Using lme4: Mixed-Effects Modeling in R

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Outline

**Organizing and plotting data; simple, scalar random effects**

- Models for longitudinal data
- Interactions of grouping factors and other covariates
- Evaluating the log-likelihood
- Generalized Linear Mixed Models
- Item Response Models as GLMMs
- Nonlinear Mixed Models
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Web sites associated with the workshop

www.stat.wisc.edu/~bates/UseR2008  Materials for the course
www.R-project.org  Main web site for the R Project
cran.R-project.org  Comprehensive R Archive Network primary site
cran.us.R-project.org  Main U.S. mirror for CRAN
R-forge.R-project.org  R-Forge, development site for many public R packages. This is also the URL of the repository for installing the development versions of the lme4 and Matrix packages, if you are so inclined.
lme4.R-forge.R-project.org  development site for the lme4 package
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Organizing data in R

- Standard rectangular data sets (columns are variables, rows are observations) are stored in *R* as *data frames*.
- The columns can be *numeric* variables (e.g. measurements or counts) or *factor* variables (categorical data) or *ordered* factor variables. These types are called the *class* of the variable.
- The *str* function provides a concise description of the structure of a data set (or any other class of object in R). The *summary* function summarizes each variable according to its class. Both are highly recommended for routine use.
- Entering just the name of the data frame causes it to be printed. For large data frames use the *head* and *tail* functions to view the first few or last few rows.
R packages

- Packages incorporate functions, data and documentation.
- You can produce packages for private or in-house use or you can contribute your package to the Comprehensive R Archive Network (CRAN), http://cran.us.R-project.org
- We will be using the *lme4* package from CRAN. Install it from the *Packages* menu item or with
  > install.packages("lme4")
- You only need to install a package once. If a new version becomes available you can update (see the menu item).
- To use a package in an R session you attach it using
  > require(lme4)
  or
  > library(lme4)
  (This usage causes widespread confusion of the terms “package” and “library”.)
Accessing documentation

- To be added to CRAN, a package must pass a series of quality control checks. In particular, all functions and data sets must be documented. Examples and tests can also be included.

- The `data` function provides names and brief descriptions of the data sets in a package.

```r
> data(package = "lme4")
```

Data sets in package ’lme4’:

- **Dyestuff**  
  Yield of dyestuff by batch
- **Dyestuff2**  
  Yield of dyestuff by batch
- **Pastes**  
  Paste strength by batch and cask
- **Penicillin**  
  Variation in penicillin testing
- **cake**  
  Breakage angle of chocolate cakes
- **cbpp**  
  Contagious bovine pleuropneumonia
- **sleepstudy**  
  Reaction times in a sleep deprivation study

- Use `?` followed by the name of a function or data set to view its documentation. If the documentation contains an example section, you can execute it with the `example` function.
Lattice graphics

- One of the strengths of R is its graphics capabilities.
- There are several styles of graphics in R. The style in Deepayan Sarkar’s *lattice* package is well-suited to the type of data we will be discussing.
- I will not show every piece of code used to produce the data graphics. The code is available in the script files for the slides (and sometimes in the example sections of the data set’s documentation).
The Dyestuff data set

- The Dyestuff data are a balanced one-way classification of the Yield of dyestuff from samples produced from six Batches of an intermediate product. See ?Dyestuff.

```r
> str(Dyestuff)
'data.frame': 30 obs. of 2 variables:
$ Batch: Factor w/ 6 levels "A","B","C","D",..: 1 1 1 1 1 2 2 2 2 2 ...
$ Yield: num 1545 1440 1440 1520 1580 ...

> summary(Dyestuff)
Batch      Yield
 A:5  Min.  :1440
 B:5  1st Qu.:1469
 C:5  Median :1530
 D:5  Mean   :1528
 E:5  3rd Qu.:1575
 F:5  Max.   :1635
```
The effect of the batches

- To emphasize that Batch is categorical, we use letters instead of numbers to designate the levels.
- Because there is no inherent ordering of the levels of Batch, we will reorder the levels if, say, doing so can make a plot more informative.
- The particular batches observed are just a selection of the possible batches and are entirely used up during the course of the experiment.
- It is not particularly important to estimate and compare yields from these batches. Instead we wish to estimate the variability in yields due to batch-to-batch variability.
- The Batch factor will be used in random-effects terms in models that we fit.
• The line joins the mean yields of the six batches, which have been reordered by increasing mean yield.
• The vertical positions are jittered slightly to reduce overplotting. The lowest yield for batch A was observed on two distinct preparations from that batch.
A mixed-effects model for the dyestuff yield

> fm1 <- lmer(Yield ~ 1 + (1 | Batch), Dyestuff)
> print(fm1)

Linear mixed model fit by REML
Formula: Yield ~ 1 + (1 | Batch)
Data: Dyestuff
AIC BIC logLik deviance REMLdev
 325.7 329.9 -159.8 327.4 319.7
Random effects:
Groups   Name     Variance  Std.Dev.
Batch   (Intercept) 1763.7  41.996
Residual                        2451.3  49.511
Number of obs: 30, groups: Batch, 6
Fixed effects:
   Estimate  Std. Error t value
(Intercept) 1527.50     19.38  78.81

• Fitted model fm1 has one fixed-effect parameter, the mean yield, and one random-effects term, generating a simple, scalar random effect for each level of Batch.
Extracting information from the fitted model

- `fm1` is an object of class "mer" (mixed-effects representation).
- There are many extractor functions that can be applied to such objects.

```r
> fixef(fm1)

(Intercept)      
1527.5

> ranef(fm1, drop = TRUE)

$Batch
          A       B     C       D       E      F
Batch A -17.60596 0.39124 28.56079 -23.08338 56.73033 -44.99302

> fitted(fm1)

[1] 1509.9 1509.9 1509.9 1509.9 1509.9 1527.9 1527.9 1527.9
[9] 1527.9 1527.9 1556.1 1556.1 1556.1 1556.1 1556.1 1504.4
[17] 1504.4 1504.4 1504.4 1504.4 1584.2 1584.2 1584.2 1584.2
[25] 1584.2 1482.5 1482.5 1482.5 1482.5 1482.5 1482.5
```
Definition of linear mixed-effects models

- A mixed-effects model incorporates two vector-valued random variables: the response, $\mathbf{Y}$, and the random effects, $\mathbf{B}$. We observe the value, $\mathbf{y}$, of $\mathbf{Y}$. We do not observe the value of $\mathbf{B}$.
- In a \textit{linear mixed-effects model} the conditional distribution, $\mathbf{Y} \mid \mathbf{B}$, and the marginal distribution, $\mathbf{B}$, are independent, multivariate normal (or “Gaussian”) distributions,

\[
(\mathbf{Y} \mid \mathbf{B} = \mathbf{b}) \sim \mathcal{N}(X\beta + Z\mathbf{b}, \sigma^2 I), \quad \mathbf{B} \sim \mathcal{N}(\mathbf{0}, \sigma^2 \Sigma), \quad (\mathbf{Y} \mid \mathbf{B}) \perp \mathbf{B}.
\]

- The scalar $\sigma$ is the \textit{common scale parameter}; the $p$-dimensional $\beta$ is the \textit{fixed-effects parameter}; the $n \times p \ X$ and the $n \times q \ Z$ are known, fixed \textit{model matrices}; and the $q \times q$ \textit{relative variance-covariance matrix} $\Sigma(\theta)$ is a positive semidefinite, symmetric $q \times q$ matrix that depends on the parameter $\theta$. 

Conditional modes of the random effects

- Technically we do not provide “estimates” of the random effects because they are not parameters.
- One answer to the question, “so what are those numbers anyway?” is that they are BLUPs (Best Linear Unbiased Predictors) but that answer is not informative and the concept does not generalize.
- A better answer is that those values are the conditional means, $\mathbb{E}[\mathbf{B} | \mathbf{Y} = \mathbf{y}]$, evaluated at the estimated parameters. Regrettably, we can only evaluate the conditional means for linear mixed models.
- However, these values are also the conditional modes and that concept does generalize to other types of mixed models.
• For linear mixed models we can evaluate the means and standard deviations of the conditional distributions $\mathcal{B}_j | \mathbf{Y}, j = 1, \ldots, q$. We show these in the form of a 95% prediction interval, with the levels of the grouping factor arranged in increasing order of the conditional mean.

• These are sometimes called “caterpillar plots”.

Caterpillar plot for fm1
Mixed-effects model formulas

- In `lmer` the model is specified by the `formula` argument. As in most R model-fitting functions, this is the first argument.
- The model formula consists of two expressions separated by the `∼` symbol.
- The expression on the left, typically the name of a variable, is evaluated as the response.
- The right-hand side consists of one or more `terms` separated by `+` symbols.
- A random-effects term consists of two expressions separated by the vertical bar, (`|`) symbol (read as “given” or “by”). Typically, such terms are enclosed in parentheses.
- The expression on the right of the `|` is evaluated as a factor, which we call the grouping factor for that term.
Simple, scalar random-effects terms

- In a *simple, scalar* random-effects term, the expression on the left of the ‘|’ is ‘1’. Such a term generates one random effect (i.e. a scalar) for each level of the grouping factor.

- Each random-effects term contributes a set of columns to $Z$. For a simple, scalar r.e. term these are the indicator columns for the levels of the grouping factor. The transpose of the Batch indicators is

```r
> with(Dyestuff, as(Batch, "sparseMatrix"))

6 x 30 sparse Matrix of class "dgCMatrix"
A 1 1 1 1 1 . . . . . . . . . . . . . . . . . . . . . . . . . . B . . . . 1 1 1 1 1 . . . . . . . . . . . . . . . . . . . . C . . . . . . . . . . 1 1 1 1 1 . . . . . . . . . . . . . . . . . . . . D . . . . . . . . . . . . . . . . . . . . 1 1 1 1 1 . . . . . . . . E . . . . . . . . . . . . . . . . . . . . 1 1 1 1 1 . . . . . . . . . . . . . . F . . . . . . . . . . . . . . . . . . . . . . . . . . . . . 1 1 1 1 1
```
Formulation of the marginal variance matrix

- In addition to determining $Z$, the random effects terms determine the form and parameterization of the relative variance-covariance matrix, $\Sigma(\theta)$.
- The parameterization is based on a modified “LDL′′” Cholesky factorization
  \[ \Sigma = TSS′T′ \]
  where $T$ is a $q \times q$ unit lower triangular matrix and $S$ is a $q \times q$ diagonal scale matrix with nonnegative diagonal elements.
- $\Sigma$, $T$ and $S$ are all block-diagonal, with blocks corresponding to the random-effects terms.
- The diagonal block of $T$ for a scalar random effects term is the identity matrix, $I$, and the block in $S$ is a nonnegative multiple of $I$. 
Verbose fitting, extracting $T$ and $S$

- The optional argument `verbose = TRUE` causes `lmer` to print iteration information during the optimization of the parameter estimates.

- The quantity being minimized is the *profiled deviance* of the model. The deviance is negative twice the log-likelihood. It is profiled in the sense that it is a function of $\theta$ only — $\beta$ and $\sigma$ are at their conditional estimates.

- If you want to see exactly how the parameters $\theta$ generate $\Sigma$, use `expand` to obtain a list with components `sigma`, $T$ and $S$. The list also contains a permutation matrix $P$ whose role we will discuss later.

