Fitting Linear Mixed-Effects Models Using lme4

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Abstract

Maximum likelihood or restricted maximum likelihood (REML) estimates of the parameters in linear mixed-effects models can be determined using the \texttt{lmer} function in the \texttt{lme4} package for R. As for most model-fitting functions in R, the model is described in an \texttt{lmer} call by a formula, in this case including both fixed- and random-effects terms. The formula and data together determine a numerical representation of the model from which the profiled deviance or the profiled REML criterion can be evaluated as a function of some of the model parameters. The appropriate criterion is optimized, using one of the constrained optimization functions in R, to provide the parameter estimates. We describe the structure of the model, the steps in evaluating the profiled deviance or REML criterion, and the structure of classes or types that represents such a model. Sufficient detail is included to allow specialization of these structures by users who wish to write functions to fit specialized linear mixed models, such as models incorporating pedigrees or smoothing splines, that are not easily expressible in the formula language used by \texttt{lmer}.

Keywords: sparse matrix methods, linear mixed models, penalized least squares, Cholesky decomposition.

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1. Introduction

The \texttt{lme4} package \citep{Bates:2014} for R \citep{R.Core.Team:2015} provides functions to fit and analyze linear mixed models, generalized linear mixed models and nonlinear mixed models. In each of these names, the term “mixed” or, more fully, “mixed effects”, denotes a model that incorporates both fixed- and random-effects terms in a linear predictor expression from which the conditional mean of the response can be evaluated. In this paper we describe the formulation and representation of linear mixed models. The techniques used for generalized linear and nonlinear mixed models will be described separately, in a future paper.

At present, the main alternative to \texttt{lme4} for mixed modeling in R is the \texttt{nlme} package \citep{Pinheiro:2014}. The main features distinguishing \texttt{lme4} from \texttt{nlme} are (1) more efficient linear algebra tools, giving improved performance on large problems; (2) simpler syntax and more efficient implementation for fitting models with crossed random effects; (3) the implementation of profile likelihood confidence intervals on random-effects parameters; and (4) the ability to fit generalized linear mixed models (al-
though in this paper we restrict ourselves to linear mixed models). The main advantage of \texttt{nlme} relative to \texttt{lme4} is a user interface for fitting models with structure in the residuals (various forms of heteroscedasticity and autocorrelation) and in the random-effects covariance matrices (e.g., compound symmetric models). With some extra effort, the computational machinery of \texttt{lme4} can be used to fit structured models that the basic \texttt{lmer} function cannot handle (see Appendix A).

The development of general software for fitting mixed models remains an active area of research with many open problems. Consequently, the \texttt{lme4} package has evolved since it was first released, and continues to improve as we learn more about mixed models. However, we recognize the need to maintain stability and backward compatibility of \texttt{lme4} so that it continues to be broadly useful. In order to maintain stability while continuing to advance mixed-model computation, we have developed several additional frameworks that draw on the basic ideas of \texttt{lme4} but modify its structure or implementation in various ways. These descendants include the \texttt{MixedModels} package (Bates 2015) in Julia (Bezanson, Karpinski, Shah, and Edelman 2012), the \texttt{lme4pureR} package (Bates and Walker 2013) in R, and the \texttt{flexLambda} development branch of \texttt{lme4}. The current article is largely restricted to describing the current stable version of the \texttt{lme4} package (1.1-7), with Appendix A describing hooks into the computational machinery that are designed for extension development. The \texttt{gamm4} (Wood and Scheipl 2014) and \texttt{blme} (Dorie 2015; Chung, Rabe-Hesketh, Dorie, Gelman, and Lin 2013) packages currently make use of these hooks.

Another goal of this article is to contrast the approach used by \texttt{lme4} with previous formulations of mixed models. The expressions for the profiled log-likelihood and profiled REML (restricted maximum likelihood) criteria derived in Section 3.4 are similar to those presented in Bates and DebRoy (2004) and, indeed, are closely related to “Henderson’s mixed-model equations” (Henderson Jr. 1982). Nonetheless there are subtle but important changes in the formulation of the model and in the structure of the resulting penalized least squares (PLS) problem to be solved (Section 3.6). We derive the current version of the PLS problem (Section 3.2) and contrast this result with earlier formulations (Section 3.5).

This article is organized into four main sections (Sections 2, 3, 4, and 5), each of which corresponds to one of the four largely separate modules that comprise \texttt{lme4}. Before describing the details of each module, we describe the general form of the linear mixed model underlying \texttt{lme4} (Section 1.1); introduce the \texttt{sleepstudy} data that will be used as an example throughout (Section 1.2); and broadly outline \texttt{lme4}’s modular structure (Section 1.3).

### 1.1. Linear mixed models

Just as a linear model is described by the distribution of a vector-valued random response variable, $\mathcal{Y}$, whose observed value is $y_{\text{obs}}$, a linear mixed model is described by the distribution of two vector-valued random variables: $\mathcal{Y}$, the response, and $\mathcal{B}$, the vector of random effects. In a linear model the distribution of $\mathcal{Y}$ is multivariate normal,

$$
\mathcal{Y} \sim \mathcal{N}(X\beta + o, \sigma^2 W^{-1}),
$$

where $n$ is the dimension of the response vector, $W$ is a diagonal matrix of known prior weights, $\beta$ is a $p$-dimensional coefficient vector, $X$ is an $n \times p$ model matrix, and $o$ is a vector of known prior offset terms. The parameters of the model are the coefficients $\beta$ and the scale parameter $\sigma$. 
In a linear mixed model it is the conditional distribution of $Y$ given $B = b$ that has such a form,

$$ (Y | B = b) \sim N(X\beta + Zb + o, \sigma^2W^{-1}), $$

where $Z$ is the $n \times q$ model matrix for the $q$-dimensional vector-valued random-effects variable, $B$, whose value we are fixing at $b$. The unconditional distribution of $B$ is also multivariate normal with mean zero and a parameterized $q \times q$ variance-covariance matrix, $\Sigma$,

$$ B \sim N(0, \Sigma). $$

As a variance-covariance matrix, $\Sigma$ must be positive semidefinite. It is convenient to express the model in terms of a relative covariance factor, $\Lambda_\theta$, which is a $q \times q$ matrix, depending on the variance-component parameter, $\theta$, and generating the symmetric $q \times q$ variance-covariance matrix, $\Sigma$, according to

$$ \Sigma_\theta = \sigma^2\Lambda_\theta\Lambda_\theta^\top, $$

where $\sigma$ is the same scale factor as in the conditional distribution (2).

Although Equations 2, 3, and 4 fully describe the class of linear mixed models that \texttt{lme4} can fit, this terse description hides many important details. Before moving on to these details, we make a few observations:

- This formulation of linear mixed models allows for a relatively compact expression for the profiled log-likelihood of $\theta$ (Section 3.4, Equation 35).
- The matrices associated with random effects, $Z$ and $\Lambda_\theta$, typically have a sparse structure with a sparsity pattern that encodes various model assumptions. Sections 2.3 and 3.7 provide details on these structures, and how to represent them efficiently.
- The interface provided by \texttt{lme4}'s \texttt{lmer} function is slightly less general than the model described by Equations 2, 3, and 4. To take advantage of the entire range of possibilities, one may use the modular functions (Sections 1.3 and Appendix A) or explore the experimental \texttt{flexLambda} branch of \texttt{lme4} on Github.

