binomial-logit

In class 2, we met the Binomial-logit model for binary data:

$$
y_i \sim Bin(1, p_i), logit(p_i) = \alpha + \beta(x_i - \bar{x})
$$

Here logit (p_i) is the link function equal to log $\left(\frac{p_i}{1-p_i}\right)$ $1-p_i$) and transforms the bounded probabilities into an unbounded space

If we have non-binary data we just change the likelihood:

$$
y_i \sim Bin(N_i, p_i), logit(p_i) = \alpha + \beta(x_i - \bar{x})
$$

In a hierarchical version of this model, we vary the *latent parameters* α and β and give them prior distributions

The swiss willow tit data

swt = **read.csv**('../data/swt.csv', stringsAsFactors = TRUE) **head**(swt)

A hierarchical model

In Suppose we want to fit a model on the sum $y_i = \text{rep.1} + \text{rep.2} + \text{rep.3}$:

$$
y_i \sim Bin(N_i, p_i), logit(p_i) = \alpha_{\text{altitude}_i} + \beta_{\text{altitude}_i}(x_i - \bar{x})
$$

where x_i is the percentage of forest cover

- \triangleright What prior distributions should we use for *α* and *β*?
- \triangleright Useful side note: A value of 10 on the logit scale leads to a probability of about 1, and a value of -10 leads to a probability of about 0 (you can test this by typing inv. $logit(10)$) so I wouldn't expect the value of $logit(p_i)$ to ever get much bigger than 10 or smaller than -10
- I have no idea whether we are more likely to find these birds in high percentage forest or low, so I'm happy to think that *β* might be around zero, and be positive or negative. Forest cover ranges from 0 to 100 so that suggests that *β* is very unlikely to be bigger than 0.1 or smaller than -0.1. Perhaps $\beta \sim \mathcal{N}(0,0.1^2)$ is a good prior
- It looks to me like the intercept is very unlikely to be outside the range $(-10, 10)$ so perhaps $\alpha \sim \mathcal{N}(0, 5^2)$ is appropriate

JAGS code

```
jags code = 'model{
  # Likelihood
  for(i \text{ in } 1:N) {
    y[i] ~ dbin(p[i], N exp[i])
    logit(p[i]) \leftarrow alpha[alt[i]] + beta[alt[i]] * (x[i] - mean(x))}
  # Priors
  for(j in 1:N alt) {
    alpha[j] ~ dnorm(mu_alpha, sigma_alpha^-2)
    beta[j] ~ dnorm(mu_beta, sigma_beta^-2)
  }
  mu alpha \sim dnorm(0, 5^{\sim}-2)mu beta ~ dnorm(0, 0.1^{\circ}-2)sigma alpha ~\sim dt(0, 5^{\sim}-2, 1)T(0, )sigma beta \sim dt(0, 5^{\degree}-2, 1)T(0,)}
```
Model fit - intercepts

Model fit - Slopes

Model fit - estimated mean proportions

Type 2: Poisson HGLMs

- \triangleright For a Poisson distribution there is no upper bound on the number of counts
- \triangleright We just change the likelihood (to Poisson) and the link function (to log):

$$
y_i \sim Po(\lambda_i), \log(\lambda_i) = \alpha + \beta(x_i - \bar{x}))
$$

- \triangleright We can now add our hierarchical layers into α and β , or. . .
- **I** Another way we can add an extra layer is by giving $log(\lambda_i)$ a probability distribution rather than setting it to a value
- \blacktriangleright This is a way of introducing *over-dispersion*, i.e. saying that the data are more variable than that expected by a standard Poisson distribution with our existing covariates

An over-dispersed model

 \blacktriangleright The over-dispersed model looks like:

$$
y_i \sim Po(\lambda_i), \log(\lambda_i) \sim N(\alpha + \beta(x_i - \bar{x}), \sigma^2)
$$

where σ is the over-dispersion parameter

- \triangleright We now need to estimate prior distributions for *α*, *β*, and *σ*
- \triangleright We will use the SWT data again, but pretend that we didn't know that they had gone out N times looking for the birds

JAGS code for OD Poisson

```
jags_code = '
model{
  # Likelihood
  for(i \text{ in } 1:N) {
    y[i] ~ dpois(exp(log lambda[i]))
    log lambda[i] ~ dnorm(alpha + beta * (x[i] - mean(x)),
          sigma<sup>-2</sup>)
  }
  alpha \sim dnorm(0, 5^{\sim}-2)beta \sim dnorm(0, 0.1^{\circ}-2)sigma \sim dt(0, 5^{\degree}-2, 1)T(0,)}
```
Model run

Notes about OD Poisson model

 \triangleright The way to think about OD models is via the data generating process.