- $T$, $S$ and $\Sigma$ can be very large but are always highly patterned. The `image` function can be used to examine their structure.
Obtain the verbose output for fitting `fm1`

```r
> invisible(update(fm1, verbose = TRUE))
```

<table>
<thead>
<tr>
<th>Iteration</th>
<th>Profiled Deviance</th>
<th>Parameter Vector θ</th>
</tr>
</thead>
<tbody>
<tr>
<td>0</td>
<td>319.76562</td>
<td>0.730297</td>
</tr>
<tr>
<td>1</td>
<td>319.73549</td>
<td>0.962389</td>
</tr>
<tr>
<td>2</td>
<td>319.65735</td>
<td>0.869461</td>
</tr>
<tr>
<td>3</td>
<td>319.65441</td>
<td>0.844025</td>
</tr>
<tr>
<td>4</td>
<td>319.65428</td>
<td>0.848469</td>
</tr>
<tr>
<td>5</td>
<td>319.65428</td>
<td>0.848327</td>
</tr>
<tr>
<td>6</td>
<td>319.65428</td>
<td>0.848324</td>
</tr>
</tbody>
</table>

- The first number on each line is the iteration count — iteration 0 is at the starting value for θ.
- The second number is the profiled deviance — the criterion to be minimized at the estimates.
- The third and subsequent numbers are the parameter vector θ.
Extract $T$ and $S$

- As previously indicated, $T$ and $S$ from $fm1$ are boring.

```r
> (efm1 <- expand(fm1))$S

6 x 6 diagonal matrix of class "ddiMatrix"

[1,] 0.84823 . . . . .
[2,] . 0.84823 . . . .
[3,] . . 0.84823 . . .
[4,] . . . 0.84823 . .
[5,] . . . . 0.84823 .
[6,] . . . . . 0.84823

> efm1$T

6 x 6 sparse Matrix of class "dtCMatrix"

[1,] 1 . . . . .
[2,] . 1 . . . .
[3,] . . 1 . . .
[4,] . . . 1 . .
[5,] . . . . 1 .
[6,] . . . . . 1
```
Reconstructing $\Sigma$

\[
> \ (\text{fm1S} <- \text{tcrossprod}(\text{efm1}$T \%\% \text{efm1}$S))
\]

6 x 6 sparse Matrix of class "dsCMatrix"

\[
\begin{bmatrix}
[1,] & 0.71949 & . & . & . & . \\
[2,] & . & 0.71949 & . & . & . \\
[3,] & . & . & 0.71949 & . & . \\
[4,] & . & . & . & 0.71949 & . \\
[5,] & . & . & . & . & 0.71949 \\
[6,] & . & . & . & . & . & 0.71949
\end{bmatrix}
\]
REML estimates versus ML estimates

- The default parameter estimation criterion for linear mixed models is restricted (or “residual”) maximum likelihood (REML).

- Maximum likelihood (ML) estimates (sometimes called “full maximum likelihood”) can be requested by specifying `REML = FALSE` in the call to `lmer`.

- Generally REML estimates of variance components are preferred. ML estimates are known to be biased. Although REML estimates are not guaranteed to be unbiased, they are usually less biased than ML estimates.

- Roughly, the difference between REML and ML estimates of variance components is comparable to estimating $\sigma^2$ in a fixed-effects regression by $SSR/(n - p)$ versus $SSR/n$, where $SSR$ is the residual sum of squares.

- For a balanced, one-way classification like the Dyestuff data, the REML and ML estimates of the fixed-effects are identical.
Re-fitting the model for ML estimates

```r
> (fm1M <- update(fm1, REML = FALSE))

Linear mixed model fit by maximum likelihood
Formula: Yield ~ 1 + (1 | Batch)
Data: Dyestuff
AIC   BIC   logLik deviance REMLdev
333.3 337.5  -163.7  327.3  319.7
Random effects:
Groups   Name     Variance  Std.Dev.
Batch    (Intercept) 1388.1   37.258
Residual             2451.3   49.511
Number of obs: 30, groups: Batch, 6
Fixed effects:
             Estimate Std. Error t value
(Intercept) 1527.50     17.69  86.33
(The extra parentheses around the assignment cause the value to be printed. Generally the results of assignments are not printed.)
```
Recap of the Dyestuff model

• The model is fit as
  \[
  \text{lmer(formula = Yield \sim 1 + (1 \mid \text{Batch}), data = \text{Dyestuff})}
  \]

• There is one random-effects term, \((1 \mid \text{Batch})\), in the model formula. It is a simple, scalar term for the grouping factor \text{Batch} with \(n_1 = 6\) levels. Thus \(q = 6\).

• The model matrix \(Z\) is the \(30 \times 6\) matrix of indicators of the levels of \text{Batch}.

• The relative variance-covariance matrix, \(\Sigma\), is a nonnegative multiple of the \(6 \times 6\) identity matrix \(I_6\).

• The fixed-effects parameter vector, \(\beta\), is of length \(p = 1\). All the elements of the \(30 \times 1\) model matrix \(X\) are unity.
The Penicillin data (see also the ?Penicillin description)

```r
> str(Penicillin)
'data.frame': 144 obs. of 3 variables:
$ diameter: num 27 23 26 23 23 21 27 23 26 23 ...
$ plate   : Factor w/ 24 levels "a","b","c","d",..: 1 1 1 1 1 1 2 2 2 2 ...
$ sample  : Factor w/ 6 levels "A","B","C","D",..: 1 2 3 4 5 6 1 2 3 4 ...

> xtabs(~sample + plate, Penicillin)

<table>
<thead>
<tr>
<th>plate</th>
<th>sample</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>a b c d e f g h i j k l m n o p q r s t u v w x</td>
</tr>
<tr>
<td>A</td>
<td>1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1</td>
</tr>
<tr>
<td>B</td>
<td>1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1</td>
</tr>
<tr>
<td>C</td>
<td>1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1</td>
</tr>
<tr>
<td>D</td>
<td>1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1</td>
</tr>
<tr>
<td>E</td>
<td>1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1</td>
</tr>
<tr>
<td>F</td>
<td>1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1</td>
</tr>
</tbody>
</table>

• These are measurements of the potency (measured by the diameter of a clear area on a Petri dish) of penicillin samples in a balanced, unreplicated two-way crossed classification with the test medium, plate.
Penicillin data plot

Diameter of growth inhibition zone (mm)
Model with crossed simple random effects for Penicillin

```r
> (fm2 <- lmer(diameter ~ 1 + (1 | plate) + (1 | sample),
+ Penicillin))
```

Linear mixed model fit by REML  
Formula: diameter ~ 1 + (1 | plate) + (1 | sample)  
Data: Penicillin

AIC  BIC logLik deviance REMLdev  
338.9 350.7 -165.4 332.3 330.9

Random effects:  
Groups   Name        Variance  Std.Dev.  
plate    (Intercept) 0.71691   0.84671  
sample   (Intercept) 3.73030   1.93140  
Residual            0.30242   0.54992  
Number of obs: 144, groups: plate, 24; sample, 6

Fixed effects:  
(Intercept)   22.9722   0.8085   28.41
Fixed and random effects for fm2

- The model for the $n = 144$ observations has $p = 1$ fixed-effects parameter and $q = 30$ random effects from $k = 2$ random effects terms in the formula.

```r
> fixef(fm2)

(Intercept) 
  22.972

> ranef(fm2, drop = TRUE)

$plate

<table>
<thead>
<tr>
<th></th>
<th>a</th>
<th>b</th>
<th>c</th>
<th>d</th>
<th>e</th>
<th>f</th>
</tr>
</thead>
<tbody>
<tr>
<td>a</td>
<td>0.804547</td>
<td>0.804547</td>
<td>0.181672</td>
<td>0.337391</td>
<td>0.025953</td>
<td>-0.441203</td>
</tr>
<tr>
<td>b</td>
<td>-1.375516</td>
<td>0.804547</td>
<td>-0.752641</td>
<td>-0.752641</td>
<td>0.960266</td>
<td>0.493109</td>
</tr>
<tr>
<td>c</td>
<td>1.427422</td>
<td>0.493109</td>
<td>0.960266</td>
<td>0.025953</td>
<td>-0.285484</td>
<td>-0.285484</td>
</tr>
<tr>
<td>d</td>
<td>-1.375516</td>
<td>0.960266</td>
<td>-0.908360</td>
<td>-0.285484</td>
<td>-0.596922</td>
<td>-1.219797</td>
</tr>
</tbody>
</table>

$sample

<table>
<thead>
<tr>
<th></th>
<th>A</th>
<th>B</th>
<th>C</th>
<th>D</th>
<th>E</th>
<th>F</th>
</tr>
</thead>
<tbody>
<tr>
<td>A</td>
<td>2.187057</td>
<td>-1.010476</td>
<td>1.937898</td>
<td>-0.096895</td>
<td>-0.013842</td>
<td>-3.003742</td>
</tr>
</tbody>
</table>
Prediction intervals for random effects
Model matrix $Z$ for fm2

- Because the model matrix $Z$ is generated from $k = 2$ simple, scalar random effects terms, it consists of two sets of indicator columns.
- The structure of $Z'$ is shown below. (Generally we will show the transpose of these model matrices - they fit better on slides.)
Models with crossed random effects

• Many people believe that mixed-effects models are equivalent to hierarchical linear models (HLMs) or “multilevel models”. This is not true. The plate and sample factors in fm2 are crossed. They do not represent levels in a hierarchy.

• There is no difficulty in defining and fitting models with crossed random effects (meaning random-effects terms whose grouping factors are crossed). However, fitting models with crossed random effects can be somewhat slower.

• The crucial calculation in each lmer iteration is evaluation of the sparse, lower triangular, Cholesky factor, $L(\theta)$, that satisfies

$$L(\theta)L(\theta)' = P(A(\theta)A(\theta)' + I_q)P'$$

from $A(\theta)' = ZT(\theta)S(\theta)$. Crossing of grouping factors increases the number of nonzeros in $AA'$ and also causes some “fill-in” when creating $L$ from $A$. 
All HLMs are mixed models but not vice-versa

- Even though Raudenbush and Bryk (2002) do discuss models for crossed factors in their HLM book, such models are not hierarchical.
- Experimental situations with crossed random factors, such as "subject" and "stimulus", are common. We can, and should, model such data according to its structure.
- In longitudinal studies of subjects in social contexts (e.g. students in classrooms or in schools) we almost always have partial crossing of the subject and the context factors, meaning that, over the course of the study, a particular student may be observed in more than one class but not all students are observed in all classes. The student and class factors are neither fully crossed nor strictly nested.
- For longitudinal data, "nested" is only important if it means "nested across time". "Nested at a particular time" doesn’t count.
- The lme4 package in R is different from most other software for fitting mixed models in that it handles fully crossed and partial crossed random effects gracefully.
Images of some of the $q \times q$ matrices for fm2

- Because both random-effects terms are scalar terms, $T$ is a block-diagonal matrix of two blocks, both of which are identity matrices. Hence $T = I_q$.
- For this model it is also the case that $P = I_q$.
- $S$ consists of two diagonal blocks, both of which are multiples of an identity matrix. The multiples are different.
Recap of the Penicillin model

- The model formula is
  \[
  \text{diameter} \sim 1 + (1 \mid \text{plate}) + (1 \mid \text{sample})
  \]

- There are two random-effects terms, \((1 \mid \text{plate})\) and \((1 \mid \text{sample})\). Both are simple, scalar \((q_1 = q_2 = 1)\) random effects terms, with \(n_1 = 24\) and \(n_2 = 6\) levels, respectively. Thus \(q = q_1 n_1 + q_2 n_2 = 30\).

- The model matrix \(Z\) is the \(144 \times 30\) matrix created from two sets of indicator columns.

- The relative variance-covariance matrix, \(\Sigma\), is block diagonal in two blocks that are nonnegative multiples of identity matrices. The matrices \(AA'\) and \(L\) show the crossing of the factors. \(L\) has some fill-in relative to \(AA'\).

- The fixed-effects parameter vector, \(\beta\), is of length \(p = 1\). All the elements of the \(144 \times 1\) model matrix \(X\) are unity.
The Pastes data (see also the ?Pastes description)

> str(Pastes)

'data.frame': 60 obs. of 4 variables:
  $ strength: num 62.8 62.6 60.1 62.3 62.7 63.1 60 61.4 57.5 56.9 ...
  $ batch : Factor w/ 10 levels "A","B","C","D",..: 1 1 1 1 1 1 2 2 2 2 ...
  $ cask   : Factor w/ 3 levels "a","b","c": 1 1 2 2 3 3 1 1 2 2 ...
  $ sample : Factor w/ 30 levels "A:a","A:b","A:c",..: 1 1 2 2 3 3 4 4 ...