### 1.2. Example

Throughout our discussion of \texttt{lme4}, we will work with a data set on the average reaction time per day for subjects in a sleep deprivation study (Belenky et al. 2003). On day 0 the subjects had their normal amount of sleep. Starting that night they were restricted to 3 hours of sleep per night. The response variable, Reaction, represents average reaction times in milliseconds (ms) on a series of tests given each Day to each Subject (Figure 1),

```r
> str(sleepstudy)
'data.frame': 180 obs. of 3 variables:
$ Reaction: num 250 259 251 321 357 ... $ Days : num 0 1 2 3 4 5 6 7 8 9 ... $ Subject : Factor w/ 18 levels "308","309","310",:: 1 1 1 1 1 1 1 1 ..
```
Figure 1: Average reaction time versus days of sleep deprivation by subject. Subjects ordered (from left to right starting on the top row) by increasing slope of subject-specific linear regressions.

Each subject’s reaction time increases approximately linearly with the number of sleep-deprived days. However, subjects also appear to vary in the slopes and intercepts of these relationships, which suggests a model with random slopes and intercepts. As we shall see, such a model may be fitted by minimizing the REML criterion (Equation 40) using

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

The estimates of the standard deviations of the random effects for the intercept and the slope are 24.74 ms and 5.92 ms/day. The fixed-effects coefficients, $\beta$, are 251.4 ms and 10.47 ms/day for the intercept and slope. In this model, one interpretation of these fixed effects is that they are the estimated population mean values of the random intercept and slope (Section 2.2).

We have chosen the `sleepstudy` example because it is a relatively small and simple example to illustrate the theory and practice underlying `lmer`. However, `lmer` is capable of fitting more complex mixed models to larger data sets. For example, we direct the interested reader to `RShowDoc("lmerperf", package = "lme4")` for examples that more thoroughly exercise the performance capabilities of `lmer`.

### 1.3. High-level modular structure

The `lmer` function is composed of four largely independent modules. In the first module, a mixed-model formula is parsed and converted into the inputs required to specify a linear mixed model (Section 2). The second module uses these inputs to construct an R function which takes the covariance parameters, $\theta$, as arguments and returns negative twice the log profiled
### Module R function Description

<table>
<thead>
<tr>
<th>Module</th>
<th>R function</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Formula module</td>
<td><code>lFormula</code></td>
<td>Accepts a mixed-model formula, data, and other user inputs, and returns a list of objects required to fit a linear mixed model.</td>
</tr>
<tr>
<td>Objective function module</td>
<td><code>mkLmerDevfun</code></td>
<td>Accepts the results of <code>lFormula</code> and returns a function to calculate the deviance (or restricted deviance) as a function of the covariance parameters, $\theta$.</td>
</tr>
<tr>
<td>Optimization module</td>
<td><code>optimizeLmer</code></td>
<td>Accepts a deviance function returned by <code>mkLmerDevfun</code> and returns the results of the optimization of that deviance function.</td>
</tr>
<tr>
<td>Output module</td>
<td><code>mkMerMod</code></td>
<td>Accepts an optimized deviance function and packages the results into a useful object.</td>
</tr>
</tbody>
</table>

Table 1: The high-level modular structure of `lmer`.

likelihood or the REML criterion (Section 3). The third module optimizes this objective function to produce maximum likelihood (ML) or REML estimates of $\theta$ (Section 4). Finally, the fourth module provides utilities for interpreting the optimized model (Section 5).

To illustrate this modularity, we recreate the `fm1` object by a series of four modular steps: the formula module,

```r
> parsedFormula <- lFormula(formula = Reaction ~ Days + (Days | Subject),
                        data = sleepstudy)
```

the objective function module,

```r
> devianceFunction <- do.call(mkLmerDevfun, parsedFormula)
```

the optimization module,

```r
> optimizerOutput <- optimizeLmer(devianceFunction)
```

and the output module,

```r
> mkMerMod(   rho = environment(devianceFunction),
          opt = optimizerOutput,
          reTrms = parsedFormula$reTrms,
          fr = parsedFormula$fr)
```

### 2. Formula module
2.1. Mixed-model formulas

Like most model-fitting functions in R, `lmer` takes as its first two arguments a formula specifying the model and the data with which to evaluate the formula. This second argument, data, is optional but recommended and is usually the name of an R data frame. In the R `lm` function for fitting linear models, formulas take the form `resp ~ expr`, where `resp` determines the response variable and `expr` is an expression that specifies the columns of the model matrix. Formulas for the `lmer` function contain special random-effects terms,

```r
> resp ~ FEexpr + (REexpr1 | factor1) + (REexpr2 | factor2) + ...
```

where `FEexpr` is an expression determining the columns of the fixed-effects model matrix, `X`, and the random-effects terms, `(REexpr1 | factor1)` and `(REexpr2 | factor2)`, determine both the random-effects model matrix, `Z` (Section 2.3.1), and the structure of the relative covariance factor, `A_\theta` (Section 2.3.2). In principle, a mixed-model formula may contain arbitrarily many random-effects terms, but in practice the number of such terms is typically low.

2.2. Understanding mixed-model formulas

Before describing the details of how `lme4` parses mixed-model formulas (Section 2.3), we provide an informal explanation and then some examples. Our discussion assumes familiarity with the standard R modeling paradigm (Chambers 1993).

Each random-effects term is of the form `(expr | factor)`. The expression `expr` is evaluated as a linear model formula, producing a model matrix following the same rules used in standard R modeling functions (e.g., `lm` or `glm`). The expression `factor` is evaluated as an R factor. One way to think about the vertical bar operator is as a special kind of interaction between the model matrix and the grouping factor. This interaction ensures that the columns of the model matrix have different effects for each level of the grouping factor. What makes this a special kind of interaction is that these effects are modeled as unobserved random variables, rather than unknown fixed parameters. Much has been written about important practical and philosophical differences between these two types of interactions (e.g., Henderson Jr. 1982; Gelman 2005). For example, the random-effects implementation of such interactions can be used to obtain shrinkage estimates of regression coefficients (e.g., Efron and Morris 1977), or account for lack of independence in the residuals due to block structure or repeated measurements (e.g., Laird and Ware 1982).

Table 2 provides several examples of the right-hand-sides of mixed-model formulas. The first example, `(1 | g)`, is the simplest possible mixed-model formula, where each level of the grouping factor, `g`, has its own random intercept. The mean and standard deviation of these intercepts are parameters to be estimated. Our description of this model incorporates any nonzero mean of the random effects as fixed-effects parameters. If one wishes to specify that a random intercept has a priori known means, one may use the `offset` function as in the second model in Table 2. This model contains no fixed effects, or more accurately the fixed-effects model matrix, `X`, has zero columns and \( \beta \) has length zero.

