# Week 7: Mixed Models MATH-516 Applied Statistics

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#### Section 1

Course Organization (Update)

#### Content

- Week 1: Intro
  - Project 1: Snow Data
- Week 2: Linear Models Practical Recap
- Week 3: Logistic Regression
  - Project 2: Online Shopping Data
- Week 4: Generalized Linear Models
- Week 5: Poisson Regression
  - Project 3: Premier League Data
- Week 6: more on GLMs
- Week 7: Linear Mixed Models
  - Project 4: U.S. Presidential Elections
- Free Week: Easter Holidays
- Week 8: Linear Mixed and Multilevel Models

# Content (cont.)

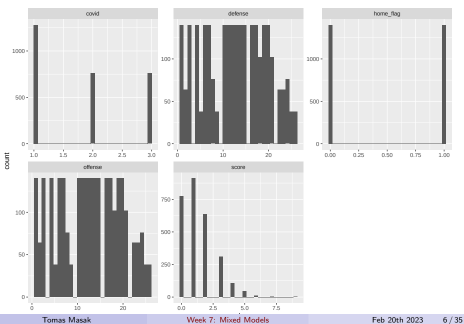
- Week 9: Time Series
  - Project 5: Global Warming
- Week 10: Time Series Regression
- Week 11: Functional Data Analysis
  - Project 7: First Wave of Covid in the US
- Week 12: Functional PCA
- Week 13: Statistical Consulting
- Week 14: Oral Exam
  - discussing your submitted projects

**Evaluation**: remains the same as announced on Week 1, with Statistical Consulting replacing Project 7 (active participation + writing up a suggested solution to a presented problem). Work on Projects 6 and 7 can be combined (to be considered a single submission) and will not be examined.

#### Section 2

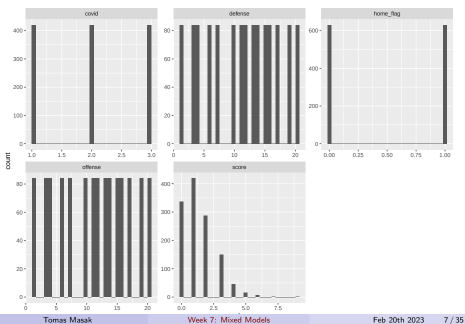
Project 3: Premier League

#### Data Visualized



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### Balanced Data



## Models (Balanced Data)

```
m <- glm(score-covid*home_flag+defense+offense, data=subDat, family="poisson")
msub <- glm(score-home_flag+defense+offense, data=subDat, family="poisson")
anova(m,msub,test="LRT")

## Analysis of Deviance Table
##</pre>
```

```
## Model 1: score - covid * home_flag + defense + offense
## Model 2: score - home_flag + defense + offense
## Resid. Df Resid. Dev Df Deviance Pr(>Chi)
## 1 1226 1352.8
## 2 1230 1357.5 -4 -4.7254 0.3167
```

## covid:home\_flag 5.77 2 0.055895 . ## covid:defense 60.04 35 0.005292 \*\*

## covid:offense ## ---

36.35 35 0.405718

## Signif. codes: 0 '\*\*\*' 0.001 '\*\*' 0.05 '.' 0.1 ' ' 1

```
m <- glm(score~covid*(home flag+defense+offense), data=Data, family="poisson")
msub <- glm(score~home flag+defense+offense, data=Data, family="poisson")
anova(m.msub.test="LRT")
## Analysis of Deviance Table
##
## Model 1: score ~ covid * (home flag + defense + offense)
## Model 2: score ~ home flag + defense + offense
     Resid. Df Resid. Dev Df Deviance Pr(>Chi)
         2674
                2952.6
## 1
         2748 3057.7 -74 -105.14 0.0101 *
## 2
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
library(car)
Anova (m, type=2)
## Analysis of Deviance Table (Type II tests)
##
## Response: score
                  LR Chisq Df Pr(>Chisq)
                    0.91 2 0.633917
## covid
                   16.99 1 3.761e-05 ***
## home_flag
                 184.35 25 < 2.2e-16 ***
## defense
## offense
               345.44 25 < 2.2e-16 ***
```