> xtabs(~batch + sample, Pastes, sparse = TRUE)

10 x 30 sparse Matrix of class "dgCMatrix"
A 2 2 2 . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . .
B . . . 2 2 2 . . . . . . . . . . . . . . . . . . . . . . . . . . . . . .
C . . . . 2 2 2 . . . . . . . . . . . . . . . . . . . . . . . . . . . . . .
D . . . . . . 2 2 2 . . . . . . . . . . . . . . . . . . . . . . . . . . . .
E . . . . . . . . . 2 2 2 . . . . . . . . . . . . . . . . . . . . . . . . .
F . . . . . . . . . . . . 2 2 2 . . . . . . . . . . . . . . . . . . . . . .
G . . . . . . . . . . . . . . . . . . . 2 2 2 . . . . . . . . . . . . . . .
H . . . . . . . . . . . . . . . . . . . . . . . . . . . 2 2 2 . . . . . . .
I . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . .
J . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . .
Structure of the Pastes data

- The `sample` factor is nested within the `batch` factor. Each sample is from one of three casks selected from a particular batch.
- Note that there are 30, not 3, distinct samples.
- We can label the casks as ‘a’, ‘b’ and ‘c’ but then the `cask` factor by itself is meaningless (because cask ‘a’ in batch ‘A’ is unrelated to cask ‘a’ in batches ‘B’, ‘C’, ...). The `cask` factor is only meaningful within a `batch`.
- Only the `batch` and `cask` factors, which are apparently crossed, were present in the original data set. `cask` may be described as being nested within `batch` but that is not reflected in the data. It is *implicitly nested*, not explicitly nested.
- You can save yourself a lot of grief by immediately creating the explicitly nested factor. The recipe is

```r
> Pastes <- within(Pastes, sample <- (batch:cask)[drop = TRUE])
```
Avoid implicitly nested representations

- The `lme4` package allows for very general model specifications. It does not require that factors associated with random effects be hierarchical or “multilevel” factors in the design.

- The same model specification can be used for data with nested or crossed or partially crossed factors. Nesting or crossing is determined from the structure of the factors in the data, not the model specification.

- You can avoid confusion about nested and crossed factors by following one simple rule: ensure that different levels of a factor in the experiment correspond to different labels of the factor in the data.

- Samples were drawn from 30, not 3, distinct casks in this experiment. We should specify models using the `sample` factor with 30 levels, not the `cask` factor with 3 levels.
A model with nested random effects

> (fm3 <- lmer(strength ~ 1 + (1 | batch) + (1 | sample),
+       Pastes))

Linear mixed model fit by REML
Formula: strength ~ 1 + (1 | batch) + (1 | sample)
   Data: Pastes
AIC  BIC logLik deviance REMLdev
 255 263.4 -123.5 248.0 247
Random effects:
 Groups     Name       Variance  Std.Dev.
sample     (Intercept) 8.43378   2.90410
 batch      (Intercept) 1.65692   1.28721
 Residual       0.67801   0.82341
Number of obs: 60, groups: sample, 30; batch, 10
Fixed effects:
          Estimate Std. Error t value
(Intercept)  60.0533     0.6768    88.7
Batch-to-batch variability is low compared to sample-to-sample variability.
Dimensions and relationships in fm3

- There are $n = 60$ observations, $p = 1$ fixed-effects parameter, $k = 2$ simple, scalar random-effects terms ($q_1 = q_2 = 1$) with grouping factors having $n_1 = 30$ and $n_2 = 10$ levels.
- Because both random-effects terms are scalar terms, $T = I_{40}$ and $S$ is block-diagonal in two diagonal blocks of sizes 30 and 10, respectively. $Z$ is generated from two sets of indicators.
Images of some of the $q \times q$ matrices for fm3

- The permutation $P$ has two purposes: reduce fill-in and "post-order" the columns to keep nonzeros near the diagonal.
- In a model with strictly nested grouping factors there will be no fill-in. The permutation $P$ is chosen for post-ordering only.
Eliminate the random-effects term for batch?

- We have seen that there is little batch-to-batch variability beyond that induced by the variability of samples within batches.
- We can fit a reduced model without that term and compare it to the original model.
- Somewhat confusingly, model comparisons from likelihood ratio tests are obtained by calling the `anova` function on the two models. (Put the simpler model first in the call to `anova`.)
- Sometimes likelihood ratio tests can be evaluated using the REML criterion and sometimes they can’t. Instead of learning the rules of when you can and when you can’t, it is easiest always to refit the models with `REML = FALSE` before comparing.
Comparing ML fits of the full and reduced models

> fm3M <- update(fm3, REML = FALSE)
> fm4M <- lmer(strength ~ 1 + (1 | sample), Pastes, + REML = FALSE)
> anova(fm4M, fm3M)

Data: Pastes

Models:
fm4M: strength ~ 1 + (1 | sample)
fm3M: strength ~ 1 + (1 | batch) + (1 | sample)

<table>
<thead>
<tr>
<th>Df</th>
<th>AIC</th>
<th>BIC</th>
<th>logLik</th>
<th>Chisq</th>
<th>Chi Df</th>
<th>Pr(&gt;Chisq)</th>
</tr>
</thead>
<tbody>
<tr>
<td>fm4M 3</td>
<td>254.40</td>
<td>260.69</td>
<td>-124.20</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>fm3M 4</td>
<td>255.99</td>
<td>264.37</td>
<td>-124.00</td>
<td>0.4072</td>
<td>1</td>
<td>0.5234</td>
</tr>
</tbody>
</table>
p-values of LR tests on variance components

- The likelihood ratio is a reasonable criterion for comparing these two models. However, the theory behind using a $\chi^2$ distribution with 1 degree of freedom as a reference distribution for this test statistic does not apply in this case. The null hypothesis is on the boundary of the parameter space.

- Even at the best of times, the p-values for such tests are only approximate because they are based on the asymptotic behavior of the test statistic. To carry the argument further, all results in statistics are based on models and, as George Box famously said, “All models are wrong; some models are useful.”
LR tests on variance components (cont’d)

- In this case the problem with the boundary condition results in a p-value that is larger than it would be if, say, you compared this likelihood ratio to values obtained for data simulated from the null hypothesis model. We say these results are “conservative”.

- As a rule of thumb, the p-value for the $\chi^2$ test on a simple, scalar term is roughly twice as large as it should be.

- In this case, dividing the p-value in half would not affect our conclusion.
Updated model, REML estimates

```r
> (fm4 <- update(fm4M, REML = TRUE))
Linear mixed model fit by REML
Formula: strength ~ 1 + (1 | sample)
   Data: Pastes
AIC  BIC logLik deviance REMLdev
253.6 259.9 -123.8 248.4  247.6
Random effects:
Groups     Name      Variance  Std.Dev.
sample  (Intercept)  9.9767     3.1586
         Residual      0.6780     0.8234
Number of obs: 60, groups: sample, 30
Fixed effects:
     Estimate Std. Error t value
(Intercept)  60.0533    0.5864  102.4
```
Recap of the analysis of the Pastes data

• The data consist of $n = 60$ observations on $q_1 = 30$ samples nested within $q_2 = 10$ batches.
• The data are labelled with a cask factor with 3 levels but that is an implicitly nested factor. Create the explicit factor sample and ignore cask from then on.
• Specification of a model for nested factors is exactly the same as specification of a model with crossed or partially crossed factors — provided that you avoid using implicitly nested factors.
• In this case the batch factor was inert — it did not “explain” substantial variability in addition to that attributed to the sample factor. We therefore prefer the simpler model.
• At the risk of “beating a dead horse”, notice that, if we had used the cask factor in some way, we would still need to create a factor like sample to be able to reduce the model. The cask factor is only meaningful within batch.
Recap of simple, scalar random-effects terms

- For the `lmer` function (and also for `glmer` and `nlmer`) a simple, scalar random effects term is of the form $(1|F)$.
- The number of random effects generated by the $i$th such term is the number of levels, $n_i$, of $F$ (after dropping “unused” levels — those that do not occur in the data. The idea of having such levels is not as peculiar as it may seem if, say, you are fitting a model to a subset of the original data.)
- Such a term contributes $n_i$ columns to $Z$. These columns are the indicator columns of the grouping factor.
- Such a term contributes a diagonal block $I_{n_i}$ to $T$. If all random effects terms are scalar terms then $T = I$.
- Such a term contributes a diagonal block $c_i I_{n_i}$ to $S$. The multipliers $c_i$ can be different for different terms. The term contributes exactly one element (which is $c_i$) to $\theta$. 
This is all very nice, but . . .

- These methods are interesting but the results are not really new. Similar results are quoted in *Statistical Methods in Research and Production*, which is a very old book.
- The approach described in that book is actually quite sophisticated, especially when you consider that the methods described there, based on observed and expected mean squares, are for hand calculation — in pre-calculator days!
- Why go to all the trouble of working with sparse matrices and all that if you could get the same results with paper and pencil? The one-word answer is *balance*.
- Those methods depend on the data being balanced. The design must be completely balanced and the resulting data must also be completely balanced.
- Balance is fragile. Even if the design is balanced, a single missing or questionable observation destroys the balance. Observational studies (as opposed to, say, laboratory experiments) cannot be expected to yield balanced data sets.
A large observational data set

- A large U.S. university (not mine) provided data on the grade point score (`gr.pt`) by student (`id`), instructor (`instr`) and department (`dept`) from a 10 year period. I regret that I cannot make these data available to others.

- These factors are unbalanced and partially crossed.

```
> str(anon.grades.df)
'data.frame': 1721024 obs. of 9 variables:
$ instr : Factor w/ 7964 levels "10000","10001",..: 1 1 1 1 1 1 1 1 1 1 ...
$ dept : Factor w/ 106 levels "AERO","AFAM",..: 43 43 43 43 43 43 43 43 43 43 ...
$ id : Factor w/ 54711 levels "900000001","900000002",..: 12152 1405 23882 18875 18294 20922 4150 13540 5499 6425 ...
$ nclass : num 40 29 33 13 47 49 37 14 21 20 ...
$ vgpa : num NA NA NA NA NA NA NA NA NA ...
$ rawai : num 2.88 -1.15 -0.08 -1.94 3.00 ...
$ gr.pt : num 4 1.7 2 0 3.7 1.7 2 4 2 2.7 ...
$ section : Factor w/ 70366 levels "19959 AERO0011A001",..: 18417 18417 ...
$ semester: num 19989 19989 19989 19989 19989 19972 ...
```
A preliminary model

Linear mixed model fit by REML
Formula: gr.pt ~ (1 | id) + (1 | instr) + (1 | dept)
Data: anon.grades.df

AIC BIC logLik deviance REMLdev
3447389 3447451 -1723690 3447374 3447379

Random effects:
Groups   Name     Variance  Std.Dev.
id       (Intercept) 0.3085  0.555
instr    (Intercept) 0.0795  0.282
dept     (Intercept) 0.0909  0.301
Residual                  0.4037  0.635

Number of obs: 1685394, groups: id, 54711; instr, 7915; dept, 102

Fixed effects:

  Estimate  Std. Error    t value
(Intercept) 3.1996       0.0314      102
Comments on the model fit

- \( n = 1685394, \ p = 1, \ k = 3, \ n_1 = 54711, \ n_2 = 7915, \ n_3 = 102, \ q_1 = q_2 = q_3 = 1, \ q = 62728 \)
- This model is sometimes called the “unconditional” model in that it does not incorporate covariates beyond the grouping factors.
- It takes less than an hour to fit an ”unconditional” model with random effects for student (\textit{id}), instructor (\textit{inst}) and department (\textit{dept}) to these data.
- Naturally, this is just the first step. We want to look at possible time trends and the possible influences of the covariates.
- This is an example of what “large” and “unbalanced” mean today. The size of the data sets and the complexity of the models in mixed modeling can be formidable.
Outline

Organizing and plotting data; simple, scalar random effects

Models for longitudinal data

Interactions of grouping factors and other covariates

Evaluating the log-likelihood

Generalized Linear Mixed Models

Item Response Models as GLMMs

Nonlinear Mixed Models
Simple longitudinal data

- **Repeated measures** data consist of measurements of a response (and, perhaps, some covariates) on several experimental (or observational) units.
- Frequently the experimental (observational) unit is Subject and we will refer to these units as “subjects”. However, the methods described here are not restricted to data on human subjects.
- **Longitudinal** data are repeated measures data in which the observations are taken over time.
- We wish to characterize the response over time within subjects and the variation in the time trends between subjects.
- Frequently we are not as interested in comparing the particular subjects in the study as much as we are interested in modeling the variability in the population from which the subjects were chosen.
Sleep deprivation data

- This laboratory experiment measured the effect of sleep deprivation on cognitive performance.
- There were 18 subjects, chosen from the population of interest (long-distance truck drivers), in the 10 day trial. These subjects were restricted to 3 hours sleep per night during the trial.
- On each day of the trial each subject’s reaction time was measured. The reaction time shown here is the average of several measurements.
- These data are *balanced* in that each subject is measured the same number of times and on the same occasions.
Reaction time versus days by subject
Comments on the sleep data plot

- The plot is a “trellis” or “lattice” plot where the data for each subject are presented in a separate panel. The axes are consistent across panels so we may compare patterns across subjects.