We may also construct models with multiple grouping factors. For example, if the observations are grouped by `g2`, which is nested within `g1`, then the third formula in Table 2 can be used to model variation in the intercept. A common objective in mixed modeling is to account
### Table 2: Examples of the right-hand-sides of mixed-effects model formulas. The names of grouping factors are denoted \( g \), \( g_1 \), and \( g_2 \), and covariates and *a priori* known offsets as \( x \) and \( o \).

<table>
<thead>
<tr>
<th>Formula</th>
<th>Alternative</th>
<th>Meaning</th>
</tr>
</thead>
<tbody>
<tr>
<td>((1 \mid g))</td>
<td>(1 + (1 \mid g))</td>
<td>Random intercept with fixed mean.</td>
</tr>
<tr>
<td>(0 + \text{offset}(o) + (1 \mid g))</td>
<td>(-1 + \text{offset}(o) + (1 \mid g))</td>
<td>Random intercept with <em>a priori</em> means.</td>
</tr>
<tr>
<td>((1 \mid g_1/g_2))</td>
<td>((1 \mid g_1) + (1 \mid g_1:g_2))</td>
<td>Intercept varying among ( g_1 ) and ( g_2 ) within ( g_1 ).</td>
</tr>
<tr>
<td>((1 \mid g_1) + (1 \mid g_2))</td>
<td>(1 + (1 \mid g_1) + (1 \mid g_2))</td>
<td>Intercept varying among ( g_1 ) and ( g_2 ).</td>
</tr>
<tr>
<td>(x + (x \mid g))</td>
<td>(1 + x + (1 + x \mid g))</td>
<td>Correlated random intercept and slope.</td>
</tr>
<tr>
<td>(x + (x \mid</td>
<td>\mid g))</td>
<td>(1 + x + (1 \mid g) + (0 + x \mid g))</td>
</tr>
</tbody>
</table>

for such nested (or hierarchical) structure. However, one of the most useful aspects of *lme4* is that it can be used to fit random effects associated with non-nested grouping factors. For example, suppose the data are grouped by fully crossing two factors, \( g_1 \) and \( g_2 \), then the fourth formula in Table 2 may be used. Such models are common in item response theory, where *subject* and *item* factors are fully crossed (Doran, Bates, Bliese, and Dowling 2007). In addition to varying intercepts, we may also have varying slopes (e.g., the *sleepstudy* data, Section 1.2). The fifth example in Table 2 gives a model where both the intercept and slope vary among the levels of the grouping factor.

#### Specifying uncorrelated random effects

By default, *lme4* assumes that all coefficients associated with the same random-effects term are correlated. To specify an uncorrelated slope and intercept (for example), one may either use double-bar notation, \((x \mid| \mid g)\), or equivalently use multiple random-effects terms, \(x + (1 \mid g) + (0 + x \mid g)\), as in the final example of Table 2. For example, if one examined the results of model \( fm1 \) of the *sleepstudy* data (Section 1.2) using summary(fm1), one would see that the estimated correlation between the slope for \( Days \) and the intercept is fairly low (0.066) (See Section 5.2.2 below for more on how to extract the random-effects covariance matrix.) We may use double-bar notation to fit a model that excludes a correlation parameter:

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

Although mixed models where the random slopes and intercepts are assumed independent are commonly used to reduce the complexity of random-slopes models, they do have one subtle drawback. Models in which the slopes and intercepts are allowed to have a nonzero correlation (e.g., \( fm1 \)) are invariant to additive shifts of the continuous predictor (\( Days \) in this case). This invariance breaks down when the correlation is constrained to zero; any shift in the predictor will necessarily lead to a change in the estimated correlation, and in the likelihood and predictions of the model. For example, we can eliminate the correlation in
Symbol Size Description
\(n\) Length of the response vector, \(Y\)
\(p\) Number of columns of fixed-effects model matrix, \(X\)
\(q = \sum_{i}^{k} q_{i}\) Number of columns of random-effects model matrix, \(Z\)
\(p_{i}\) Number of columns of the raw model matrix, \(X_{i}\)
\(\ell_{i}\) Number of levels of the grouping factor indices, \(i_{i}\)
\(q_{i} = p_{i}\ell_{i}\) Number of columns of the term-wise model matrix, \(Z_{i}\)
\(k\) Number of random-effects terms
\(m_{i} = \frac{(p_{i} + 1)}{2}\) Number of covariance parameters for term \(i\)
\(m = \sum_{i}^{k} m_{i}\) Total number of covariance parameters

Table 3: Dimensions of linear mixed models. The subscript \(i = 1, \ldots, k\) denotes a specific random-effects term.

<table>
<thead>
<tr>
<th>Symbol</th>
<th>Size</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>(X_{i})</td>
<td>(n \times p_{i})</td>
<td>Raw random-effects model matrix</td>
</tr>
<tr>
<td>(J_{i})</td>
<td>(n \times \ell_{i})</td>
<td>Indicator matrix of grouping factor indices</td>
</tr>
<tr>
<td>(X_{ij})</td>
<td>(p_{i} \times 1)</td>
<td>Column vector containing (j)th row of (X_{i})</td>
</tr>
<tr>
<td>(J_{ij})</td>
<td>(\ell_{i} \times 1)</td>
<td>Column vector containing (j)th row of (J_{i})</td>
</tr>
<tr>
<td>(i_{i})</td>
<td>(n)</td>
<td>Vector of grouping factor indices</td>
</tr>
<tr>
<td>(Z_{i})</td>
<td>(n \times q_{i})</td>
<td>Term-wise random-effects model matrix</td>
</tr>
<tr>
<td>(\theta)</td>
<td>(m)</td>
<td>Covariance parameters</td>
</tr>
<tr>
<td>(T_{i})</td>
<td>(p_{i} \times p_{i})</td>
<td>Lower triangular template matrix</td>
</tr>
<tr>
<td>(\Lambda_{i})</td>
<td>(q_{i} \times q_{i})</td>
<td>Term-wise relative covariance factor</td>
</tr>
</tbody>
</table>

Table 4: Symbols used to describe the structure of the random-effects model matrix and the relative covariance factor. The subscript \(i = 1, \ldots, k\) denotes a specific random-effects term.

\(\text{fm1}\) simply by adding an amount equal to the ratio of the estimated among-subject standard deviations multiplied by the estimated correlation (i.e., \(\sigma_{\text{slope}} / \sigma_{\text{intercept}} \cdot \rho_{\text{slope:intercept}}\)) to the \texttt{Days} variable. The use of models such as \(\text{fm2}\) should ideally be restricted to cases where the predictor is measured on a ratio scale (i.e., the zero point on the scale is meaningful, not just a location defined by convenience or convention), as is the case here.

2.3. Algebraic and computational account of mixed-model formulas

The fixed-effects terms of a mixed-model formula are parsed to produce the fixed-effects model matrix, \(X\), in the same way that the \texttt{R} \texttt{lm} function generates model matrices. However, a mixed-model formula incorporates \(k \geq 1\) random-effects terms of the form \((r \mid f)\) as well. These \(k\) terms are used to produce the random-effects model matrix, \(Z\) (Equation 2; Section 2.3.1), and the structure of the relative covariance factor, \(\Lambda_{\theta}\) (Equation 4; Section 2.3.2), which are matrices that typically have a sparse structure. We now describe how one might construct these matrices from the random-effects terms, considering first a single term, \((r \mid f)\), and then generalizing to multiple terms. Tables 3 and 4 summarize the matrices and vectors that determine the structure of \(Z\) and \(\Lambda_{\theta}\).

