Mixed effects models Wheels within wheels

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Part 1: Mixed effects models

Motivation

- What are mixed effects models?
 - Scary math (matrix algebra)
 - Variance partitioning
 - Fixed effects vs. Random effects
- Working with random effects
 - Fixed vs. random effects
 - Model validation
- Exercise

What are mixed effects models?

Many different names:

- Mixed effects models
- 2 Random effects models
- 3 Hierarchical models
- 4 Empirical/Bayesian hierarchical models
- 6 Latent variable models
- 6 Split-plot models¹

I like the term heirarchical models, as this is the closest to what I will teach you

¹Earlier form of variance partitioning

Scary math

Unfortunately, we need a review of matrix algebra in order to explain this:

• This is a matrix:

$$A = \begin{bmatrix} 1 & 4 & 7 \\ 2 & 5 & 8 \\ 3 & 6 & 9 \end{bmatrix}$$

• This is a vector:

$$b = \begin{bmatrix} 1 & 2 & 3 \end{bmatrix}$$

• Multiplying them looks like this:

$$A \times b = Ab = 1 \times \begin{bmatrix} 1 \\ 2 \\ 3 \end{bmatrix} + 2 \times \begin{bmatrix} 4 \\ 5 \\ 6 \end{bmatrix} + 3 \times \begin{bmatrix} 7 \\ 8 \\ 9 \end{bmatrix} = \begin{bmatrix} 30 \\ 36 \\ 42 \end{bmatrix}$$

Why do we call them "linear models"?

- Linear mapping of coefficients onto a model matrix (from your data)
- Coefficients:

$$eta = egin{bmatrix} 0.1 & 1.8 & -0.03 \end{bmatrix}$$

Model matrix:

$$X = egin{bmatrix} 1 & 1 & 10 \ 1 & 1 & 12 \ 1 & 0 & 9 \ dots & dots & dots \end{pmatrix}$$

Multiplying them looks like:

$$\hat{y} = Xeta = egin{bmatrix} 1.60 \\ 1.54 \\ -0.17 \\ dots \end{bmatrix}$$

This is exactly what R does to fit models:

head(dat)

```
## y x site
## 1 1.5101095 -4.248450 g
## 2 3.7190900 5.766103 j
## 3 -4.3737644 -1.820462 f
## 4 30.1459331 7.660348 n
## 5 0.2777422 8.809346 o
## 6 -3.6978175 -9.088870 p

m1 <- lm(y~x,data=dat) #Uses x to predict y
summary(m1)
```

```
##
## Call:
## lm(formula = v \sim x. data = dat)
## Residuals:
       Min
                10 Median
                                         Max
## -18.2574 -4.1262 0.0296 3.1854 25.2780
## Coefficients:
              Estimate Std. Error t value Pr(>|t|)
## (Intercept) 2.7306 0.5772 4.731 4.92e-06 ***
## x
                0.7213
                       0.1020 7.074 4.60e-11 ***
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
## Residual standard error: 7.3 on 158 degrees of freedom
## Multiple R-squared: 0.2405, Adjusted R-squared: 0.2357
## F-statistic: 50.04 on 1 and 158 DF, p-value: 4.6e-11
```

This is exactly what R does to fit models (cont.):

```
head(model.matrix(m1))
    (Intercept)
             1 -4 248450
             1 5.766103
             1 -1.820462
             1 7.660348
             1 8.809346
## 5
             1 -9.088870
## 6
coef(m1)
## (Intercept)
    2 7305689 0 7212867
pred2 <- model.matrix(m1) %*% coef(m1) #predicted = matrix * coefs</pre>
head(data.frame(pred1=predict(m1),pred2)) #same thing!
        pred1
                  pred2
    -0.3337812 -0.3337812
     6.8895819 6.8895819
     1.4174942 1.4174942
     8.2558758 8.2558758
    9.0846325 9.0846325
## 6 -3.8251119 -3.8251119
```

Groups are coded by "dummy variables" (0s and 1s)

```
m2 <- lm(v~site.data=dat) #Use site to predict v
head(model.matrix(m2)) #0s and 1s used to identify groups
    (Intercept) siteb sitec sited sitee sitef siteg siteh sitei sitej sitek sitel
## 1
## 2
## 3
## 4
## 5
## 6
    sitem siten siteo sitep
## 1
## 2
## 3
## 4
             0
## 5
## 6
coef(m2) #This uses the 1st site as the "control" group
```

```
## (Intercept)
                     siteb
                                sitec
                                            sited
                                                        sitee
                                                                    sitef
     7.192416 -11.998464 -14.632803
                                         1.983649
                                                    -7.765354
                                                                -4.523079
                                sitei
                                                        sitek
                                                                    sitel
##
        siteg
                     siteh
                                            sitej
    -3.439621
                -8.280601
                            -4.306456
                                        -4.085855
                                                    -5.663021
                                                                -5.155112
        sitem
                     siten
                                siteo
                                            siten
##
    -6.226642
                 8.403599
                            -8.626661
                                       -10.934182
```

Structure of LMs... now with matrices!