## Models (Full Data)

## [1] 30

```
minter <- glm(score~covid*(home flag+defense)+offense, data=Data, family="poisson")
msub <- glm(score~home_flag+covid*defense+offense, data=Data, family="poisson")
anova(m.msub.test="LRT")
## Analysis of Deviance Table
##
## Model 1: score ~ covid * (home flag + defense + offense)
## Model 2: score ~ home_flag + covid * defense + offense
    Resid. Df Resid. Dev Df Deviance Pr(>Chi)
          2674
                  2952.6
         2711
               2994.7 -37 -42.102 0.2596
anova(minter.msub.test="LRT")
## Analysis of Deviance Table
##
## Model 1: score ~ covid * (home flag + defense) + offense
## Model 2: score ~ home_flag + covid * defense + offense
    Resid. Df Resid. Dev Df Deviance Pr(>Chi)
## 1
         2709
                  2988 9
## 2
          2711
                  2994.7 -2 -5.7544 0.05629 .
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
sum(is.na(coefficients(m)))
```

 we are on the edge of significance with a model that has too few observations to rely on asymptotics and to estimate all the parameters

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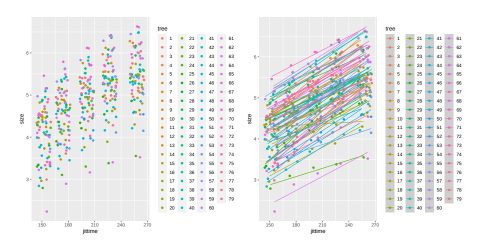
#### Section 3

Example: Tree Growth

- log-size (log-height+2log-diameter) of 79 (Sitka spruce) trees measured repeatedly in about 1-month intervals
  - each tree measured 5-times
  - 54 trees grown in ozone-enriched environment (treat=1) and 25 were control

```
## size time tree treat
## 1 4.51 152 1 ozone
## 2 4.98 174 1 ozone
## 3 5.41 201 1 ozone
## 4 5.90 227 1 ozone
## 5 6.15 258 1 ozone
## 6 4.24 152 2 ozone
```

## Data Displayed



Right: individual line for every tree corresponding to model

y ~ tree\*time

#### Models

```
m1 <- lm(size~(time+I(time^2))*tree,data=Sitka)
m0 <- lm(size~(time+I(time^2)), data=Sitka)
anova(m0,m1)</pre>
```

```
Res.Df RSS Df Sum of Sq F Pr(>F)
1 391 157.107
2 237 6.267 154 150.84 37.043 < 2.2e-16 ***
```

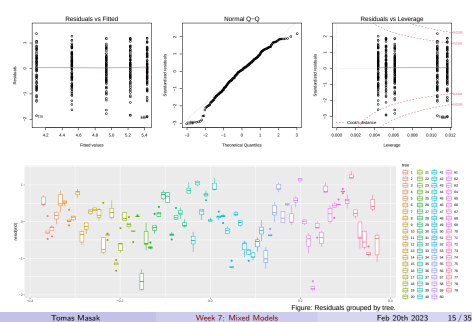
- mO allows for a separate curve for the two treatment groups
- m1 allows for a separate curve for every tree (so m0 is a submodel of m1)

#### Problems:

- m1 cannot be simplified to m0, but the effect of interest (treat)
   cannot be fitted without this simplification, because every single tree is either treatment or control
  - also what if we had low number of observations for some trees and couldn't afford to fit m1?
- but assumptions of m0 are clearly violated

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## Diagnostics



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#### Section 4

#### Linear Mixed Models

#### Definition

- regressions with a large no. of coefficients some of which are themselves being modelled
- extend the linear model

$$Y = \mathbf{X}\beta + \epsilon, \quad \epsilon \sim \mathcal{N}_N(0, \sigma^2 \mathbf{I}_N)$$

to the linear mixed model

$$Y = \mathbf{X}\beta + \mathbf{Z}b + \epsilon, \quad b \sim \mathcal{N}_q(0, \mathbf{C}), \quad \epsilon \sim \mathcal{N}_n(0, \sigma^2 \mathbf{I}_n)$$