The expression, \(r\), is a linear model formula that evaluates to an \texttt{R} model matrix, \(X_{i}\), of size \(n \times p_{i}\), called the \textit{raw random-effects model matrix} for term \(i\). A term is said to be a
**scalar** random-effects term when \( p_i = 1 \), otherwise it is **vector-valued**. For a simple, **scalar** random-effects term of the form \((1 \mid f)\), \(X_i\) is the \( n \times 1 \) matrix of ones, which implies a random intercept model.

The expression \( f \) evaluates to an R factor, called the **grouping factor**, for the term. For the \( i \)th term, we represent this factor mathematically with a vector \( i_i \) of **factor indices**, which is an \( n \)-vector of values from 1, \ldots, \( \ell_i \).¹ Let \( J_i \) be the \( n \times \ell_i \) matrix of indicator columns for \( i_i \). Using the **Matrix** package (Bates and Maechler 2015) in R, we may construct the transpose of \( J_i \) from a factor vector, \( f \), by coercing \( f \) to a ‘**sparseMatrix**’ object. For example,

```r
> (f <- gl(3, 2))
[1] 1 1 2 2 3 3
Levels: 1 2 3
> (Ji <- t(as(f, Class = "sparseMatrix")))

6 x 3 sparse Matrix of class "dgCMatrix"
     1 2 3
[1,] 1 . .
[2,] 1 . .
[3,] . 1 .
[4,] . 1 .
[5,] . . 1
[6,] . . 1
```

When \( k > 1 \) we order the random-effects terms so that \( \ell_1 \geq \ell_2 \geq \cdots \geq \ell_k \); in general, this ordering reduces “fill-in” (i.e., the proportion of elements that are zero in the lower triangle of \( \Lambda_\theta^T Z^T W Z \Lambda_\theta + I \) but not in the lower triangle of its left Cholesky factor, \( L_\theta \), described below in Equation 18). This reduction in fill-in provides more efficient matrix operations within the penalized least squares algorithm (Section 3.2).

**Constructing the random-effects model matrix**

The \( i \)th random-effects term contributes \( q_i = \ell_i p_i \) columns to the model matrix \( Z \). We group these columns into a matrix, \( Z_i \), which we refer to as the **term-wise model matrix** for the \( i \)th term. Thus \( q \), the number of columns in \( Z \) and the dimension of the random variable, \( B \), is

\[
q = \sum_{i=1}^{k} q_i = \sum_{i=1}^{k} \ell_i p_i.
\]

Creating the matrix \( Z_i \) from \( X_i \) and \( J_i \) is a straightforward concept that is, nonetheless, somewhat awkward to describe. Consider \( Z_i \) as being further decomposed into \( \ell_i \) blocks of \( p_i \) columns. The rows in the first block are the rows of \( X_i \) multiplied by the 0/1 values in

¹In practice, fixed-effects model matrices and random-effects terms are evaluated with respect to a **model frame**, ensuring that any expressions for grouping factors have been coerced to factors and any unused levels of these factors have been dropped. That is, \( \ell_i \), the number of levels in the grouping factor for the \( i \)th random-effects term, is well-defined.
the first column of $J_i$ and similarly for the subsequent blocks. With these definitions we may define the term-wise random-effects model matrix, $Z_i$, for the $i$th term as a transposed Khatri-Rao product,

$$Z_i = (J_i^T \ast X_i^T)^T = \begin{bmatrix}
J_{i1}^T \otimes X_{i1}^T \\
J_{i2}^T \otimes X_{i2}^T \\
\vdots \\
J_{in}^T \otimes X_{in}^T
\end{bmatrix},$$

(6)

where $\ast$ and $\otimes$ are the Khatri-Rao\(^2\) (Khatri and Rao 1968) and Kronecker products, and $J_{ij}^T$ and $X_{ij}^T$ are row vectors of the $j$th rows of $J_i$ and $X_i$. These rows correspond to the $j$th sample in the response vector, $Y_i$, and thus $j$ runs from $1, \ldots, n$. The Matrix package for R contains a KhatriRao function, which can be used to form $Z_i$. For example, if we begin with a raw model matrix,

```r
> (Xi <- cbind(1, rep.int(c(-1, 1), 3L)))
[,1] [,2]
[1,] 1 -1
[2,] 1 1
[3,] 1 -1
[4,] 1 1
[5,] 1 -1
[6,] 1 1
```

then the term-wise random-effects model matrix is,

```r
> (Zi <- t(KhatriRao(t(Ji), t(Xi))))
```

```
6 x 6 sparse Matrix of class "dgCMatrix"

[1,] 1 -1 . . . .
[2,] 1 1 . . . .
[3,] . . 1 -1 . .
[4,] . . 1 1 . .
[5,] . . . . 1 -1
[6,] . . . . 1 1
```

In particular, for a simple, scalar term, $Z_i$ is exactly $J_i$, the matrix of indicator columns. For other scalar terms, $Z_i$ is formed by element-wise multiplication of the single column of $X_i$ by each of the columns of $J_i$.

Because each $Z_i$ is generated from indicator columns, its cross-product, $Z_i^T Z_i$ is block-diagonal consisting of $\ell_i$ diagonal blocks each of size $p_i$.\(^3\) Note that this means that when

\(^2\)Note that the original definition of the Khatri-Rao product is more general than the definition used in the Matrix package, which is the definition we use here.

\(^3\)To see this, note that by the properties of Kronecker products we may write the cross-product matrix $Z_i^T Z_i$ as $\sum_{j=1}^{n} J_{ij} J_{ij}^T \otimes X_{ij} X_{ij}^T$. Because $J_{ij}$ is a unit vector along a coordinate axis, the cross-product $J_{ij} J_{ij}^T$ is a $p_i \times p_i$ matrix of all zeros except for a single 1 along the diagonal. Therefore, the cross-products, $X_{ij} X_{ij}^T$, will be added to one of the $\ell_i$ blocks of size $p_i \times p_i$ along the diagonal of $Z_i^T Z_i$.\)
\(k = 1\) (i.e., there is only one random-effects term, and \(Z_i = Z\)). \(Z^T Z\) will be block diagonal. These block-diagonal properties allow for more efficient sparse matrix computations (Section 3.7).