• All linear models take the form:

$$\hat{y} = X\beta = b_0 1 + b_1 x_1 ... + b_i x_i$$

 $y \sim Normal(\hat{y}, \sigma)$

- y is a vector of data you want to predict
- \hat{y} is a vector of *predicted values* for y
- $X = \{1, x_1...\}$ is a matrix of *predictors* for y
- $\beta = \{b_0, b_1, ...\}$ is a vector of *coefficients*
- $y \sim Normal(\hat{y}, \sigma)$ means:
 - "y follows a Normal distribution with mean \hat{y} and SD σ "

Fixed effects vs. Random effects

Say that X is a model matrix coding for 10 sites², and y is something we're interested in predicting

$$\hat{y} = b_0 + X\beta$$

 $y \sim Normal(\hat{y}, \sigma)$

- Site coefficients (β) are unrelated to each other
- σ is the SD of *residuals*
- Site is a fixed effect

$$\hat{y} = b_0 + X\zeta$$
 $y \sim Normal(\hat{y}, \sigma)$
 $\zeta \sim Normal(0, \sigma_{site})$

- Site coefficients (ζ) are related to each other via a *Normal* distribution
- σ is the SD of *residuals*, σ_{site} is the SD of *sites*
- Site is a random effect

²Intercept is a separate variable

Mixed effects = fixed + random effects

A mixed effects model has both **fixed** and **random** effects

$$\hat{y} = X\beta + U\zeta$$
 $y \sim Normal(\hat{y}, \sigma)$
 $\zeta \sim Normal(0, \sigma_{site})$

- X =fixed effects matrix (e.g. intercept, temperature)
- β = fixed effects coefficients
- U = random effects matrix (e.g. sites)
- ζ = random effects coefficients
- σ , σ_{site} = variance terms

Mixed effect model example

Let's go back to our earlier example:

- We're interested in predicting *y* using *x* (fixed effects)
- Data was collected at a number of sites, which may affect y "somehow"
- Effect of each site is normally distributed

head(dat)

Mixed effect model example

```
summary(mm1)
## Linear mixed model fit by REML ['lmerMod']
## Formula: v ~ x + (1 | site)
     Data: dat
##
## REML criterion at convergence: 1040.4
##
## Scaled residuals:
       Min
                       Median
                                            Max
## -2 50816 -0 71380 -0 02682 0 69401 3 01951
##
## Random effects:
   Groups Name
                         Variance Std Dev
            (Intercept) 26.78
                                  5.175
                         31.68
                                  5.628
   Residual
## Number of obs: 160, groups: site, 16
##
## Fixed effects:
               Estimate Std. Error t value
## (Intercept) 2.00331
                           1 38297
                                     1 449
               0.76192
## x
                           0.08083
                                     9.426
## Correlation of Fixed Effects:
    (Intr)
## x -0.002
```

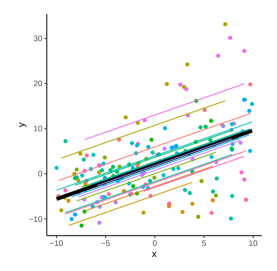
Results from 1mer model:

- Random effects:
 - residual and site variance (σ, σ_{site})
- Fixed effects:
 - Intercept and slope estimates (β)
 - No d.f. and p-value ³
 - If you need p-values for parameters, you can use the *lmerTest* package (or just calculate them yourself using means/SEs)

³'lme4' author doesn't think they can be calculated. I somewhat agree

Mixed effect model results

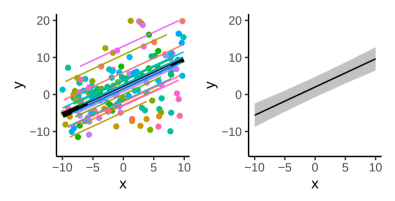
- In a random intercepts model, the regression line of x on y is allowed to move up or down around the main regression line for each site
- These changes in intercepts are normally distributed



Mixed effect model results (cont.)