- X and Z are known design matrices
- $\beta \in \mathbb{R}^p$  are fixed (non-random) parameters (effects)
- $b \in \mathbb{R}^q$  are random effects with mean 0 and covariance matrix  ${\bf C}$  independent of  $\epsilon$
- parameters:  $\beta$ , **C** and  $\sigma^2$

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## Fitting the Model (ML method)

- the linear mixed model has its log-likelihood  $\ell(\beta, \mathbf{C}, \sigma^2)$
- if we knew C, we could rewrite the model to

$$\mathbf{Y} = \mathbf{X}\beta + e, \quad e \sim \mathcal{N}_{n}(0, \sigma^{2}\mathbf{W}), \quad \mathbf{W} = \mathbf{I}_{n} + \frac{1}{\sigma^{2}}\mathbf{Z}\mathbf{C}\mathbf{Z}^{\top}$$

then the solution  $\widehat{\beta}_{\bf C}$  and  $\widehat{\sigma}_{\bf C}^2$  would be given explicitly by weighted least squares

- ullet imagine a reparametrization "if we knew  ${f C}/\sigma^2$ " instead
- consider the profile log-likelihood for C:  $\ell_p(C) = \ell(\widehat{\beta}_C, C, \widehat{\sigma}_C^2)$

**Algorithm**: starting from an initial  $\mathbf{C}^{(0)}$  alternate until convergence for  $l=1,2,\ldots$  between

- calculation of  $\widehat{\beta}_{\mathbf{C}^{(l-1)}}$  and  $\widehat{\sigma}_{\mathbf{C}^{(l-1)}}^2$  by weighted least squares
- ② updating  $\mathbf{C}^{(I)} = \arg\min_{\mathbf{C}} \ell_p(\mathbf{C})$  by Newton's method (itself iterative)
  - since Newton works well given good starting values, one rather runs EM
    algorithm for a while (treating b as unobserved data) before switching to
    this scheme

# Fitting the Model (REML method)

- ullet start by integrating out eta from the log-likelihood
  - it actually has a closed form and is equivalent to working with likelihood for  $\mathbf{A}Y$  such that  $\mathbb{E}\mathbf{A}Y=0$ , i.e. it is just a simple transformation of the problem
- use iterative solver
  - closed form for  $\sigma^2$
  - inner iteration for C
- ullet finally obtain eta as with the ML method

#### ML vs. REML:

- REML is often preferred (and set as default) since it can lead to unbiased variance estimators (which ML never does)
- but REML depends on parametrization of the fixed effects
  - if one wants to compare models with different X using likelihood criteria,
     ML needs to be used!
- ML and REML are asymptotically equivalent

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## Example: Tree Growth

Rewrite the model in a form that shows the grouping:

$$Y_k = \mathbf{X}_k \beta + \mathbf{Z}_k b_k + \epsilon, \quad b_k \sim \mathcal{N}_q(0, \mathbf{C}), \quad \epsilon_k \sim \mathcal{N}_{N_k}(0, \sigma^2 \mathbf{I}_{N_k})$$

- $b_i$  and  $\epsilon_i$  are i.i.d. for k = 1, ..., K
- $\{b_i\}_{k=1,\ldots,K} \perp \{\epsilon_k\}_{k=1,\ldots,K}$

Specifically, in the tree growth example m1 can be replaced by:

- *K* is the no. of trees
- $N_k = 5$  (for all k) is the number of measurements per tree

$$Y_k = \begin{pmatrix} Y_{k1} \\ \vdots \\ Y_{k5} \end{pmatrix}$$
  $\mathbf{X}_k = \begin{pmatrix} 1 & t_{k1} & t_{k1}^2 \\ \vdots & \vdots & \\ 1 & t_{k5} & t_{k5}^2 \end{pmatrix} = \mathbf{Z}_k$ 

$$\beta = \begin{pmatrix} \beta_0 \\ \beta_1 \\ \beta_2 \end{pmatrix} \qquad b_k = \begin{pmatrix} b_{k0} \\ b_{k1} \\ b_{k2} \end{pmatrix} \qquad \mathbf{C} = \begin{pmatrix} \sigma_0^2 & \sigma_{01} & \sigma_{02} \\ \sigma_{01} & \sigma_1^2 & \sigma_{12} \\ \sigma_{02} & \sigma_{12} & \sigma_2^2 \end{pmatrix}$$