The full random-effects model matrix, \(Z\), is constructed from \(k \geq 1\) blocks,

\[
Z = \begin{bmatrix} Z_1 & Z_2 & \ldots & Z_k \end{bmatrix}.
\]

(7)

By transposing Equation 7 and substituting in Equation 6, we may represent the structure of the transposed random-effects model matrix as follows,

\[
Z^T = \begin{bmatrix}
J_{11} \otimes X_{11} & J_{12} \otimes X_{12} & \ldots & J_{1n} \otimes X_{1n} \\
J_{21} \otimes X_{21} & J_{22} \otimes X_{22} & \ldots & J_{2n} \otimes X_{2n} \\
\vdots & \vdots & \ddots & \vdots \\
J_{n1} \otimes X_{n1} & J_{n2} \otimes X_{n2} & \ldots & J_{nn} \otimes X_{nn}
\end{bmatrix}
\]

(8)

Note that the proportion of elements of \(Z^T\) that are structural zeros is

\[
\frac{\sum_{i=1}^{k} p_i (\ell_i - 1)}{\sum_{i=1}^{k} p_i}.
\]

(9)

Therefore, the sparsity of \(Z^T\) increases with the number of grouping factor levels. As the number of levels is often large in practice, it is essential for speed and efficiency to take account of this sparsity, for example by using sparse matrix methods, when fitting mixed models (Section 3.7).

**Constructing the relative covariance factor**

The \(q \times q\) covariance factor, \(\Lambda_\theta\), is a block diagonal matrix whose \(i\)th diagonal block, \(\Lambda_i\), is of size \(q_i\), \(i = 1, \ldots, k\). We refer to \(\Lambda_i\) as the term-wise relative covariance factor. Furthermore, \(\Lambda_i\) is a homogeneous block diagonal matrix with each of the \(\ell_i\) lower-triangular blocks on the diagonal being a copy of a \(p_i \times p_i\) lower-triangular template matrix, \(T_i\). The covariance parameter vector, \(\theta\), of length \(m_i = \left(\frac{p_i + 1}{2}\right)\), consists of the elements in the lower triangle of \(T_i\), \(i = 1, \ldots, k\). To provide a unique representation we require that the diagonal elements of the \(T_i\), \(i = 1, \ldots, k\) be non-negative.

The template, \(T_i\), can be constructed from the number \(p_i\) alone. In R code we denote \(p_i\) as \(\text{nc}\). For example, if we set \(\text{nc} \leftarrow 3\), we could create the template for term \(i\) as,

\[
\begin{align*}
\text{\textgreater{} (rowIndices} & \leftarrow \text{rep}(1:}\text{nc}, 1:}\text{nc)) \\
[1] & \begin{bmatrix} 1 & 2 & 2 & 3 & 3 & 3 \end{bmatrix} \\
\text{\textgreater{} (colIndices} & \leftarrow \text{sequence}(1:}\text{nc)) \\
[1] & \begin{bmatrix} 1 & 1 & 2 & 1 & 2 \end{bmatrix} \\
\text{\textgreater{} (template} & \leftarrow \text{sparseMatrix(rowIndices, colIndices,} \\
\text{ x = 1} * \text{(rowIndices} == \text{colIndices))))
\end{align*}
\]
3 x 3 sparse Matrix of class "dgCMatrix"

[1,] 1 . .
[2,] 0 1 .
[3,] 0 0 1

Note that the rowIndices and colIndices fill the entire lower triangle, which contains the initial values of the covariance parameter vector, \( \theta \),

\[
\Lambda^\top_i
\]

For example, if we set the number of levels, \( \ell_i \), to two, \( nl \leftarrow 2 \), we could create the transposed term-wise relative covariance factor, \( \Lambda^\top_i \), using the .bdiag function in the Matrix package,

\[
\Lambda^\top_i \leftarrow .bdiag(rep(list(t(template)), nl))
\]

6 x 6 sparse Matrix of class "dgTMatrix"

[1,] 1 0 0 . .
[2,] . 1 0 . .
[3,] . . 1 . .
[4,] . . . 1 0 0
[5,] . . . . 1 0
[6,] . . . . . 1

For a model with a single random-effects term, \( \Lambda^\top \) would be the initial transposed relative covariance factor \( \Lambda^\top_{\theta} \) itself.

The transposed relative covariance factor, \( \Lambda^\top_{\theta} \), that arises from parsing the formula and data is set at the initial value of the covariance parameters, \( \theta \). However, during model fitting, it needs to be updated to a new \( \theta \) value at each iteration (see Section 3.6.1). This is achieved by constructing a vector of indices, \( Lind \), that identifies which elements of \( \theta \) should be placed in which elements of \( \Lambda^\top \),

\[
Lind \leftarrow rep(LindTemplate, nl)
\]

For example, if we randomly generate a new value for \( \theta \),
we may update \( \Lambda_{\text{dat}} \) as follows,

\[
> \text{Lambdati}@x \leftarrow \text{thetanew}[Lind]
\]

Section 3.6 describes the process of updating the relative covariance factor in more detail.

### 3. Objective function module

#### 3.1. Model reformulation for improved computational stability

In our initial formulation of the linear mixed model (Equations 2, 3, and 4), the covariance parameter vector, \( \theta \), appears only in the marginal distribution of the random effects (Equation 3). However, from the perspective of computational stability and efficiency, it is advantageous to reformulate the model such that \( \theta \) appears only in the conditional distribution for the response vector given the random effects. Such a reformulation allows us to work with singular covariance matrices, which regularly arise in practice (e.g., during intermediate steps of the nonlinear optimizer, Section 4).

The reformulation is made by defining a **spherical** random-effects variable, \( U \), with distribution

\[
U \sim \mathcal{N}(0, \sigma^2 I_q).
\]

If we set,

\[
B = \Lambda_{\theta} U,
\]

then \( B \) will have the desired \( \mathcal{N}(0, \Sigma_{\theta}) \) distribution (Equation 3). Although it may seem more natural to define \( U \) in terms of \( B \) we must write the relationship as in Equation 11 to allow for singular \( \Lambda_{\theta} \). The conditional distribution (Equation 2) of the response vector given the random effects may now be reformulated as,

\[
(Y | U = u) \sim \mathcal{N}(\mu_{Y|U=u}, \sigma^2 W^{-1}),
\]

where

\[
\mu_{Y|U=u} = X \beta + Z \Lambda_{\theta} u + o
\]

is a vector of linear predictors, which can be interpreted as a conditional mean (or mode). Similarly, we also define \( \mu_{U|Y=y_{\text{obs}}} \) as the conditional mean (or mode) of the spherical random effects given the observed value of the response vector. Note also that we use the \( u \) symbol throughout to represent a specific value of the random variable, \( U \).

#### 3.2. Penalized least squares

Our computational methods for maximum likelihood fitting of the linear mixed model involve repeated applications of the PLS method. In particular, the PLS problem is to minimize the penalized weighted residual sum-of-squares,

\[
r^2(\theta, \beta, u) = \rho^2(\theta, \beta, u) + \|u\|^2,
\]

\({\mathcal{N}(\mu, \sigma^2 I)}\) distributions are called “spherical” because contours of the probability density are spheres.
over $\begin{bmatrix} u \\ \beta \end{bmatrix}$, where,

$$\rho^2(\theta, \beta, u) = \left\| W^{1/2} \left[ y_{\text{obs}} - \mu_{\mathcal{Y}|u=u}\right] \right\|^2,$$

is the weighted residual sum-of-squares. This notation makes explicit the fact that $r^2$ and $\rho^2$ both depend on $\theta$, $\beta$, and $u$. The reason for the word “penalized” is that the term, $\|u\|^2$, in Equation 14 penalizes models with larger magnitude values of $u$.