For plotting, we want a partial effects plot that marginalizes across sites (i.e. "What does the trend look like at the average site?")

 ggpredict works well for this. If you want partial residuals, you'll have to add them in yourself using predict and residual



Random slopes + intercepts

Suppose that y wasn't just higher or lower at each site, but that the effect of x on y was higher or lower at each site

$$\hat{y} = X\beta + U\zeta_{int} + U_{x}\zeta_{slope}$$
 $y \sim Normal(\hat{y}, \sigma)$
 $\zeta_{int} \sim Normal(0, \sigma_{int})$
 $\zeta_{slope} \sim Normal(0, \sigma_{slope})$

- X =fixed effects matrix (e.g. intercept, temperature)
- β = fixed effects coefficients
- *U* = random intercept matrix (e.g. sites)
- $U_x = \text{random slopes matrix (e.g. temperature)}$
- ζ_{int} , ζ_{slope} = random intercept and slope coefficients
- σ , σ_{int} , σ_{slope} = variance terms

Random slope and intercept example:

```
#Intercept varies with site, and slope of x can
    also vary with site (both hierarchical)
mm2 \leftarrow lmer(v \sim x + (x|site), data=dat)
summary(mm2)
## Linear mixed model fit by REML ['lmerMod']
## Formula: v ~ x + (x | site)
     Data: dat
##
## REML criterion at convergence: 900.6
##
## Scaled residuals:
       Min
                     Median
## -2.10500 -0.64857 0.02414 0.61137 2.22996
## Random effects:
   Groups Name
                       Variance Std.Dev. Corr
            (Intercept) 35,2210 5,9347
                        0.7889 0.8882
                                        0.82
                        9.3162 3.0522
   Residual
## Number of obs: 160, groups: site, 16
##
## Fixed effects:
              Estimate Std. Error t value
## (Intercept) 2.0383
                          1.5091
                                 1.351
               0.6438
                          0.2275
                                  2.830
## Y
## Correlation of Fixed Effects:
    (Intr)
## x 0.790
```

Results from 1mer model:

- Random effects:
 - residual, slope, and site variance (σ , σ_{int} , σ_{slope})
 - Correlation b/w intercept and slope = 0.82
 - Sites with higher intercept also have a higher slope
- Fixed effects:
 - Intercept and slope estimates

Model matrices

X

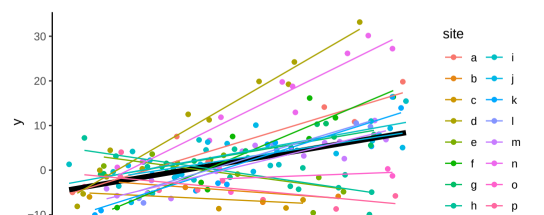
U

Ux

sitea siteb sitec sited sitee sitef siteg siteh sitei sitej sitek

Mixed effect model results

- Regression line of x on y is allowed to move up or down around the main regression line for each site (random intercepts)
- Slope of regression line can be more or less steep for each site (random slopes)
- Changes in intercepts and slopes are normally distributed, and in this example, are correlated with each other



Why do we need to do any of this?

"My supervisor told me to just use site as a fixed effect. Why can't I do that?"

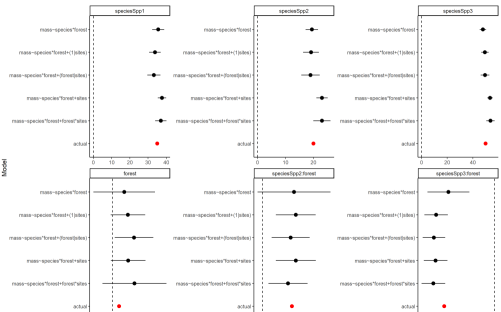
- You can do it this way, but you may encounter the following problems:
 - You lose the partial pooling that occurs in mixed effects models = Worse estimates of site effects!
 - You lose 1 d.f. for each site = Type II error ↑ = You may not find the fixed effect of interest, even if it's there!
 - Sites with low sample sizes may cause your models to break
 - People⁴who have read statistics books published after 1980 may ask questions
- However, if you have a low number of sites (1-10), fixed effects may work better
 - Hard to estimate σ_{site} if number of sites is low
 - If stakes are high, it may be better to be more conservative about site intercepts
 - Easier to interpret (p-values, ANOVA, etc.)