- A reference line fit by simple linear regression to the panel’s data has been added to each panel.

- The aspect ratio of the panels has been adjusted so that a typical reference line lies about 45° on the page. We have the greatest sensitivity in checking for differences in slopes when the lines are near ±45° on the page.

- The panels have been ordered not by subject number (which is essentially a random order) but according to increasing intercept for the simple linear regression. If the slopes and the intercepts are highly correlated we should see a pattern across the panels in the slopes.
Assessing the linear fits

- In most cases a simple linear regression provides an adequate fit to the within-subject data.
- Patterns for some subjects (e.g. 350, 352 and 371) deviate from linearity but the deviations are neither widespread nor consistent in form.
- There is considerable variation in the intercept (estimated reaction time without sleep deprivation) across subjects – 200 ms. up to 300 ms. – and in the slope (increase in reaction time per day of sleep deprivation) – 0 ms./day up to 20 ms./day.
- We can examine this variation further by plotting confidence intervals for these intercepts and slopes. Because we use a pooled variance estimate and have balanced data, the intervals have identical widths.
- We again order the subjects by increasing intercept so we can check for relationships between slopes and intercepts.
95% conf int on within-subject intercept and slope

These intervals reinforce our earlier impressions of considerable variability between subjects in both intercept and slope but little evidence of a relationship between intercept and slope.
A preliminary mixed-effects model

- We begin with a linear mixed model in which the fixed effects $[\beta_1, \beta_2]'$ are the representative intercept and slope for the population and the random effects $b_i = [b_{i1}, b_{i2}]', i = 1, \ldots, 18$ are the deviations in intercept and slope associated with subject $i$.

- The random effects vector, $b$, consists of the 18 intercept effects followed by the 18 slope effects.
Fitting the model

> (fm1 <- lmer(Reaction ~ Days + (Days | Subject),
+     sleepstudy))

Linear mixed model fit by REML
Formula: Reaction ~ Days + (Days | Subject)
Data: sleepstudy
AIC  BIC  logLik deviance REMLdev
1756 1775  -871.8  1752  1744
Random effects:
Groups     Name   Variance  Std.Dev.  Corr
Subject   (Intercept)  612.095   24.7405
            Days      35.071    5.9221   0.065
Residual          654.944   25.5919
Number of obs: 180, groups: Subject, 18
Fixed effects:
             Estimate  Std. Error    t value
(Intercept)  251.405     6.825    36.84
Days           10.467     1.546     6.77

Correlation of Fixed Effects:
             (Intr)
Days  -0.138
Terms and matrices

- The term \textit{Days} in the formula generates a model matrix $X$ with two columns, the intercept column and the numeric \textit{Days} column. (The intercept is included unless suppressed.)
- The term \textit{(Days|Subject)} generates a vector-valued random effect (intercept and slope) for each of the 18 levels of the \textit{Subject} factor.
A model with uncorrelated random effects

- The data plots gave little indication of a systematic relationship between a subject’s random effect for slope and his/her random effect for the intercept. Also, the estimated correlation is quite small.

- We should consider a model with uncorrelated random effects. To express this we use two random-effects terms with the same grouping factor and different left-hand sides. In the formula for an \texttt{lmer} model, distinct random effects terms are modeled as being independent. Thus we specify the model with two distinct random effects terms, each of which has \texttt{Subject} as the grouping factor. The model matrix for one term is intercept only (1) and for the other term is the column for \texttt{Days} only, which can be written 0+Days. (The expression \texttt{Days} generates a column for \texttt{Days} and an intercept. To suppress the intercept we add 0+ to the expression; -1 also works.)
A mixed-effects model with independent random effects

Linear mixed model fit by REML
Formula: Reaction ~ Days + (1 | Subject) + (0 + Days | Subject)
Data: sleepstudy

AIC BIC logLik deviance REMLdev
1754 1770 -871.8 1752 1744

Random effects:
Groups   Name     Variance Std.Dev.
Subject  (Intercept)  627.577  25.0515
Subject   Days       35.852   5.9876
Residual             653.594  25.5655

Number of obs: 180, groups: Subject, 18

Fixed effects:

             Estimate Std. Error t value
(Intercept)   251.405     6.885   36.51
Days          10.467      1.559    6.71

Correlation of Fixed Effects:
       (Intr)
Days -0.184
Comparing the models

- Model `fm1` contains model `fm2` in the sense that if the parameter values for model `fm1` were constrained so as to force the correlation, and hence the covariance, to be zero, and the model were re-fit, we would get model `fm2`.

- The value 0, to which the correlation is constrained, is not on the boundary of the allowable parameter values.

- In these circumstances a likelihood ratio test and a reference distribution of a $\chi^2$ on 1 degree of freedom is suitable.

```r
> anova(fm2, fm1)

Data: sleepstudy
Models:
  fm2: Reaction ~ Days + (1 | Subject) + (0 + Days | Subject)
  fm1: Reaction ~ Days + (Days | Subject)

Df  AIC      BIC    logLik Chisq Chi Df Pr(>Chisq)
fm2  5 1762.05 1778.01  -876.02          
fm1  6 1763.99 1783.14  -875.99  0.0609 1 0.805
```
Conclusions from the likelihood ratio test

• Because the large p-value indicates that we would not reject $f_{m2}$ in favor of $f_{m1}$, we prefer the more parsimonious $f_{m2}$.

• This conclusion is consistent with the AIC (Akaike’s Information Criterion) and the BIC (Bayesian Information Criterion) values for which “smaller is better”.

• We can also use a Bayesian approach, where we regard the parameters as themselves being random variables, is assessing the values of such parameters. A currently popular Bayesian method is to use sequential sampling from the conditional distribution of subsets of the parameters, given the data and the values of the other parameters. The general technique is called Markov chain Monte Carlo sampling.

• The \texttt{lme4} package has a function called \texttt{mcmcsamp} to evaluate such samples from a fitted model. At present, however, there seem to be a few “infelicities”, as Bill Venables calls them, in this function.
Likelihood ratio tests on variance components

- As for the case of a covariance, we can fit the model with and without the variance component and compare the fit quality.
- As mentioned previously, the likelihood ratio is a reasonable test statistic for the comparison but the “asymptotic” reference distribution of a $\chi^2$ does not apply because the parameter value being tested is on the boundary.
- The p-value computed using the $\chi^2$ reference distribution should be conservative (i.e. greater than the p-value that would be obtained through simulation).

```r
> fm3 <- lmer(Reaction ~ Days + (1 | Subject), sleepstudy)
> anova(fm3, fm2)
```

Data: sleepstudy
Models:

<table>
<thead>
<tr>
<th></th>
<th>Df</th>
<th>AIC</th>
<th>BIC</th>
<th>logLik</th>
<th>Chisq</th>
<th>Chi Df</th>
<th>Pr(&gt;Chisq)</th>
</tr>
</thead>
<tbody>
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<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>fm2</td>
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<td>1778.01</td>
<td>-876.02</td>
<td>42.053</td>
<td>1</td>
<td>8.885e-11</td>
</tr>
</tbody>
</table>
```
Conditional modes of the random effects

```r
> (rr2 <- ranef(fm2))

$Subject

       (Intercept)   Days
 308  1.5138208  9.3232133
 309 -40.3749111 -8.5989182
 310 -39.1816685 -5.3876345
 330  24.5182902 -4.9684963
 331  22.9140342 -3.1938381
 332   9.2219310 -0.3084836
 333  17.1560764 -0.2871973
 334  -7.4515943  1.1159563
 335  0.5774084 -10.9056432
 337  34.7689489  8.6273638
 349 -25.7541538  1.2806475
 350 -13.8642113  6.7561991
 351   4.9156060 -3.0750414
 352  20.9294541  3.5121076
 369   3.2587508  0.8730251
 370 -26.4752093  4.9836364
 371  0.9055256 -1.0052631
 372  12.4219020  1.2583667
```
Scatterplot of the conditional modes
Comparing within-subject coefficients

- For this model we can combine the conditional modes of the random effects and the estimates of the fixed effects to get conditional modes of the within-subject coefficients.

- These conditional modes will be “shrunk” towards the fixed-effects estimates relative to the estimated coefficients from each subject’s data. John Tukey called this “borrowing strength” between subjects.

- Plotting the shrinkage of the within-subject coefficients shows that some of the coefficients are considerably shrunk toward the fixed-effects estimates.

- However, comparing the within-group and mixed model fitted lines shows that large changes in coefficients occur in the noisy data. Precisely estimated within-group coefficients are not changed substantially.
Estimated within-group coefficients and BLUPs

Mixed model

Days
(Intercept)
200
220
240
260
280
0 5 10 15 20

Within–group
Each set of prediction intervals have constant width because of the balance in the experiment.
Conclusions from the example

• Carefully plotting the data is enormously helpful in formulating the model.

• It is relatively easy to fit and evaluate models to data like these, from a balanced designed experiment.

• We consider two models with random effects for the slope and the intercept of the response w.r.t. time by subject. The models differ in whether the (marginal) correlation of the vector of random effects per subject is allowed to be nonzero.

• The “estimates” (actually, the conditional modes) of the random effects can be considered as penalized estimates of these parameters in that they are shrunk towards the origin.

• Most of the prediction intervals for the random effects overlap zero.
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Interactions of covariates and grouping factors

• For longitudinal data, having a random effect for the slope w.r.t. time by subject is reasonably easy to understand.

• Although not generally presented in this way, these random effects are an interaction term between the grouping factor for the random effect (subject) and the time covariate.

• We can also define interactions between discrete covariates in the fixed-effects terms and a random-effects grouping factor. However, there is more than one way to define such an interaction.

• Different ways of expressing such interactions lead to different numbers of random effects.

• Models with interactions defined in different ways have levels of complexity, affecting both their expressive power and the ability to estimate all the parameters in the model.
Machines data

- Milliken and Johnson (1989) provide (probably artificial) data on an experiment to measure productivity according to the machine being used for a particular operation.
- In the experiment, a sample of six different operators used each of the three machines on three occasions — a total of nine runs per operator.
- These three machines were the specific machines of interest and we model their effect as a fixed-effect term.
- The operators represented a sample from the population of potential operators. We model this factor, (Worker), as a random effect.
- This is a replicated “subject/stimulus” design with a fixed set of stimuli that are themselves of interest. (In other situations the stimuli may be a sample from a population of stimuli.)
Comments on the data plot

- There are obvious differences between the scores on different machines.
- It seems likely that Worker will be a significant random effect, especially when considering the low variation within replicates.
- There also appears to be a significant Worker:Machine interaction. Worker 6 has a very different pattern w.r.t. machines than do the others.
- We can approach the interaction in one of two ways: define simple, scalar random effects for Worker and for the Worker:Machine interaction or define vector-valued random effects for Worker.
Random effects for subject and subject/stimulus

```r
> print(fm1 <- lmer(score ~ Machine + (1 | Worker) +
+     (1 | Worker:Machine), Machines), corr = FALSE)

Linear mixed model fit by REML
Formula: score ~ Machine + (1 | Worker) + (1 | Worker:Machine)
Data: Machines
AIC  BIC logLik deviance REMLdev
227.7 239.6 -107.8 225.5 215.7
Random effects:
Groups Name Variance Std.Dev.
Worker:Machine (Intercept) 13.90963 3.72956
Worker (Intercept) 22.85526 4.78072
Residual 0.92464 0.96158
Number of obs: 54, groups: Worker:Machine, 18; Worker, 6
Fixed effects:
  Estimate Std. Error  t value
(Intercept)   52.356     2.486   21.062
MachineB       7.967     2.177    3.659
MachineC   13.917     2.177    6.393
```
Vector-valued random effects by subject

> print(fm2 <- lmer(score ~ Machine + (0 + Machine | Worker), Machines), corr = FALSE)

Linear mixed model fit by REML
Formula: score ~ Machine + (0 + Machine | Worker)
   Data: Machines

      AIC   BIC logLik deviance REMLdev
228.3 248.2 -104.2  216.6   208.3

Random effects:
Groups   Name    Variance  Std.Dev.  Corr
Worker  MachineA  16.64098  4.07934
        MachineB  74.39564  8.62529  0.803
        MachineC  19.26648  4.38936  0.623  0.771
Residual        0.92463  0.96158

Number of obs: 54, groups: Worker, 6

Fixed effects:

       Estimate Std. Error  t value
(Intercept)    52.356      1.681   31.150
MachineB        7.967      2.421    3.291
MachineC      13.917      1.540    9.037
Comparing the model fits

- Although not obvious from the specifications, the model fits are nested. If the variance-covariance matrix for the vector-valued random effects has a special form, called *compound symmetry*, the model reduces to model $fm1$.

- The p-value from this comparison is borderline significant.