In the so-called “pseudo-data” approach we write the penalized weighted residual sum-of-squares as the squared length of a block matrix equation,

$$r^2(\theta, \beta, u) = \left\| W^{1/2} \begin{bmatrix} y_{\text{obs}} - o \\ 0 \\ X^T W \Lambda \theta + I_q \end{bmatrix} \right\|^2,$$

This pseudo-data approach shows that the PLS problem may also be thought of as a standard least squares problem for an extended response vector, which implies that the minimizing value, $\left[ \mu_{\mathcal{Y}|y=y_{\text{obs}}} \right]$, satisfies the normal equations,

$$\begin{bmatrix} \Lambda^\top \theta Z^\top W (y_{\text{obs}} - o) \\ X^\top W (y_{\text{obs}} - o) \end{bmatrix} = \begin{bmatrix} \Lambda^\top \theta Z^\top W Z \Lambda \theta + I \\ X^\top W Z \Lambda \theta \end{bmatrix} \begin{bmatrix} \mu_{\mathcal{Y}|y=y_{\text{obs}}} \\ \beta \end{bmatrix},$$

where $\mu_{\mathcal{Y}|y=y_{\text{obs}}}$ is the conditional mean of $\mathcal{Y}$ given that $Y = y_{\text{obs}}$. Note that this conditional mean depends on $\theta$, although we do not make this dependency explicit in order to reduce notational clutter.

The cross-product matrix in Equation 17 can be Cholesky decomposed,

$$\begin{bmatrix} \Lambda^\top \theta Z^\top W Z \Lambda \theta + I \\ X^\top W Z \Lambda \theta \end{bmatrix} = \begin{bmatrix} L^\top \theta & 0 \\ R^\top_{ZX} & R^\top_X \end{bmatrix} \begin{bmatrix} L \theta \\ R_{ZX} \end{bmatrix}.$$

We may use this decomposition to rewrite the penalized weighted residual sum-of-squares as,

$$r^2(\theta, \beta, u) = r^2(\theta) + \left\| L^\top \theta (u - \mu_{\mathcal{Y}|y=y_{\text{obs}}}) + R_{ZX} (\beta - \hat{\beta}_\theta) \right\|^2 + \left\| R_X (\beta - \hat{\beta}_\theta) \right\|^2,$$

where we have simplified notation by writing $r^2(\theta, \hat{\beta}_\theta, \mu_{\mathcal{Y}|y=y_{\text{obs}}})$ as $r^2(\theta)$. This is an important expression in the theory underlying lme4. It relates the penalized weighted residual sum-of-squares, $r^2(\theta, \beta, u)$, with its minimum value, $r^2(\theta)$. This relationship is useful in the next section where we integrate over the random effects, which is required for maximum likelihood estimation.

### 3.3. Probability densities

The residual sums-of-squares discussed in the previous section can be used to express various probability densities, which are required for maximum likelihood estimation of the linear
mixed model\(^5\),

\[
\begin{align*}
f_{Y\mid U}(y_{\text{obs}}\mid u) &= \frac{|W|^{1/2}}{(2\pi\sigma^2)^{n/2}} \exp \left[ -\frac{r^2(\theta, \beta, u)}{2\sigma^2} \right], \quad (20) \\
f_u(u) &= \frac{1}{(2\pi\sigma^2)^{q/2}} \exp \left[ -\frac{\|u\|^2}{2\sigma^2} \right], \quad (21) \\
f_{Y\mid U}(y_{\text{obs}}, u) &= \frac{|W|^{1/2}}{(2\pi\sigma^2)^{(n+q)/2}} \exp \left[ -\frac{r^2(\theta, \beta, u)}{2\sigma^2} \right], \quad (22) \\
f_{U\mid Y}(u\mid y_{\text{obs}}) &= \frac{f_{Y\mid U}(y_{\text{obs}}, u)}{f_Y(y_{\text{obs}})}, \quad (23)
\end{align*}
\]

where

\[
f_Y(y_{\text{obs}}) = \int f_{Y\mid U}(y_{\text{obs}}, u) du. \quad (25)
\]

The log-likelihood to be maximized can therefore be expressed as,

\[
L(\theta, \beta, \sigma^2\mid y_{\text{obs}}) = \log f_Y(y_{\text{obs}}). \quad (26)
\]

The integral in Equation 25 may be more explicitly written as,

\[
f_Y(y_{\text{obs}}) = \frac{|W|^{1/2}}{(2\pi\sigma^2)^{(n+q)/2}} \exp \left[ -\frac{r^2(\theta) - \frac{\|L_\theta^\top (u - \mu_{U\mid Y=y_{\text{obs}}}) + R_Z X (\beta - \hat{\beta}_\theta)\|^2}{2\sigma^2}}{2\sigma^2} \right] \\
\int \exp \left[ -\frac{\|v\|^2}{2\sigma^2} \right] \left| L_\theta \right|^{-1} dv, \quad (27)
\]

which can be evaluated with the change of variables,

\[
v = L_\theta^\top (u - \mu_{U\mid y_{\text{obs}}}) + R_Z X (\beta - \hat{\beta}_\theta). \quad (28)
\]

The Jacobian determinant of the transformation from \(u\) to \(v\) is \(\left| L_\theta \right|\). Therefore we are able to write the integral as,

\[
f_Y(y_{\text{obs}}) = \frac{|W|^{1/2}}{(2\pi\sigma^2)^{(n+q)/2}} \exp \left[ -\frac{r^2(\theta) - \frac{\|R_Z X (\beta - \hat{\beta}_\theta)\|^2}{2\sigma^2}}{2\sigma^2} \right] \\
\int \exp \left[ -\frac{\|v\|^2}{2\sigma^2} \right] \left| L_\theta \right|^{-1} dv, \quad (29)
\]

which by the properties of exponential integrands becomes,

\[
\exp L(\theta, \beta, \sigma^2\mid y_{\text{obs}}) = f_Y(y_{\text{obs}}) = \frac{|W|^{1/2} \left| L_\theta \right|^{-1}}{(2\pi\sigma^2)^{n/2}} \exp \left[ -\frac{r^2(\theta) - \frac{\|R_Z X (\beta - \hat{\beta}_\theta)\|^2}{2\sigma^2}}{2\sigma^2} \right]. \quad (30)
\]

\(^5\)These expressions only technically hold at the observed value, \(y_{\text{obs}}\), of the response vector, \(Y\).
3.4. Evaluating and profiling the deviance and the REML criterion

We are now in a position to understand why the formulation in Equations 2 and 3 is particularly useful. We are able to explicitly profile $\beta$ and $\sigma$ out of the log-likelihood (Equation 26), to find a compact expression for the profiled deviance (negative twice the profiled log-likelihood) and the profiled REML criterion as a function of the relative covariance parameters, $\theta$, only. Furthermore these criteria can be evaluated quickly and accurately.