⁴e.g. me

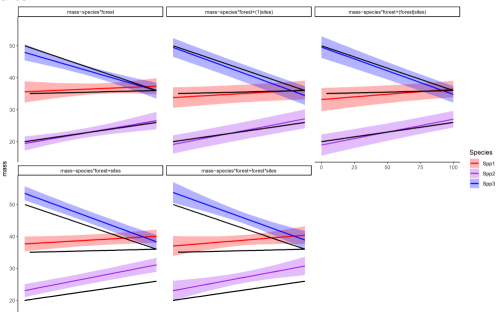
A challenger approaches!

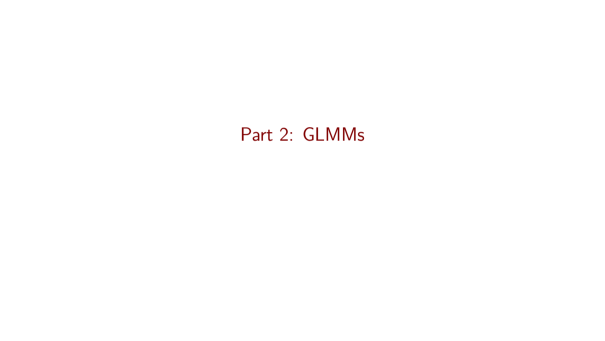
- You're interested in how forest cover influences the mass of three bat species.
 Maybe some of the species do better in forests?
- You've weighed a bunch of bats across different forest covers (batMass.csv). However, these were collected across 15 separate sites. Perhaps some of the variation is just caused by the site?
- Fit a mixed effects model with the fixed effects you're interested in (forest cover, species), and include site as a random effect (intercept or slope)
- Your supervisor doesn't like hierarchical models, and tells you to just use site as another fixed term in an 1m model. Do you get different results if you use their approach?

Results:



Results:





Motivation

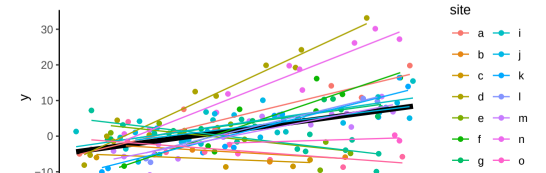
- How do I check if model results are valid?
 - Residual checks
 - Hypothesis testing

Mixed effect model example

Let's go back to our earlier example:

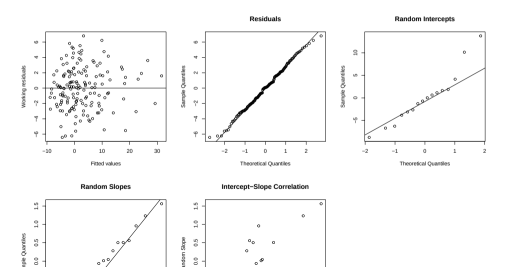
 $lmer(y \sim x + (x|site), data = dat)$

- We're interested in predicting *y* using *x* (fixed effects)
- Data was collected at a number of sites, which may affect y
- Effect of each site is normally distributed (random intercept)
- Effect of site on slope of x is normally distributed (random slope)



Validation

 Similar to linear models, but we also check whether the random intercepts are normally distributed



Hypothesis testing

Is this fixed effect important? (e.g. ANOVA)

- Use likelihood-based test via drop1 (likelihood ratio test, AIC)
- Be careful to fit model with REML = FALSE!

```
lmm1 <- update(lmm1,REML=FALSE) #Refit model using ML rather than REML
drop1(lmm1,test='Chisq') #x has a strong effect</pre>
```

Hypothesis testing (cont.)

How do I know this effect is different from x?

Use Wald Z-test (2-sided p-value from Z-test)

```
lmm1 <- update(lmm1,REML=TRUE) #Reset to REML
meanEst <- fixef(lmm1)[2] #Get mean
seEst <- sqrt(vcov(lmm1)[2,2]) #Get standard error
(1-pnorm(meanEst/seEst,0,1))*2 #p-value from 2-sided Z-test</pre>
```

```
## x
## 0.004659069
```

• glht from library(multcomp) works with lmer models if you are comparing between coefficients (e.g. "Is treatment A different from B and C?")

What if my response variable is non-normal?