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## Example & Predictors of Random Effects

Fixed effect model: 
$$\mathbb{E}[Y_{ki} \mid X_{ki} = t_{ki}] = \beta_{k0} + \beta_{k1}t_{k1} + \beta_{k2}t_{ki}^2$$

• or something similar depending on the parametrization of tree, e.g. with contr.sum it would be for all but the last tree:

$$\mathbb{E}[Y_{ki} \mid X_{ki} = t_{ki}] = (\beta_0 + \beta_{k0}) + (\beta_1 + \beta_{k1})t_{ki} + (\beta_2 + \beta_{k2})t_{ki}^2$$

Mixed effect model:

$$\mathbb{E}[Y_{ki} \mid X_{ki} = t_{ki}, b_k] = (\beta_0 + b_{k0}) + (\beta_1 + b_{k1})t_{ki} + (\beta_2 + b_{k2})t_{ki}^2$$

 $\bullet \cos(Y_{ki}, Y_{kj}) = \cos(b_{k0} + b_{k1}t_{ki} + b_{k2}t_{ki}^2, b_{k0} + b_{k1}t_{kj} + b_{k2}t_{kj}^2) \neq 0$ 

In general (no need to read too much into the formula):

$$\widehat{b}_k = \mathbf{C} \mathbf{Z}_k^\top (\mathbf{Z}_k \mathbf{C} \mathbf{Z}_k^\top + \sigma^2 \mathbf{I}_{N_k})^{-1} (Y_k - \mathbf{X}_k^\top \widehat{\beta})$$

- called *predictors* since these are not parameters but random variables, but they are also sort of *shrinkage estimators*, because
- vaguely:  $\hat{b}_{kl}$  is somewhere between 0 and the  $\hat{\beta}_{kl}$  in the contr.sum parametrization above

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## Example: Tree Growth

Consider a simpler model, where only the intercept is random:

$$\mathbb{E}[Y_{ki} \mid X_{ki} = t_{ki}, b_k] = (\beta_0 + b_k) + \beta_{k1}t_{ki}$$

and the corresponding fixed-effect-only model y ~ tree+time.

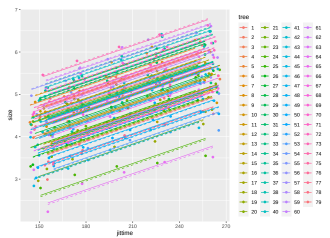


Figure: Tree Growth data and lines by fixed-effect-only model (dashed) and random intercept model (solid).

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## **Uncertainty Quantification**

Let  $\theta \in \mathbb{R}^r$  denote the vector of parameters determining **C**.

**Theorem.** Under validity of the model above and the MLE regularity conditions, we have for  $K \to \infty$ :

• estimators  $\widehat{\beta}, \widehat{\theta}, \widehat{\sigma}^2$  are consistent,

$$\sqrt{K} \begin{pmatrix} \widehat{\beta} - \beta \\ \widehat{\theta} - \theta \\ \widehat{\sigma}^2 - \sigma^2 \end{pmatrix} \to \mathcal{N}_{p+r+1}(0, \mathbf{J}^{-1}), \text{ where } \mathbf{J} = \begin{pmatrix} \mathbf{J}_{\beta} & 0 & 0 \\ 0 & \mathbf{J}_{\theta} & \mathbf{J}_{\theta, \sigma^2} \\ 0 & \mathbf{J}_{\theta, \sigma^2} & \mathbf{J}_{\sigma^2} \end{pmatrix}$$

is the Fisher information matrix,

when  $\widehat{\ell}$  denotes the maximized log-likelihood of the model and  $\widehat{\ell}_0$  denotes the maximized log-likelihood of a submodel then

$$2[\widehat{\ell}-\widehat{\ell}_0] \to \chi_m^2$$

where m is the difference in the no. of parameters between the model and the submodel.