To estimate the parameters, $\theta$, $\beta$, and $\sigma^2$, we minimize negative twice the log-likelihood, which can be written as,

$$-2\mathcal{L}(\theta, \beta, \sigma^2 | y_{obs}) = \log \left| \frac{L_\theta}{W} \right|^2 + n \log(2\pi\sigma^2) + \frac{r^2(\theta)}{\sigma^2} + \frac{\|R_x(\beta - \hat{\beta}_\theta)\|^2}{\sigma^2}. \quad (31)$$

It is very easy to profile out $\beta$, because it enters into the ML criterion only through the final term, which is zero if $\beta = \hat{\beta}_\theta$, where $\hat{\beta}_\theta$ is found by solving the penalized least-squares problem in Equation 16. Therefore we can write a partially profiled ML criterion as,

$$-2\mathcal{L}(\theta, \sigma^2 | y_{obs}) = \log \left| \frac{L_\theta}{W} \right|^2 + n \log(2\pi\sigma^2) + \frac{r^2(\theta)}{\sigma^2}. \quad (32)$$

This criterion is only partially profiled because it still depends on $\sigma^2$. Differentiating this criterion with respect to $\sigma^2$ and setting the result equal to zero yields,

$$0 = \frac{n}{\hat{\sigma}_\theta^2} - \frac{r^2(\theta)}{\hat{\sigma}_\theta^2}, \quad (33)$$

which leads to a maximum profiled likelihood estimate,

$$\hat{\sigma}_\theta^2 = \frac{r^2(\theta)}{n}. \quad (34)$$

This estimate can be substituted into the partially profiled criterion to yield the fully profiled ML criterion,

$$-2\mathcal{L}(\theta | y_{obs}) = \log \left| \frac{L_\theta}{W} \right|^2 + n \left[ 1 + \log \left( \frac{2\pi r^2(\theta)}{n} \right) \right]. \quad (35)$$

This expression for the profiled deviance depends only on $\theta$. Although $q$, the number of columns in $Z$ and the size of $\Sigma_\theta$, can be very large indeed, the dimension of $\theta$ is small, frequently less than 10. The lme4 package uses generic nonlinear optimizers (Section 4) to optimize this expression over $\theta$ to find its maximum likelihood estimate.

The REML criterion

The REML criterion can be obtained by integrating the marginal density for $Y$ with respect to the fixed effects (Laird and Ware 1982),

$$\int f_Y(y_{obs})d\beta = \left| W^{1/2} \right|^{-1} \exp \left[ -\frac{r^2(\theta)}{2\sigma^2} \right] \int \exp \left[ -\frac{\|R_x(\beta - \hat{\beta}_\theta)\|^2}{2\sigma^2} \right] d\beta, \quad (36)$$
which can be evaluated with the change of variables,
\[ v = R_X (\beta - \hat{\beta}_\theta). \] (37)

The Jacobian determinant of the transformation from \( \beta \) to \( v \) is \( |R_X| \). Therefore we are able to write the integral as,
\[ \int f_y(y_{obs})d\beta = |W|^{1/2} |L_\theta|^{-1} \exp \left[ -r^2(\theta) \right] \int \exp \left[ -\frac{\|v\|^2}{2\sigma^2} \right] |R_X|^{-1} dv, \] (38)

which simplifies to,
\[ \int f_y(y_{obs})d\beta = \frac{|W|^{1/2} |L_\theta|^{-1} |R_X|^{-1}}{(2\pi\sigma^2)^{(n-p)/2}} \exp \left[ -\frac{r^2(\theta)}{2\sigma^2} \right]. \] (39)

Minus twice the log of this integral is the (unprofiled) REML criterion,
\[ -2\mathcal{L}_R(\theta, \sigma^2 | y_{obs}) = \log \frac{|L_\theta|^2 |R_X|^2}{|W|} + (n-p) \log(2\pi\sigma^2) + \frac{r^2(\theta)}{\sigma^2}. \] (40)

Note that because \( \beta \) gets integrated out, the REML criterion cannot be used to find a point estimate of \( \beta \). However, we follow others in using the maximum likelihood estimate, \( \hat{\beta}_\theta \), at the optimum value of \( \theta = \hat{\theta} \). The REML estimate of \( \sigma^2 \) is,
\[ \hat{\sigma}^2 = \frac{r^2(\hat{\theta})}{n-p}, \] (41)

which leads to the profiled REML criterion,
\[ -2\mathcal{L}_R(\theta | y_{obs}) = \log \frac{|L_\theta|^2 |R_X|^2}{|W|} + (n-p) \left[ 1 + \log \left( \frac{2\pi r^2(\theta)}{n-p} \right) \right]. \] (42)

### 3.5. Changes relative to previous formulations

We compare the PLS problem as formulated in Section 3.2 with earlier versions and describe why we use this version. What have become known as “Henderson’s mixed-model equations” are given as Equation 6 of Henderson Jr. (1982) and would be expressed as,
\[ \begin{bmatrix} X^TX/\sigma^2 & X^TZ/\sigma^2 \\ Z^TX/\sigma^2 & Z^TZ/\sigma^2 + \Sigma^{-1} \end{bmatrix} \begin{bmatrix} \hat{\beta}_\theta \\ \mu_{B|y=y_{obs}} \end{bmatrix} = \begin{bmatrix} X^Ty_{obs}/\sigma^2 \\ Z^Ty_{obs}/\sigma^2 \end{bmatrix}, \] (43)
in our notation (ignoring weights and offsets, without loss of generality). The matrix written as \( R \) in Henderson Jr. (1982) is \( \sigma^2 I_n \) in our formulation of the model. Bates and DebRoy (2004) modified the PLS equations to
\[ \begin{bmatrix} Z^TZ + \Omega & Z^TX \\ X^TZ & X^TX \end{bmatrix} \begin{bmatrix} \mu_{B|y=y_{obs}} \\ \hat{\beta}_\theta \end{bmatrix} = \begin{bmatrix} Z^Ty_{obs} \\ X^Ty_{obs} \end{bmatrix}. \] (44)
where $\mathbf{\Omega}_\theta = \left(\Lambda_\theta^\top \Lambda_\theta\right)^{-1} = \sigma^2 \mathbf{\Sigma}^{-1}$ is the relative precision matrix for a given value of $\theta$. They also showed that the profiled log-likelihood can be expressed (on the deviance scale) as

$$-2\mathcal{L}(\theta) = \log \left(\frac{\mathbf{Z}^\top \mathbf{Z} + \mathbf{\Omega}}{|\mathbf{\Omega}|}\right) + n \left[1 + \log \left(\frac{2\pi r_\theta^2}{n}\right)\right]. \quad (45)$$

The primary difference between Equation 43 and Equation 44 is the order of the blocks in the system matrix. The PLS problem can be solved using a Cholesky factor of the system matrix

The primary difference between Equation 43 and Equation 44 is the order of the blocks in the system matrix. The PLS problem can be solved using a Cholesky factor of the system matrix with the blocks in either order. The advantage of using the arrangement in Equation 44 is to allow for evaluation of the profiled log-likelihood. To evaluate $|\mathbf{Z}^\top \mathbf{Z} + \mathbf{\Omega}|$ from the Cholesky factor that block must be in the upper-left corner, not the lower right. Also, $\mathbf{Z}$ is sparse whereas $\mathbf{X}$ is usually dense. It is straightforward to exploit the sparsity of $\mathbf{Z}^\top \mathbf{Z}$ in the Cholesky factorization when the block containing this matrix is the first block to be factored. If $\mathbf{X}^\top \mathbf{X}$ is the first block to be factored, it is much more difficult to preserve sparsity.