```r
> fm2M <- update(fm2, REML = FALSE)
> fm1M <- update(fm1, REML = FALSE)
> anova(fm2M, fm1M)
```

Data: Machines
Models:

```
fm1M: score ~ Machine + (1 | Worker) + (1 | Worker:Machine)
fm2M: score ~ Machine + (0 + Machine | Worker)
```

```
Df  AIC   BIC  logLik  Chisq Chi Df Pr(>Chisq)
fm1M  6 237.27 249.20  -112.64
fm2M 10 236.42 256.31  -108.21   8.8516 4 0.06492
```
Model comparisons eliminating the unusual combination

- In a case like this we may want to check if a single, unusual combination (Worker 6 using Machine “B”) causes the more complex model to appear necessary. We eliminate that unusual combination.

```r
> Machines1 <- subset(Machines, Worker != "6" | Machine != "B")

> xtabs(~Machine + Worker, Machines1)

Worker
Machine 1 2 3 4 5 6
A 3 3 3 3 3 3
B 3 3 3 3 3 0
C 3 3 3 3 3 3
```
Machines data after eliminating the unusual combination.
Model comparisons without the unusual combination

\[
> \text{fm1aM} \leftarrow \text{lmer}(\text{score} \sim \text{Machine} + (1 \mid \text{Worker}) + (1 \mid \text{Worker}:\text{Machine}), \text{Machines1}, \text{REML} = \text{FALSE})
\]

\[
> \text{fm2aM} \leftarrow \text{lmer}(\text{score} \sim \text{Machine} + (0 + \text{Machine} \mid \text{Worker}), \text{Machines1}, \text{REML} = \text{FALSE})
\]

\[
> \text{anova(fm2aM, fm1aM)}
\]

Data: Machines1
Models:
fm1aM: score ~ Machine + (1 | Worker) + (1 | Worker:Machine)
fm2aM: score ~ Machine + (0 + Machine | Worker)

<table>
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<th>Df</th>
<th>AIC</th>
<th>BIC</th>
<th>logLik</th>
<th>Chisq</th>
<th>Chi Df</th>
<th>Pr(&gt;Chisq)</th>
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<td>220.145</td>
<td>-98.277</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>fm2aM</td>
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<td>227.607</td>
<td>-94.144</td>
<td>8.2655</td>
<td>4</td>
<td>0.08232</td>
</tr>
</tbody>
</table>
Trade-offs when defining interactions

• It is important to realize that estimating scale parameters (i.e. variances and covariances) is considerably more difficult than estimating location parameters (i.e. means or fixed-effects coefficients).

• A vector-valued random effect term having $q_i$ random effects per level of the grouping factor requires $q_i(q_i + 1)/2$ variance-covariance parameters to be estimated. A simple, scalar random effect for the interaction of a “random-effects” factor and a “fixed-effects” factor requires only 1 additional variance-covariance parameter.

• Especially when the “fixed-effects” factor has a moderate to large number of levels, the trade-off in model complexity argues against the vector-valued approach.

• One of the major sources of difficulty in using the lme4 package is the tendency to overspecify the number of random effects per level of a grouping factor.
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Nonlinear Mixed Models
Definition of linear mixed models

- As previously stated, we define a linear mixed model in terms of two random variables: the $n$-dimensional $\mathbf{Y}$ and the $q$-dimensional $\mathbf{B}$
- The probability model specifies the conditional distribution
$$
(\mathbf{Y}|\mathbf{B} = \mathbf{b}) \sim \mathcal{N}(X\beta + Z\mathbf{b}, \sigma^2 I)
$$
and the unconditional distribution
$$
\mathbf{B} \sim \mathcal{N}(0, \sigma^2 \Sigma(\theta)), \quad (\mathbf{Y}|\mathbf{B}) \perp \mathbf{B}
$$
as independent, multivariate Gaussian distributions depending on the parameters $\beta$, $\theta$ and $\sigma$.
- The relative variance-covariance matrix for $\mathbf{B}$, written $\Sigma(\theta)$, can be factored as
$$
\Sigma(\theta) = T(\theta)S(\theta)S(\theta)T(\theta)' = (TS)(TS)'.
$$
We say that the product $T(\theta)S(\theta)$ is a left square-root factor of $\Sigma(\theta)$. 
The conditional distribution, $\mathbf{Y|B}$

- The mean of the conditional distribution, $\mathbf{Y|B}$, is a linear function of $\beta$ and $b$.

$$
\mu_{\mathbf{Y|B}}(b) = \mathbb{E}[\mathbf{Y|B} = b] = \eta = X\beta + Zb
$$

- For generalized linear models we will distinguish between the conditional mean, $\mu_{\mathbf{Y|B}}(b)$, which may be bounded, and the linear predictor, $\eta$, which is always unbounded. For linear mixed models, $\mu_{\mathbf{Y|B}}(b) = \eta$.

- Components of $\mathbf{Y}$ are *conditionally independent*, given $\mathbf{B}$. That is, the conditional distribution, $(\mathbf{Y|B} = b)$, is determined by the (scalar) distribution of each component.

- Hence, the conditional distribution, $(\mathbf{Y|B} = b)$, is completely determined by the conditional mean, $\mu_{\mathbf{Y|B}}$, and the common scale parameter, $\sigma$. 
The unscaled conditional density of \( \mathcal{B}|\mathcal{Y} = y \)

- Because it is \( y \), not \( b \), that we observe, we are interested in evaluating the other conditional distribution, \( (\mathcal{B}|\mathcal{Y} = y) \). We will write its density as \( [\mathcal{B}|\mathcal{Y}](b|y) \) (it is always continuous, even when, as in some GLMMs, \( \mathcal{Y} \) is discrete).
- Given \( y, \theta, \beta \) and, if used, \( \sigma \), we can evaluate \( [\mathcal{B}|\mathcal{Y}](b|y) \), up to a scale factor, as \( [\mathcal{Y}|\mathcal{B}](y|b) [\mathcal{B}](b) \).
- The inverse of the scale factor,

\[
\int_{\mathbb{R}^q} [\mathcal{Y}|\mathcal{B}](y|b) [\mathcal{B}](b) \, db,
\]

is exactly the likelihood, \( L(\theta, \beta, \sigma^2|y) \) (or \( L(\theta, \beta, |y) \) when \( \sigma \) is not used).
The unscaled conditional density of $\mathbf{U} | \mathbf{Y} = \mathbf{y}$

- To simplify the integral defining the likelihood, we change the variable of integration to $\mathbf{u}$, where $\mathbf{U}$ is a vector-valued random variable with unconditional distribution $\mathbf{U} \sim \mathcal{N}(\mathbf{0}, \sigma^2 \mathbf{I}_q)$ (or $\mathbf{U} \sim \mathcal{N}(\mathbf{0}, \mathbf{I}_q)$, when $\sigma$ is not used), and $\mathbf{B} = T(\theta)S(\theta) \mathbf{P}' \mathbf{U}$.

- The linear predictor, $\eta$, which determines the conditional density, $[\mathbf{Y} | \mathbf{U}](\mathbf{y} | \mathbf{u})$, becomes

$$\eta = ZT(\theta)S(\theta)\mathbf{P}'\mathbf{u} + X\beta = A(\theta)'\mathbf{P}'\mathbf{u} + X\beta,$$

where $A(\theta)' = ZT(\theta)S(\theta)$, and likelihood

$$L(\theta, \beta | \mathbf{y}) = \int_{\mathbb{R}^q} [\mathbf{Y} | \mathbf{U}](\mathbf{y} | \mathbf{u}) [\mathbf{U}](\mathbf{u}) \, d\mathbf{u}.$$
Maximizing the unscaled density $U|Y = y$

- In our general strategy for evaluating the likelihood, $L(\theta, \beta, \sigma^2|y)$, we first maximize the unscaled density of $U|Y = y$, w.r.t. $u$.
- Both $[Y|U](y|u)$ and $[U](u)$ are spherical normal densities, which means that the components are independent with constant variance, e.g. $\text{Var}(U) = \sigma^2 I$, (“spherical” because the contours of constant density are spheres).
- That is, probability density is related to the (squared) lengths, $\|y - \mu Y|U\|^2$ and $\|u\|^2$, with the same scale factor, $\sigma^2$.
- The conditional mode of $U|Y$ – the value that maximizes the conditional density (and also the unscaled version) – does not depend on $\sigma^2$.

\[
\tilde{u}(y|\theta, \beta) = \arg \max_u [Y|U](y|u) [U](u) \\
= \arg \min_u \left( \|y - \mu Y|U\|^2 + \|u\|^2 \right)
\]
Solving for the conditional mode

- Incorporating the definition of $\mu_{Y|U}$ provides

$$\|y - \mu_{Y|U}\|^2 = \|y - A'P'u - X\beta\|^2$$

- Recall that $P$ is a permutation matrix. These have the property that $P^{-1} = P'$, allowing us to write

$$\|0 - P'u\|^2 = u'PP'u = u'u = \|u\|^2$$

- Combining these produces

$$\hat{u}(y|\theta, \beta) = \arg\min_u \left\| \begin{bmatrix} y - X\beta \\ 0 \end{bmatrix} - \begin{bmatrix} A' \\ I \end{bmatrix} P'u \right\|^2$$

Hence, $\hat{u}$ satisfies

$$P \left( AA' + I \right) P'\hat{u} = LL'\hat{u} = PA(y - X\beta)$$

where $L(\theta)$ is the sparse left Cholesky factor of $P \left( A(\theta)A(\theta)' + I \right) P'$. 
Evaluating the likelihood - linear mixed models

- Because $\mu_{Y|U}$ depends linearly on both $u$ and $\beta$, the conditional mode $\tilde{u}(\theta)$ and the conditional maximum likelihood estimate, $\hat{\beta}(\theta)$, can be determined simultaneously as the solutions to a penalized least squares problem

\[
\begin{bmatrix}
\tilde{u}(\theta) \\
\hat{\beta}(\theta)
\end{bmatrix} = \arg\min_{u,\beta} \left\| \begin{bmatrix} y \\ 0 \end{bmatrix} - \begin{bmatrix} A'P' & X \\ I & 0 \end{bmatrix} \begin{bmatrix} u \\ \beta \end{bmatrix} \right\|^2
\]

for which the solution satisfies

\[
\begin{bmatrix}
P (AA' + I) P' & PAX \\
X'A'P' & X'X
\end{bmatrix} \begin{bmatrix}
\tilde{u}(\theta) \\
\hat{\beta}(\theta)
\end{bmatrix} = \begin{bmatrix} PAy \\ X'y \end{bmatrix}
\]

- The Cholesky factor of the system matrix for the PLS problem is

\[
\begin{bmatrix}
P (AA' + I) P' & PAX \\
X'A'P' & X'X
\end{bmatrix} = \begin{bmatrix} L & 0 \\ R'^{ZX} & R'_X \\ 0 & R_X \end{bmatrix}
\]

- The dense matrices $R_{ZX}$ and $R_X$ are stored in the $RZX$ and $RX$ slots, respectively.
Special case of linear mixed models (cont’d)

• It is not necessary to solve for \( \tilde{u}(\theta) \) and \( \hat{\beta}(\theta) \). All that is needed for evaluation of the profiled log-likelihood is the penalized residual sum of squares, \( r^2 \), and the determinant

\[
|AA' + I| = |L|^2
\]

• Because \( L \) is triangular, its determinant is simply the product of its diagonal elements.

• Because \( AA' + I \) is positive definite, \( |L|^2 > 0 \).

• The profiled deviance, as a function of \( \theta \) only (\( \beta \) and \( \sigma^2 \) at their conditional estimates), is

\[
d(\theta|y) = \log(|L|^2) + n \left( 1 + \log \left( \frac{2\pi r^2}{n} \right) \right)
\]
REML results

- Although not often derived in this form, Laird and Ware showed that the REML criterion can be derived as the integral of the likelihood w.r.t. $\beta$.

- The same techniques as used to evaluate the integral w.r.t. $b$ can be used to evaluate the integral for the REML criterion. In this case the integral introduces the factor $|R_X|^2$.

- The profiled REML deviance, as a function of $\theta$ only ($\sigma$ at its conditional estimate), is

$$d_R(\theta|y) = \log(|L|^2|R_X|^2) + (n - p) \left(1 + \log \left(\frac{2\pi r^2}{n - p}\right)\right)$$
Recap

- For a linear mixed model, even one with a huge number of observations and random effects like the model for the grade point scores, evaluation of the ML or REML profiled deviance, given a value of $\theta$, is straightforward. It involves updating $T$ and $S$, then updating $A$, $L$, $R_{ZX}$, $R_X$, calculating the penalized residual sum of squares, $r$ and a couple of determinants of triangular matrices.

- The profiled deviance can be optimized as a function of $\theta$ only. The dimension of $\theta$ is usually very small. For the grade point scores there are only three components to $\theta$. 
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Generalized Linear Mixed Models

• When using linear mixed models (LMMs) we assume that the response being modeled is on a continuous scale.

• Sometimes we can bend this assumption a bit if the response is an ordinal response with a moderate to large number of levels. For example, the Scottish secondary school test results were integer values on the scale of 1 to 10.

• However, an LMM is not suitable for modeling a binary response, an ordinal response with few levels or a response that represents a count. For these we use generalized linear mixed models (GLMMs).

• To describe GLMMs we return to the representation of the response as an $n$-dimensional, vector-valued, random variable, $\mathbf{Y}$, and the random effects as a $q$-dimensional, vector-valued, random variable, $\mathbf{B}$. 
Parts of LMMs carried over to GLMMs

- Random variables
  - \( \mathbf{Y} \) the response variable
  - \( \mathbf{B} \) the (possibly correlated) random effects
  - \( \mathbf{U} \) the orthogonal random effects

- Parameters
  - \( \mathbf{\beta} \) - fixed-effects coefficients
  - \( \sigma \) - the common scale parameter (not always used)