Linear model (LM)

$$\hat{y} = X\beta$$

 $y \sim Normal(\hat{y}, \sigma)$

Linear mixed effects model (LMM)

$$\hat{y} = X\beta + U\zeta$$
 $y \sim Normal(\hat{y}, \sigma)$
 $\zeta \sim Normal(0, \sigma_{site})$

• Generalized linear model (GLM)

$$logit(\hat{oldsymbol{\phi}}) = oldsymbol{X}eta \ y \sim Binomial(\hat{oldsymbol{\phi}})$$

 Generalized linear mixed effects model (GLMM)

$$logit(\hat{\phi}) = X\beta + U\zeta$$
 $y \sim Binomial(\hat{\phi})$
 $\zeta \sim Normal(0, \sigma_{site})$

How do I fit GLMMs?

 glmer and glmer.nb from library(lme4) work for Binomial, Poisson, and Negative Binomial data

```
library(lme4)
glmm1 <- glmer.nb(y~x+(x|site),data=dat2) #Negative binomial GLMM
summary(glmm1) #glmer.nb takes a long time to run
```

- glmmTMB from library(glmmTMB) works for those above, *plus* a bunch of others
 - e.g. Zero-inflation, Beta-binomial, Spatial Models

```
library(glmmTMB)
glmm2 <- glmmTMB(y~x+(x|site),data=dat2,family=nbinom2())
summary(glmm2) #Similar results, but quicker</pre>
```

Fitting GLMMs - glmer.nb

```
## Generalized linear mixed model fit by maximum likelihood (Laplace
    Approximation) [glmerMod]
## Family: Negative Binomial(5.1294) ( log )
## Formula: y ~ x + (x | site)
     Data: dat2
       AIC
               BIC logLik deviance df.resid
##
     627.8
             646.3 -307.9 615.8
                                      154
##
## Scaled residuals:
      Min
          1Q Median
                             30
                                    Max
## -1.3745 -0.7098 -0.3946 0.5108 2.5367
##
## Random effects:
## Groups Name
                    Variance Std.Dev. Corr
## site (Intercept) 1.43500 1.1979
                     0.02878 0.1697 0.92
          v
## Number of obs: 160, groups: site, 16
##
## Fixed effects:
              Estimate Std. Error z value Pr(>|z|)
##
## (Intercept) 0.32746
                         0.32166
                                  1.018 0.3087
## v
              0.10830
                       0.04681
                                  2.314
                                         0.0207 *
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
##
## Correlation of Fixed Effects:
## (Intr)
## x 0.799
```

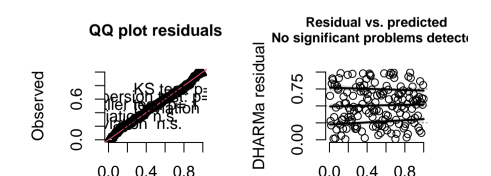
Fitting GLMMs - glmmTMB

```
## Family: nbinom2 (log)
## Formula:
                   v ~ x + (x | site)
## Data: dat2
       AIC
               BIC
                    logLik deviance df.resid
     627.8
            646.2 -307.9
                               615.8
                                         154
##
##
## Random effects:
##
## Conditional model:
## Groups Name
                     Variance Std.Dev. Corr
   site (Intercept) 1.43543 1.1981
                     0.02892 0.1701 0.92
## Number of obs: 160, groups: site, 16
##
## Dispersion parameter for nbinom2 family (): 5.12
##
## Conditional model:
             Estimate Std. Error z value Pr(>|z|)
## (Intercept) 0.34132 0.32172 1.061 0.2887
## x
              0.11026
                       0.04697
                                   2.348 0.0189 *
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
```

Residual checks on glmmTMB models

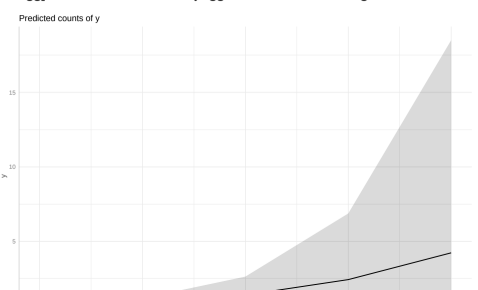
- Extract residuals and make your own plots, or use simulateResiduals from library(DHARMa) (see here)
- DHARMa also has useful functions for checking overdispersion and zero-inflation (found here)

DHARMa residual



Partial residual plots for glmmTMB models

• ggpredict() from library(ggeffects) works with glmmTMB models



Part 2: Model selection and inference

Model selection

How many terms should be in my model?

- Same principle as in regular linear models: what do you think the process is?
 - Just because a term is "not significant" doesn't mean it should be dropped out!
 - Just because a term is "significant" doesn't mean it should be left in!
 - Avoid selecting models based on R^2 or AIC alone. Avoid stargazing⁵ (hunting around for "better" p-values or AIC scores)
- To test whether terms are important in predicting your data (similar to), use likelihood-ratio tests
 - drop1(model,test='Chisq')
 - AIC tests usually say the same thing as LR tests

⁵"My God, it's full of stars!" -2001, A Space Odyssey