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## Testing for Model Components

- testing fixed effects, i.e.  $H_0: \beta_{p-m+1} = \ldots = \beta_p = 0$  against  $H_1: \neg H_0$ 
  - can be done via LRT due to point 3. of the previous theorem
  - ML needs to be used instead of REML
  - however, p-values tend to be too small, sometimes overstating importance of some effects
- testing random effects, i.e.  $H_0: \theta_{p-m+1} = \ldots = \theta_p = 0$ 
  - usually cannot be done using the previous theorem, because MLE regularity assumptions are typically not met
    - one of the components of  $\theta$  is typically the variance for one of the components of  $b_i$ 's, which lies on the edge of the parameter space
  - it can still be shown that the LR statistic still follows  $\chi^2$ -distribution, but with a smaller no. of degrees of freedom
    - p-values of the test the LRT from point 3. of the previous theorem are too big, sometimes understating importance of some effects

While solutions based on theory exist, a simpler road for us is the parametric bootstrap.

## Example: Tree Growth

```
library(lme4)
# standardize time, otherwise convergence issues
Sitka <- Sitka %>% mutate(time=(time-mean(time))/sd(time))
mm1 <- lmer(size~treat*(time+I(time^2)) + (time+I(time^2)|tree),</pre>
           data=Sitka,REML=F)
mm0 <- lmer(size~time+I(time^2) + (time+I(time^2)|tree),
           data=Sitka,REML=F)
anova(mm0,mm1)
## Data: Sitka
## Models:
## mmO: size ~ time + I(time^2) + (time + I(time^2) | tree)
## mm1: size ~ treat * (time + I(time^2)) + (time + I(time^2) | tree)
      npar AIC BIC logLik deviance Chisq Df Pr(>Chisq)
##
## mm0 10 -111.88 -72.089 65.939 -131.88
## mm1 13 -118.50 -66.772 72.249 -144.50 12.62 3 0.005535 **
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
```

- REML=F because the fixed-effect structure differs between the two models
- treatment seems significant, but... let's do the bootstrap

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## Example: Tree Growth

Bootstrap still rejects, although the p-value is doubled:

```
lrstat <- as.numeric(2*(logLik(mm1)-logLik(mm0)))</pre>
lrstats \leftarrow rep(0,1000)
for(i in 1:1000){
  set.seed(517*i)
  if(i %% 10 ==0) print(i)
  newDat <- Sitka
  newDat$size <- unlist(simulate(mm0))</pre>
  bnull <- lmer(size~time+I(time^2) + (time+I(time^2)|tree),</pre>
                 data=newDat.REML=F)
  balt <- lmer(size~treat*(time+I(time^2)) + (time+I(time^2)|tree),</pre>
                 data=newDat,REML=F)
  lrstats[i] <- as.numeric(2*(logLik(balt)-logLik(bnull)))</pre>
mean(lrstats > lrstat)
```

[1] 0.011

So we can finally conclude that ozone treatment matters :)

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## Diagnostics

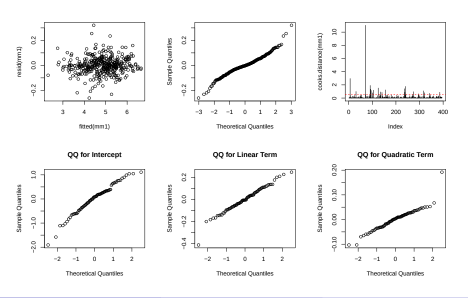
Similar to the standard linear model, with the additional

• check of normality for the random effects

```
plot(fitted(mm1),resid(mm1)) # the only thing `plot(mm1)` gives
qqnorm(resid(mm1),main="") # a bit heavy tails
plot(cooks.distance(mm1),type="h") # the old ROTs not useful here
abline(h=3*mean(cooks.distance(mm1)),col="red",lty=2) # another ROT
qqnorm(ranef(mm1)$tree[,1], main="QQ for Intercept")
qqnorm(ranef(mm1)$tree[,2], main="QQ for Linear Term")
qqnorm(ranef(mm1)$tree[,3], main="QQ for Quadratic Term")
# this is only checking marginals; multivariate GoF tests are tricky
```

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## Diagnostics



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## summary(mm1)

```
Random effects:
```

```
Groups Name Variance Std.Dev. Corr
tree (Intercept) 0.356490 0.59707
time 0.013509 0.11623 0.04
I(time^2) 0.002593 0.05092 0.12 -0.71
Residual 0.008096 0.08998
Number of obs: 395, groups: tree, 79
```