  - \( \theta \) - parameters that determine \( \text{Var}(\mathbf{B}) = \sigma^2 (\mathbf{T S})(\mathbf{T S})' \)

- Some matrices
  - \( \mathbf{X} \) the \( n \times p \) model matrix for \( \mathbf{\beta} \)
  - \( \mathbf{Z} \) the \( n \times q \) model matrix for \( \mathbf{b} \)
  - \( \mathbf{P} \) fill-reducing \( q \times q \) permutation (from \( \mathbf{Z} \))
  - \( \mathbf{S}(\theta) \) non-negative \( q \times q \) diagonal scale matrix
  - \( \mathbf{T}(\theta) \) \( q \times q \) unit lower-triangular matrix
  - \( \mathbf{A}(\theta) = (\mathbf{Z T}(\theta)\mathbf{S}(\theta))' \)
The conditional distribution, $\mathbf{Y} | \mathbf{U}$

- For GLMMs, the marginal distribution, $\mathbf{B} \sim \mathcal{N}(\mathbf{0}, \Sigma(\theta))$ is the same as in LMMs except that $\sigma^2$ is omitted. We define $\mathbf{U} \sim (\mathbf{0}, I_q)$ such that $\mathbf{B} = T(\theta)S(\theta)P'\mathbf{U}$.

- For GLMMs we retain some of the properties of the conditional distribution

  $$(\mathbf{Y} | \mathbf{U} = \mathbf{u}) \sim \mathcal{N}(\mu_{\mathbf{Y}|\mathbf{U}}(\mathbf{u}), \sigma^2 I)$$

  where $\mu_{\mathbf{Y}|\mathbf{U}}(\mathbf{u}) = \mathbf{X}\beta + \mathbf{A}'P'\mathbf{u}$

  Specifically
  - The distribution $\mathbf{Y}|\mathbf{U} = \mathbf{u}$ depends on $\mathbf{u}$ only through the conditional mean, $\mu_{\mathbf{Y}|\mathbf{U}}(\mathbf{u})$.
  - Elements of $\mathbf{Y}$ are conditionally independent. That is, the distribution of $\mathbf{Y}|\mathbf{U} = \mathbf{u}$ is completely specified by the univariate, conditional distributions, $\mathbf{Y}_i|\mathbf{U}, i = 1, \ldots, n$.
  - These univariate, conditional distributions all have the same form. They differ only in their means.

- GLMMs differ from LMMs in the form of the univariate, conditional distributions and in how $\mu_{\mathbf{Y}|\mathbf{U}}(\mathbf{u})$ depends on $\mathbf{u}$. 
Some choices of univariate conditional distributions

- Typical choices of univariate conditional distributions are:
  - The *Bernoulli* distribution for binary (0/1) data, which has probability mass function
    \[ p(y|\mu) = \mu^y (1 - \mu)^{1-y}, \quad 0 < \mu < 1, \quad y = 0, 1 \]
  - Several independent binary responses can be represented as a *binomial* response, but only if all the Bernoulli distributions have the same mean.
  - The *Poisson* distribution for count (0, 1, \ldots) data, which has probability mass function
    \[ p(y|\mu) = e^{-\mu} \frac{\mu^y}{y!}, \quad 0 < \mu, \quad y = 0, 1, 2, \ldots \]

- All of these distributions are completely specified by the conditional mean. This is different from the conditional normal (or Gaussian) distribution, which also requires the common scale parameter, \( \sigma \).
The link function, $g$

- When the univariate conditional distributions have constraints on $\mu$, such as $0 < \mu < 1$ (Bernoulli) or $0 < \mu$ (Poisson), we cannot define the conditional mean, $\mu_{Y|U}$, to be equal to the linear predictor, $X\beta + A'P'u$, which is unbounded.

- We choose an invertible, univariate link function, $g$, such that $\eta = g(\mu)$ is unconstrained. The vector-valued link function, $g$, is defined by applying $g$ component-wise.

$$\eta = g(\mu) \quad \text{where} \quad \eta_i = g(\mu_i), \quad i = 1, \ldots, n$$

- We require that $g$ be invertible so that $\mu = g^{-1}(\eta)$ is defined for $-\infty < \eta < \infty$ and is in the appropriate range ($0 < \mu < 1$ for the Bernoulli or $0 < \mu$ for the Poisson). The vector-valued inverse link, $g^{-1}$, is defined component-wise.
“Natural” link functions

• There are many choices of invertible scalar link functions, $g$, that we could use for a given set of constraints.

• For the Bernoulli and Poisson distributions, however, one link function arises naturally from the definition of the probability mass function. (The same is true for a few other, related but less frequently used, distributions, such as the gamma distribution.)

• To derive the natural link, we consider the logarithm of the probability mass function (or, for continuous distributions, the probability density function).

• For distributions in this “exponential” family, the logarithm of the probability mass or density can be written as a sum of terms, some of which depend on the response, $y$, only and some of which depend on the mean, $\mu$, only. However, only one term depends on both $y$ and $\mu$, and this term has the form $y \cdot g(\mu)$, where $g$ is the natural link.
The natural link for the Bernoulli distribution

- The logarithm of the probability mass function is

\[ \log(p(y|\mu)) = \log(1-\mu) + y \log\left(\frac{\mu}{1-\mu}\right), \quad 0 < \mu < 1, \quad y = 0, 1. \]

- Thus, the natural link function is the **logit** link

\[ \eta = g(\mu) = \log\left(\frac{\mu}{1-\mu}\right). \]

- Because \( \mu = P[Y = 1] \), the quantity \( \mu/(1-\mu) \) is the odds ratio (in the range \((0, \infty)\)) and \( g \) is the logarithm of the odds ratio, sometimes called “log odds”.

- The inverse link is

\[ \mu = g^{-1}(\eta) = \frac{e^\eta}{1 + e^\eta} = \frac{1}{1 + e^{-\eta}} \]
Plot of natural link for the Bernoulli distribution
Plot of inverse natural link for the Bernoulli distribution

\[ \mu = \frac{1}{1 + e^{-\eta}} \]

\( \eta \) values range from -5 to 5.
The natural link for the Poisson distribution

- The logarithm of the probability mass is

\[
\log(p(y|\mu)) = \log(y!) - \mu + y \log(\mu)
\]

- Thus, the natural link function for the Poisson is the \textit{log} link

\[
\eta = g(\mu) = \log(\mu)
\]

- The inverse link is

\[
\mu = g^{-1}(\eta) = e^\eta
\]
The natural link related to the variance

- For the natural link function, the derivative of its inverse is the variance of the response.
- For the Bernoulli, the natural link is the logit and the inverse link is $\mu = g^{-1}(\eta) = 1/(1 + e^{-\eta})$. Then
  \[
  \frac{d\mu}{d\eta} = \frac{e^{-\eta}}{(1 + e^{-\eta})^2} = \frac{1}{1 + e^{-\eta}} \frac{e^{-\eta}}{1 + e^{-\eta}} = \mu(1 - \mu) = \text{Var}(Y)
  \]
- For the Poisson, the natural link is the log and the inverse link is $\mu = g^{-1}(\eta) = e^\eta$. Then
  \[
  \frac{d\mu}{d\eta} = e^\eta = \mu = \text{Var}(Y)
  \]
The unscaled conditional density of $\mathbf{U} | \mathbf{Y} = y$

- As in LMMs we evaluate the likelihood of the parameters, given the data, as
  \[ L(\theta, \beta | y) = \int_{\mathbb{R}^q} [\mathbf{Y} | \mathbf{U}](y | u) [\mathbf{U}](u) \, du, \]

- The product $[\mathbf{Y} | \mathbf{U}](y | u)[\mathbf{U}](u)$ is the unscaled (or unnormalized) density of the conditional distribution $\mathbf{U} | \mathbf{Y}$.

- The density $[\mathbf{U}](u)$ is a spherical Gaussian density
  \[ \frac{1}{(2\pi)^{q/2}} e^{-\|u\|^2/2}. \]

- The expression $[\mathbf{Y} | \mathbf{U}](y | u)$ is the value of a probability mass function or a probability density function, depending on whether $\mathbf{Y}_i | \mathbf{U}$ is discrete or continuous.

- The linear predictor is $g(\mu_{\mathbf{Y} | \mathbf{U}}) = \eta = X\beta + A(\theta)'P'u$. Alternatively, we can write the conditional mean of $\mathbf{Y}$, given $\mathbf{U}$, as
  \[ \mu_{\mathbf{Y} | \mathbf{U}}(u) = g^{-1} (X\beta + A(\theta)'P'u) \]
The conditional mode of $\mathbf{U} \mid \mathbf{Y} = \mathbf{y}$

- In general the likelihood, $L(\theta, \beta \mid \mathbf{y})$ does not have a closed form. To approximate this value, we first determine the **conditional mode**

$$\tilde{u}(\mathbf{y} \mid \theta, \beta) = \arg \max_u \mathbb{E}[\mathbf{Y} \mid \mathbf{U}](\mathbf{y} \mid u) \mathbb{E}[\mathbf{U}]$$

using a quadratic approximation to the logarithm of the unscaled conditional density.

- This optimization problem is (relatively) easy because the quadratic approximation to the logarithm of the unscaled conditional density can be written as a penalized, weighted residual sum of squares,

$$\tilde{u}(\mathbf{y} \mid \theta, \beta) = \arg \min_u \left\| \begin{bmatrix} W^{1/2}(\mu)(\mathbf{y} - \mu \mathbb{E}[\mathbf{Y} \mid \mathbf{U}(u))] \\ -u \end{bmatrix} \right\|^2$$

where $W(\mu)$ is the diagonal weights matrix. The weights are the inverses of the variances of the $\mathbf{Y}_i$. 
The PIRLS algorithm

- Parameter estimates for generalized linear models (without random effects) are usually determined by iteratively reweighted least squares (IRLS), an incredibly efficient algorithm. PIRLS is the penalized version. It is iteratively reweighted in the sense that parameter estimates are determined for a fixed weights matrix $W$ then the weights are updated to the current estimates and the process repeated.

- For fixed weights we solve

$$
\min_{\mathbf{u}} \left\| \begin{bmatrix} W^{1/2} (y - \mu | \mathbf{u}(\mathbf{u})) \\ -\mathbf{u} \end{bmatrix} \right\|^2
$$

as a nonlinear least squares problem with update, $\delta_{\mathbf{u}}$, given by

$$
P (AMWMA' + I) P' \delta_{\mathbf{u}} = PAMW (y - \mu) - \mathbf{u}
$$

where $M = d\mu/d\eta$ is the (diagonal) Jacobian matrix. Recall that for the natural link, $M = \text{Var}(\mathbf{y}|\mathbf{u}) = W^{-1}$. 

The Laplace approximation to the deviance

- At convergence, the sparse Cholesky factor, $L$, used to evaluate the update is

$$LL' = P(AMWMA' + I)P'$$

or

$$LL' = P(AMA' + I)P'$$

if we are using the natural link.

- The integrand of the likelihood is approximately a constant times the density of the $\mathcal{N}(\tilde{u}, LL')$ distribution.

- On the deviance scale (negative twice the log-likelihood) this corresponds to

$$d(\beta, \theta|y) = d_g(y, \mu(\tilde{u})) + \|\tilde{u}\|^2 + \log(|L|^2)$$

where $d_g(y, \mu(\tilde{u}))$ is the GLM deviance for $y$ and $\mu$. 
Modifications to the algorithm

- Notice that this deviance depends on the fixed-effects parameters, $\beta$, as well as the variance-component parameters, $\theta$. This is because $\log(|L|^2)$ depends on $\mu_{Y|U}$ and, hence, on $\beta$. For LMMs $\log(|L|^2)$ depends only on $\theta$.

- It is likely that modifying the PIRLS algorithm to optimize simultaneously on $u$ and $\beta$ would result in a value that is very close to the deviance profiled over $\beta$.

- Another approach, which is being implemented as a Google Summer of Code project, is adaptive Gauss-Hermite quadrature (AGQ). This has a similar structure to the Laplace approximation but is based on more evaluations of the unscaled conditional density near the conditional modes. It is only appropriate for models in which the random effects are associated with only one grouping factor.
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Item Response Models

- Models for binary (or ordered categorical) data that are cross-classified according to subject and item are sometimes called *Item Response* or *IRT* (Item Response Theory) models.

- There is a long history of models for such data with many contributors. Only recently have statisticians become aware of this literature and considered how such models could be framed in the context of GLMMs.

- Even when approaching IRT models as GLMMs they were not expressed as GLMMs with crossed random effects, because of software limitations.

- Because `glmer` can fit GLMMs with crossed random effects, we can approach such models as GLMMs with random effects for subject and item.
Data from a study of verbal aggression

- Results on a study of verbal aggression, used as an example through the book *Expanatory Item Response Models*, edited by De Boeck and Wilson (Springer, 2004) are available as the data set VerbAgg, in the “long” format.
- The items correspond to scenarios for which the subject was asked if they would curse, scold or shout.
- The scenarios are classified according to the behavior mode (want versus do) and according to the situation (self-to-blame versus other-to-blame).
- The subjects are classified by sex. Each subject’s score on a separately administered anger index (STAXI) is given.
- The response was recorded on a three-level ordinal scale (“no”, “perhaps” and “yes”). We will consider a dichotomous version, $r_2$. 
Structure of VerbAgg data

- We also check that the item-level covariates and the person-level covariates are consistently defined.