- \triangleright We could compare this model to one without over dispersion via DIC (or if time, cross validation). We should also compute a posterior predictive distribution for full comparison
- In general, the parameter values (i.e. alpha and beta) tend to be more uncertain when you add in over dispersion
- \triangleright Also in the data set is a variable called dur which represents how long they spent looking for the birds. This could be added in as an offset via the likelihood:
- $y[i]$ ~ dpois(dur[i] * exp(log_lambda[i]))
- \blacktriangleright How do Bayesians deal with outliers?
- \triangleright A common view is that we should delete these observations before we run the model, but what if we can't find a reason for doing so
- \triangleright A good Bayesian will include outliers as part of the model.
- \triangleright One way of doing this is by switching from a normal distribution to a t-distribution

Normal vs t

 $curve(dnorm, from = -5, to = 5)$ $curve(dt(x, df = 1), add = TRUE, col = 'red')$ $curve(dt(x, df = 4), add = TRUE, col = 'blue')$

Polluted data

 \triangleright Suppose we had some data which looked like this:

x

There are a few observations here which look a bit odd

JAGS code for a t-model

```
jags_code = '
model{
  # Likelihood
  for(i \text{ in } 1:N) {
    y[i] ~ dt(alpha + beta * (x[i] - mean(x)),
                   sigma\hat{-2}, df[i])
    df[i] \sim dcat(p)}
  alpha \sim dnorm(0, 1^{\sim}-2)beta \sim dnorm(0, 1^{\sim}-2)signa \sim dt(0,1,1)T(0,)}
```
Fitting the model

```
jags_run = jags(data = list(N = N,
                            p = rep(1,10)/10,
                            y = y,
                            x = x,
                parameters.to.save = c('alpha',
                                        'beta',
                                        'df'),
                model.file = textConnection(jags_code))
```
Output from the model

dfs = jags_run**\$**BUGSoutput**\$**median**\$**df pars = jags_run**\$**BUGSoutput**\$**mean cols = **rainbow**(10) $plot(x, y, col = cols[dfs])$ **lines**(x, **as.numeric**(pars**\$**alpha) **+ as.numeric**(pars**\$**beta)*****(x **- mean**(x)))

Prior distributions on the degrees of freedom

- \blacktriangleright Here I've set a prior distribution on the degrees of freedom parameter to be a categorical distribution with probabilities 0.1 for $df = 1, 2, \ldots, 10$
- In Smaller values of df mean that a data point is more likely to be an outlier
- \blacktriangleright The categorical distribution automatically looks up the right df value for each probability
- \blacktriangleright This model is impossible to fit in Stan, because it contains a discrete parameter

Type 4: Ordinal data HGLMs

- \triangleright Often we have a response variable which is ordinal, e.g. disagree, neutral, agree, etc
- \blacktriangleright There are lots of different (and complicated) ways to model such data
- **•** Perhaps the easiest is to think of it as a hierarchical model with 'cut-points' on a latent linear regression

An ordinal model example

If Suppose $y_i = \{$ disagree, neutral, agree $\}$ and we make it dependent on a latent continuous variable z_i , so that :

$$
y_i = \begin{cases} \text{ agree} & \text{if } z_i > 0.5\\ \text{neutral} & \text{if } -0.5 < z_i \leq 0.5\\ \text{disagree} & \text{if } z_i \leq -0.5 \end{cases}
$$

► We then give z_i a prior distribution, e.g. $N(\beta_0 + \beta_1 x_i, \sigma^2)$

Fitting ordinal models in JAGS

```
jags_code = '
model{
  # Likelihood
  for(i \text{ in } 1:N) {
    z[i] ~ dnorm(alpha + beta * (x[i] - mean(x)),
                        signa^2-2)y[i] ~ dinterval(z[i], cuts)
  }
  alpha \sim dnorm(0, 100^{\circ}-2)beta \sim dnorm(0, 100^{\circ}-2)sigma \sim dt(0, 10^{\circ}-2, 1)T(0, )}
```
Simulating some example data

```
N = 100alpha = -1beta = 0.2sigma = 0.51set.seed(123)
x = runif(N, 0, 10)cuts = c(-0.5, 0.5)z = rnorm(N, alpha + beta * (x - mean(x)), sigma)y = findInterval(z, cuts)
```
Simulated data - plot

x

Fitting in JAGS - needs initial values

}

```
jags_inits = function() {
  z = \text{runif}(N, -0.5, 0.5)z[y == 0] = \text{runif}(\text{sum}(y == 0), -1, -0.5)z[y==2] = runif(sum(y==2), 0.5, 1)
  return(list(z = z))jags run = jags(data = list(N = N,y = y,
                              x = x,
                              cuts = cuts).
                 inits = jags_inits,
                 parameters.to.save = c('alpha',
                                           'beta',
                                           'sigma'),
                 model.file = textConnection(jags_code))
```
Output

print(jags_run)

```
## Inference for Bugs model at "7", fit using jags,
## 3 chains, each with 2000 iterations (first 1000 discarded)
## n.sims = 3000 iterations saved
## mu.vect sd.vect 2.5% 25% 50% 75% 97.5% Rhat n.eff
## alpha -1.093 0.181 -1.531 -1.188 -1.068 -0.962 -0.815 1.052 46
## beta 0.221 0.052 0.135 0.186 0.215 0.250 0.345 1.044 51
## sigma 0.559 0.128 0.365 0.468 0.538 0.627 0.859 1.027 81
## deviance 0.000 0.000 0.000 0.000 0.000 0.000 0.000 1.000 1
##
## For each parameter, n.eff is a crude measure of effective sample size,
## and Rhat is the potential scale reduction factor (at convergence, Rhat=1).
##
## DIC info (using the rule, pD = var(deviance)/2)
## pD = 0.0 and DIC = 0.0## DIC is an estimate of expected predictive error (lower deviance is better).
```
Summary

- \triangleright We have now seen a number of different types of hierarchical GLM
- \blacktriangleright Many of the ideas of hierarchical linear models transfer over, but we can explore richer behaviour with hierarchical GLMs
- \triangleright These have all used the normal, binomial or Poisson distribution at the top level, and have allowed for over-dispersion, robustness, and ordinal data, to name just three