#### Fixed effects:

```
Estimate Std. Error t value (Intercept) 5.093296 0.120072 42.419 treatozone -0.201347 0.145231 -1.386 time 0.546397 0.024638 22.177 I(time^2) -0.108450 0.014041 -7.724 treatozone:time -0.079045 0.029800 -2.652 treatozone:I(time^2) -0.009835 0.016984 -0.579
```

• the random quadratic term is very significant (LR test's p-value  $10^{-8}$  even though understate), but of a low variance and high correlation . . . problems

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#### Section 5

#### Multilevel Models

## Grouping

There can be more than one type of grouping:

$$Y_{kl} = \mathbf{X}_{kl}\beta + \mathbf{Z}_{k,l}^{(1)}b_k + \mathbf{Z}_{l,k}^{(2)}b_l + \epsilon_{kl}$$

- $Y_{kl}$  in  $\mathbb{R}^{N_{kl}}$  is the vector of individual observations belonging to
  - k-th level of grouping (1)
  - *I*-th level of grouping (2)
- e.g. if in the tree growth example we needed to group not only by tree but also by time
  - all *i*-th observations per tree could be correlated
  - e.g. if a different person would measure tree size for different times *i*, but the same person for a single *i*

Or groupings can be nested:

$$Y_{kl} = \mathbf{X}_{kl}\beta + \mathbf{Z}_{k,l}b_k + \mathbf{Z}_{kl}b_{kl} + \epsilon_{kl}$$

- these are called multilevel models:
  - k = 1, ..., K is the first-level grouping (e.g. school)
  - l = 1, ..., L is the second-level grouping (e.g. class)
  - $Y_{kl} \in \mathbb{R}^{N_{kl}}$  is the vector of individual observations (e.g. students)

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# R syntax (lme4 package)

formula	meaning
(1 group)	random group intercept
(x group) = (1+x group)	random slope of x within group with correlated intercept
(0+x group) = (-1+x group)	random slope of x within group: no variation in intercept
$(1 group) + (\theta+x group)$	uncorrelated random intercept and random slope within group
<pre>(1 site/block) = (1 site)+(1 site:block)</pre>	intercept varying among sites and among blocks within sites (nested random effects)
<pre>site+(1 site:block)</pre>	fixed effect of sites plus random variation in intercept among blocks within sites
<pre>(x site/block) = (x site)+(x site:block) =   (1 + x site)+(1+x site:block)</pre>	slope and intercept varying among sites and among blocks within sites
(x1 site)+(x2 block)	two different effects, varying at different levels
<pre>x*site+(x site:block)</pre>	fixed effect variation of slope and intercept varying among sites and random variation of slope and intercept among blocks within sites
(1 group1)+(1 group2)	intercept varying among crossed random effects (e.g. site, year)

source: link

- the linear mixed model generalizes to the GLMM in the same way that the standard linear model generalizes to the GLM
- GLMMs are not as useful, because  $\beta$ 's can only be interpreted as the population-average effects if the link g is linear:

$$\mathbb{E}[Y_n \mid X_n] = \mathbb{E}g^{-1}(X_n^\top \beta + Z_n \top b_n)$$

- cannot go inside  $g^{-1}$  with the expectation unless it is linear, i.e. cannot get rid of  $b_n$  unless  $g^{-1}$  is linear
- only Gaussian models have the linear link as the canonical link
- if one does not use a canonical link, issues may arise
  - numerical convergence issues
  - $\bullet$  calculated estimators are not guaranteed to be MLEs  $\Rightarrow$  theory does not work, etc.

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### Section 6

Project 4

#### Data

- data on U.S. Presidential elections between 1948 and 1992
- two-party vote, i.e. any other candidate than the Democratic and the Republican parties disregarded
- the data are already heavily pre-processed with many derived variables
  - details below
  - cotains polls, etc.

In the 90s, there was a general belief that U.S. presidential elections are easy to understand using linear models and the kind of data we have.

#### **Goal of the project**: Endorse or refute the general belief by:

- start by exploring the data and building a linear model
- consider whether observations in a given a year and/or in a given region within year are correlated
- motivate using mixed models and build one

More detailed description of data and tasks can be found in ./Misc/Projec-4\_assignment.

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