```r
> str(VerbAgg)
'data.frame': 7584 obs. of 9 variables:
$ Anger : int 20 11 17 21 17 21 39 21 24 16 ... 
$ Gender: Factor w/ 2 levels "M","F": 2 2 1 1 1 1 1 1 1 1 ...
$ item  : Factor w/ 24 levels "S1wantcurse",...: 1 1 1 1 1 1 1 1 1 1 ...
$ resp  : Ord.factor w/ 3 levels "no"<"perhaps"<...: 1 1 2 2 2 3 3 1 1 1 ...
$ id    : Factor w/ 316 levels "1","2","3","4",...: 1 2 3 4 5 6 7 8 9 10 ...
$ btype : Factor w/ 3 levels "curse","scold",...: 1 1 1 1 1 1 1 1 1 ...
$ situ  : Factor w/ 2 levels "other","self": 1 1 1 1 1 1 1 1 1 ...
$ mode  : Factor w/ 2 levels "want","do": 1 1 1 1 1 1 1 1 1 ...
$ r2    : Factor w/ 2 levels "N","Y": 1 1 2 2 2 2 1 1 2 ...

> stopifnot(nrow(unique(subset(VerbAgg, select = c(item, + btype, situ, mode)))) == 24, nrow(unique(subset(VerbAgg, + select = c(id, Anger, Gender)))) == 316)
```
Influence of item-level covariates

- We can check the proportions of responses for combinations of item-level covariates

```r
> round(100 * ftable(prop.table(xtabs(~mode + situ + resp, VerbAgg), 1:2)), 1)

<table>
<thead>
<tr>
<th>resp</th>
<th>no</th>
<th>perhaps</th>
<th>yes</th>
</tr>
</thead>
<tbody>
<tr>
<td>mode</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>want</td>
<td>37.7</td>
<td>30.0</td>
<td>32.3</td>
</tr>
<tr>
<td>self</td>
<td>55.9</td>
<td>29.1</td>
<td>15.0</td>
</tr>
<tr>
<td>do</td>
<td>49.8</td>
<td>27.2</td>
<td>23.0</td>
</tr>
<tr>
<td>self</td>
<td>66.2</td>
<td>23.5</td>
<td>10.3</td>
</tr>
</tbody>
</table>
```
Influence of person-level covariates

![Graph showing the influence of person-level covariates on Anger Index (STAXI).](#)
Initial model fit

Generalized linear mixed model fit by the Laplace approximation
Formula: r2 ~ Anger * Gender + situ + btype + mode + (1 | id) + (1 | item)

Data: VerbAgg
AIC  BIC  logLik  deviance
8156 8225  -4068  8136

Random effects:
  Groups   Name   Variance  Std.Dev.
  id      (Intercept)  1.79340  1.33918
  item    (Intercept)   0.11714  0.34226

Number of obs: 7584, groups: id, 316; item, 24

Fixed effects:
   Estimate  Std. Error   z value  Pr(>|z|)
(Intercept)    0.53205    0.43378    1.227  0.22000
   Anger        0.05849    0.01947    3.003  0.00267
GenderF        0.40229    0.78232    0.514  0.60710
  situ oneself  -1.05430    0.15119   -6.973 3.09e-12
  btype scold   -1.05980    0.18415   -5.755 8.67e-09
  btype shout   -2.10383    0.18651  -11.280 < 2e-16
       mode do   -0.70703    0.15100  -4.682 2.84e-06
   Anger::GenderF -0.00411    0.03817   -0.108  0.91425
Removing non-significant gender effects

Generalized linear mixed model fit by the Laplace approximation

Formula: r2 ~ Anger + situ + btype + mode + (1 | id) + (1 | item)

Data: VerbAgg

AIC  BIC  logLik  deviance
8155  8210  -4069   8139

Random effects:

<table>
<thead>
<tr>
<th>Groups</th>
<th>Name</th>
<th>Variance</th>
<th>Std.Dev.</th>
</tr>
</thead>
<tbody>
<tr>
<td>id</td>
<td>(Intercept)</td>
<td>1.81154</td>
<td>1.34594</td>
</tr>
<tr>
<td>item</td>
<td>(Intercept)</td>
<td>0.11721</td>
<td>0.34236</td>
</tr>
</tbody>
</table>

Number of obs: 7584, groups: id, 316; item, 24

Fixed effects:

|               | Estimate | Std. Error | z value | Pr(>|z|) |
|---------------|----------|------------|---------|---------|
| (Intercept)   | 0.63927  | 0.38334    | 1.668   | 0.095392|
| Anger         | 0.05685  | 0.01682    | 3.380   | 0.000726|
| situself      | -1.05437 | 0.15123    | -6.972  | 3.12e-12 |
| btypescold    | -1.05972 | 0.18420    | -5.753  | 8.76e-09 |
| btypeshout    | -2.10391 | 0.18656    | -11.278 | < 2e-16  |
| modedo        | -0.70725 | 0.15104    | -4.683  | 2.83e-06 |
Allowing situational/behavior random effects by person

Generalized linear mixed model fit by the Laplace approximation

Formula: r2 ~ Anger + situ + btype + mode + (1 | id:btype) + (1 | id:situ) + (1 | id:mode) + (1 | id) + (1 | item)

Data: VerbAgg

AIC  BIC logLik deviance
 7751 7827  -3865  7729

Random effects:

Groups   Name          Variance  Std.Dev.
id:btype (Intercept)  1.41069  1.18772
id:mode   (Intercept)  0.80916  0.89954
id:situ   (Intercept)  0.61539  0.78447
id        (Intercept)  1.70950  1.30748
item      (Intercept)  0.17213  0.41489

Number of obs: 7584, groups: id:btype, 948; id:mode, 632; id:situ, 632;

Fixed effects:

                              Estimate  Std. Error   z value  Pr(>|z|)
(Intercept)                   0.79168    0.48183    1.643   0.100369
Anger                        0.07483    0.02107    3.551   0.000384
situself                    -1.35741    0.19199   -7.070  1.55e-12
btypescold                 -1.36066    0.24118   -5.642  1.68e-08
btypeshout                  -2.69333    0.24372  -11.051  < 2e-16
modedo                      -0.94096    0.19503   -4.825   1.40e-06
Item-specific random effects

S4DoScold
S4Docurse
S4WantShout
S1WantShout
S2WantShout
S1DoScold
S4DoShout
S1DoShout
S4wantcurse
S3WantShout
S4WantScold
S1Docurse
S2DoScold
S2WantScold
S3Docurse
S1WantScold
S2wantcurse
S3WantScold
S3DoScold
S3wantcurse
S2Docurse
S2DoShout
S3DoShout
S1wantcurse
Person-specific random effects - Intercept
Correlated random effects by person

Generalized linear mixed model fit by the Laplace approximation

Formula: r2 ~ Anger + situ + btype + mode + (1 + situ + btype + mode | id) + (1 | item)

Data: VerbAgg

AIC   BIC  logLik  deviance
7727  7880  -3842  7683

Random effects:

Groups   Name   Variance  Std.Dev.  Corr
id  (Intercept)  4.53798   2.13025
    situself    1.31457   1.14655  -0.521
    btypescold  1.62184   1.27352  -0.085  -0.247
    btypeshout  4.03040   2.00759  -0.374   0.010   0.423
    modedo     1.68339   1.29745  -0.295   0.112   0.103
item  (Intercept) 0.18221   0.42686
      0.104

Number of obs: 7584, groups: id, 316; item, 24

Fixed effects:

   Estimate Std. Error  z value  Pr(>|z|)
(Intercept)   0.98349   0.47248   2.082  0.03738
   Anger       0.06564   0.02041   3.217  0.00130
 situself     -1.37642   0.19723  -6.979 2.97e-12
Outline

Organizing and plotting data; simple, scalar random effects

Models for longitudinal data

Interactions of grouping factors and other covariates

Evaluating the log-likelihood

Generalized Linear Mixed Models

Item Response Models as GLMMs

Nonlinear Mixed Models
Nonlinear mixed-effects models (NLMM)

- The LMM and GLMM are powerful data analysis tools.
- The “common denominator” of these models is the expression for the linear predictor. The models require that the fixed effects parameters and the random effects occur linearly in

\[ \eta = X\beta + Zb = X\beta + A'u \]

- This is a versatile and flexible way of specifying empirical models, whose form is determined from the data.
- In many situations, however, the form of the model is derived from external considerations of the mechanism generating the response. The parameters in such mechanistic models often occur nonlinearly.
- Mechanistic models can emulate behavior like the response approaching an asymptote, which is not possible with models that are linear in the parameters.
The Michaelis-Menten model, $SSmicmen$

$$y = \frac{\phi_1 x}{x + \phi_2}$$

$\phi_1$ (called $V_m$ in enzyme kinetics) is the maximum reaction velocity, $\phi_2 (K)$ is the concentration at which $y = \phi_1/2$. 
The “asymptotic regression” model, SSasymp

\[ y = \phi_1 + (\phi_1 - \phi_2)e^{-\phi_3 x} \]
The logistic growth model, SSlogis

\[ y = \frac{\phi_1}{1 + e^{-(x - \phi_2)/\phi_3}} \]
Modeling repeated measures data with a nonlinear model

- Nonlinear mixed-effects models are used extensively with longitudinal pharmacokinetic data.
- For such data the time pattern of an individual’s response is determined by pharmacokinetic parameters (e.g. rate constants) that occur nonlinearly in the expression for the expected response.
- The form of the nonlinear model is determined by the pharmacokinetic theory, not derived from the data.

\[
d \cdot k_e \cdot k_a \cdot C \frac{e^{-k_et} - e^{-k_at}}{k_a - k_e}
\]

- These pharmacokinetic parameters vary over the population. We wish to characterize typical values in the population and the extent of the variation.
- Thus, we associate random effects with the parameters, \( k_a, k_e \) and \( C \) in the nonlinear model.
A simple example - logistic model of growth curves

- The Orange data set are measurements of the growth of a sample of five orange trees in a location in California.
- The response is the circumference of the tree at a particular height from the ground (often converted to “diameter at breast height”).
- The covariates are age (days) and Tree (balanced).
- A data plot indicates that the growth patterns are similar but the eventual heights vary.
- One possible growth model is the logistic growth model

\[ f(t, A, t_0, s) = \frac{A}{1 + e^{-(t-t_0)/s}} \]

which can be seen to be related to the inverse logit link function.
Orange tree growth data

![Orange tree growth data graph](image-url)
Using `nlmer`

- The nonlinear mixed-effects model is fit with the `nlmer` function in the `lme4` package.
- The formula argument for `nlmer` is in three parts: the response, the nonlinear model function depending on covariates and a set of nonlinear model (nm) parameters, and the mixed-effects formula.
- There is no longer a concept of an intercept or a 1 term in the mixed-effects model. All terms in the mixed-effects formula incorporate names of nm parameters.
- The default term for the fixed-effects is a separate “intercept” parameter for each nm parameter.
- At present, the nonlinear model must provide derivatives, in addition to the expected response. The `deriv` function can be used to create such a function from an expression.
- The starting values for the fixed effects must also be given. It is safest to phrase these as a named vector.
Model fit for orange tree data

\[
> \text{print(nm1 <- nlmer(circumference} ~ \text{SSlogis(age,} \\
+ \text{Asym, xmid, scal)} ~ \text{Asym} | \text{Tree, Orange, start} = \text{c(Asym} = \\
+ \text{xmid = 770, scal = 120)}), \text{corr} = \text{FALSE})
\]

Nonlinear mixed model fit by the Laplace approximation
Formula: circumference ~ SSlogis(age, Asym, xmid, scal) ~ Asym | Tree
Data: Orange
AIC BIC logLik deviance
273.1 280.9 -131.6 263.1
Random effects:
 Groups Name Variance Std.Dev.
 Tree Asym 1001.493 31.646
 Residual 61.513 7.843
Number of obs: 35, groups: Tree, 5
Fixed effects:
 Estimate Std. Error t value
 Asym 192.05 15.58 12.32
 xmid 727.91 34.44 21.14
 scal 348.07 26.31 13.23
Random effects for trees
Extending the model

• Model nm1 incorporates random effects for the asymptote only. The asymptote parameter occurs linearly in the model expression. When random effects are associated with only such *conditionally linear* parameters, the Laplace approximation to the deviance is exact.

• We can allow more general specifications of random effects. In practice it is difficult to estimate many variance and covariance parameters when the number of levels of the grouping factor (*Tree*) is small.

• Frequently we begin with independent random effects to see which parameters show substantial variability. Later we allow covariances.

• This is not a fool-proof modeling strategy by any means but it is somewhat reasonable.
Independent random effects for each parameter

Nonlinear mixed model fit by the Laplace approximation
Formula: circumference ~ SSlogis(age, Asym, xmid, scal) ~ (Asym | Tree)

Data: Orange

AIC  BIC  logLik  deviance
277.1 288.0  -131.5  263.1

Random effects:
Groups   Name   Variance   Std.Dev.
Tree     Asym   968.559   31.1217
Tree     xmid   0.000     0.0000
Tree     scal   342.032   18.4941
Residual    scal   59.519    7.7149

Number of obs: 35, groups: Tree, 5

Fixed effects:

<table>
<thead>
<tr>
<th>Estimate</th>
<th>Std. Error</th>
<th>t value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Asym</td>
<td>192.34</td>
<td>12.54</td>
</tr>
<tr>
<td>xmid</td>
<td>728.93</td>
<td>21.49</td>
</tr>
<tr>
<td>scal</td>
<td>350.83</td>
<td>12.85</td>
</tr>
</tbody>
</table>
Correlated random effects for Asym and scal only

Nonlinear mixed model fit by the Laplace approximation
Formula: circumference ~ SSlogis(age, Asym, xmid, scal) ~ (Asym + scal

Data: Orange
AIC   BIC logLik deviance
274.1 285.0 -130.1 260.1

Random effects:
Groups   Name Variance Std.Dev. Corr
Tree     Asym  828.134  28.777
         scal  924.774  30.410  -1.000
Residual  55.591  7.456

Number of obs: 35, groups: Tree, 5

Fixed effects:

Estimate Std. Error t value
Asym     192.39   14.23   13.52
xmid   726.95   32.00   22.72
scal   357.90   28.42   12.59
Singular variance-covariance matrix

<table>
<thead>
<tr>
<th>Tree</th>
<th>Asym</th>
<th>scal</th>
</tr>
</thead>
<tbody>
<tr>
<td>3</td>
<td>-40</td>
<td>-40</td>
</tr>
<tr>
<td>5</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>1</td>
<td>20</td>
<td>20</td>
</tr>
<tr>
<td>4</td>
<td>40</td>
<td>40</td>
</tr>
</tbody>
</table>

Interactions Theory GLMM Item Response NLMM
Simple

Tree
3
1
5
2
4
−40 −20 0 20 40
●
●
●
●
●
Asym
−40 −20 0 20 40
●
●
●
●
●
scal
Theophylline pharmacokinetics
Nonlinear mixed model fit by the Laplace approximation
Formula: conc ~ SSfol(Dose, Time, lKe, lKa, lCl) ~ (lKe + lKa + lCl | Subject)
Data: Theoph
AIC  BIC logLik deviance
374  402.8   -177    354
Random effects:
  Groups   Name   Variance  Std.Dev.  Corr
  Subject  lKe  4.9461e-14  2.2240e-07
   lKa  4.3088e-01  6.5642e-01   0.000
   lCl  2.8050e-02  1.6748e-01  0.000  -0.007
Residual 5.0094e-01  7.0777e-01
Number of obs: 132, groups: Subject, 12
Fixed effects:
  Estimate  Std. Error  t value
  lKe   -2.46559    0.05187   -47.53
  lKa    0.48216    0.19986     2.41
  lCl  -3.23041    0.05952   -54.27
Nonlinear mixed model fit by the Laplace approximation

Formula: conc ~ SSfol(Dose, Time, lKe, lKa, lCl) ~ (lKa + lCl | Subject)

Data: Theoph

AIC    BIC    logLik    deviance
368    388.2  -177      354

Random effects:

Groups   Name    Variance    Std.Dev.    Corr
Subject   lKa    0.430870    0.65641
          lCl    0.028050    0.16748  -0.007
Residual  0.500939    0.70777

Number of obs: 132, groups: Subject, 12

Fixed effects:

             Estimate    Std. Error   t value
lKe          -2.46559     0.05187   -47.53
lKa           0.48217     0.19985     2.41
lCl         -3.23041     0.05952   -54.27
Remove correlation

```r
> print(nm6 <- nlmer(conc ~ SSfol(Dose, Time, lKe, + lKa, lCl) ~ (lKa | Subject) + (lCl | Subject), + Theoph, start = Th.start), corr = FALSE)
```

Nonlinear mixed model fit by the Laplace approximation
Formula: conc ~ SSfol(Dose, Time, lKe, lKa, lCl) ~ (lKa | Subject) + (lCl | Subject)
  Data: Theoph
  AIC  BIC logLik deviance
  366 383.3  -177  354
Random effects:
  Groups   Name     Variance Std.Dev.
  Subject  lKa  0.430900   0.65643
  Subject  lCl  0.028063   0.16751
  Residual 0.500943   0.70777
Number of obs: 132, groups: Subject, 12
Fixed effects:
  Estimate Std. Error  t value
  lKe -2.46552   0.05187  -47.53
  lKa  0.48214   0.19986    2.41
  lCl -3.23036   0.05953  -54.26
Random effects for clearance and absorption
Methodology

• Evaluation of the deviance is very similar to the calculation for the generalized linear mixed model. For given parameter values $\theta$ and $\beta$ the conditional mode $\tilde{u}(\theta, \beta)$ is determined by solving a penalized nonlinear least squares problem.

• $r^2(\theta, \beta)$ and $|L|^2$ determine the Laplace approximation to the deviance.

• As for GLMMs this can (and will) be extended to an adaptive Gauss-Hermite quadrature evaluation when there is only one grouping factor for the random effects.

• The theory (and, I hope, the implementation) for the generalized nonlinear mixed model (GNLMM) is straightforward, once you get to this point. Map first through the nonlinear model function then through the inverse link function.
From linear predictor to $\mu$

- The main change in evaluating $\mu_{Y|U}$ for NLMMs is in the role of the linear predictor. If there are $s$ nonlinear model (nm) parameters and $n$ observations in total then the model matrix $X$ is $n \cdot s \times p$ and the model matrix $Z$ is $n \cdot s \times q$.

- The linear predictor, $v = X\beta + A'u$, of length $n \cdot s$, is rearranged as an $n \times s$ matrix of parameter values $\Phi$. The $i$th component of the unbounded predictor, $\eta$, is the nonlinear model evaluated for the $i$ set of covariate values with the nonlinear parameters, $\phi$, at the $i$th row of $\Phi$.

$$
\begin{align*}
    u & \rightarrow b \rightarrow v \rightarrow \Phi \rightarrow \eta \rightarrow \mu \\
    b &= T(\theta)S(\theta)P'u \\
    v &= X\beta + Zb = X\beta + A(\theta)'P'u = \text{vec}(\Phi) \\
    \eta &= f(t, \Phi) \\
    \mu &= g^{-1}\eta
\end{align*}
$$
Generalizations of PIRLS

• The reason that the PLS problem for determining the conditional modes is relatively easy is because the standard least squares-based methods for fixed-effects models are easily adapted.

• For linear mixed-models the PLS problem is solved directly. In fact, for LMMs it is possible to determine the conditional modes of the random effects and the conditional estimates of the fixed effects simultaneously.

• Parameter estimates for generalized linear models (GLMs) are (very efficiently) determined by iteratively re-weighted least squares (IRLS) so the conditional modes in a GLMM are determined by penalized iteratively re-weighted least squares (PIRLS).

• Nonlinear least squares, used for fixed-effects nonlinear regression, is adapted as penalized nonlinear least squares (PNLS) or penalized iteratively reweighted nonlinear least squares (PIRNLS) for generalized nonlinear mixed models.