The main change from the formulation in Bates and DebRoy (2004) to the current formulation is the use of a relative covariance factor, $\Lambda_\theta$, instead of a relative precision matrix, $\mathbf{\Omega}_\theta$, and solving for the mean of $\mathbf{U} | \mathbf{Y} = \mathbf{y}_{\text{obs}}$ instead of the mean of $\mathbf{B} | \mathbf{Y} = \mathbf{y}_{\text{obs}}$. This change improves stability, because the solution to the PLS problem in Section 3.2 is well-defined when $\Lambda_\theta$ is singular whereas the formulation in Equation 44 cannot be used in these cases because $\mathbf{\Omega}_\theta$ does not exist.

It is important to allow for $\Lambda_\theta$ to be singular because situations where the parameter estimates, $\theta$, produce a singular $\Lambda_\theta$ do occur in practice. And even if the parameter estimates do not correspond to a singular $\Lambda_\theta$, it may be desirable to evaluate the estimation criterion at such values during the course of the numerical optimization of the criterion.

Bates and DebRoy (2004) also provided expressions for the gradient of the profiled loglikelihood expressed as Equation 45. These expressions can be translated into the current formulation. From Equation 35 we can see that (again ignoring weights),

$$\nabla \left(-2\mathcal{L}(\theta)\right) = \nabla \log(|\mathbf{L}_\theta|^2) + \nabla \left(n \log(r^2(\theta))\right)$$

$$= \nabla \log(|\Lambda_\theta^\top \mathbf{Z} \mathbf{Z} \Lambda_\theta + \mathbf{I}|) + n \left(\nabla r^2(\theta) / r^2(\theta)\right) \quad (46)$$

The first gradient is easy to express but difficult to evaluate for the general model. The individual elements of this gradient are

$$\frac{\partial \log(|\Lambda_\theta^\top \mathbf{Z} \mathbf{Z} \Lambda_\theta + \mathbf{I}|)}{\partial \theta_i} = \text{tr} \left[\frac{\partial \left(\mathbf{L}_\theta^\top \mathbf{Z} \mathbf{Z} \Lambda_\theta + \mathbf{I}\right)}{\partial \theta_i} \left(\mathbf{L}_\theta^\top \mathbf{Z} \mathbf{Z} \Lambda_\theta + \mathbf{I}\right)^{-1}\right]$$

$$= \text{tr} \left[\left(\mathbf{Z}^\top \mathbf{Z} \frac{\partial \Lambda_\theta}{\partial \theta_i} + \frac{\partial \Lambda_\theta}{\partial \theta_i} \mathbf{Z}^\top \mathbf{Z} \Lambda_\theta\right) \left(\mathbf{Z}^\top \mathbf{Z} \Lambda_\theta + \mathbf{I}\right)^{-1}\right]. \quad (47)$$

The second gradient term can be expressed as a linear function of the residual, with individual elements of the form

$$\frac{\partial r^2(\theta)}{\partial \theta_i} = -2u \frac{\partial \Lambda_\theta}{\partial \theta_i} \mathbf{Z}^\top \left(\mathbf{y} - \mathbf{Z} \Lambda_\theta \mathbf{u} - \mathbf{X} \hat{\beta}_\theta\right), \quad (48)$$
### Table 5: Inputs into a linear mixed model.

<table>
<thead>
<tr>
<th>Name/description</th>
<th>Pseudocode</th>
<th>Math</th>
<th>Type</th>
</tr>
</thead>
<tbody>
<tr>
<td>Mapping from covariance parameters to relative co-</td>
<td>mapping</td>
<td></td>
<td>function</td>
</tr>
<tr>
<td>variance factor</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Response vector</td>
<td>y</td>
<td>$y_{\text{obs}}$ (Section 1.1)</td>
<td>double vector</td>
</tr>
<tr>
<td>Fixed-effects model matrix</td>
<td>X</td>
<td>$X$ (Equation 2)</td>
<td>double dense matrix</td>
</tr>
<tr>
<td>Transposed random-effects model matrix</td>
<td>$Z^\top$</td>
<td>$Z^\top$ (Equation 2)</td>
<td>double sparse matrix</td>
</tr>
<tr>
<td>Square-root weights matrix</td>
<td>sqrtW</td>
<td>$W^{1/2}$ (Equation 2)</td>
<td>double diagonal matrix</td>
</tr>
<tr>
<td>Offset</td>
<td>offset</td>
<td>$o$ (Equation 2)</td>
<td>double vector</td>
</tr>
</tbody>
</table>

*In previous versions of *lme4* a sparse $X$ matrix, useful for models with categorical fixed-effect predictors with many levels, could be specified; this feature is not currently available.

3.6. Penalized least squares algorithm

For efficiency, in *lme4* itself, PLS is implemented in compiled C++ code using the *Eigen* (Guennebaud, Jacob, and and others 2015) templated C++ package for numerical linear algebra. Here however, in order to improve readability we describe a version in pure R. Section 3.7 provides greater detail on the techniques and concepts for computational efficiency, which is important in cases where the nonlinear optimizer (Section 4) requires many iterations.

The PLS algorithm takes a vector of covariance parameters, $\theta$, as inputs and returns the profiled deviance (Equation 35) or the REML criterion (Equation 42). This PLS algorithm consists of four main steps:

1. Update the relative covariance factor (Section 3.6.1).
2. Solve the normal equations (Section 3.6.2).
3. Update the linear predictor and residuals (Section 3.6.3).
4. Compute and return the profiled deviance (Section 3.6.4).

PLS also requires the objects described in Table 5, which define the structure of the model. These objects do not get updated during the PLS iterations, and so it is useful to store various matrix products involving them (Table 6). Table 7 lists the objects that do get updated over the PLS iterations. The symbols in this table correspond to a version of *lme4* that is implemented entirely in R (i.e., no compiled code as in *lme4* itself). This implementation is called *lme4pureR* and is currently available on Github ([https://github.com/lme4/lme4pureR/](https://github.com/lme4/lme4pureR/)).

**PLS step I: Update relative covariance factor**
The first step of PLS is to update the relative covariance factor, $\Lambda_\theta$, from the current value of the covariance parameter vector, $\theta$. The updated $\Lambda_\theta$ is then used to update the random-effects Cholesky factor, $L_\theta$ (Equation 18). The mapping from the covariance parameters to the relative covariance factor can take many forms, but in general involves a function that takes $\theta$ into the values of the nonzero elements of $\Lambda_\theta$.

If $\Lambda_\theta$ is stored as an object of class ‘dgCMatrix’ from the Matrix package for R, then we may update $\Lambda_\theta$ from $\theta$ by,

```r
> Lambdat@x[] <- mapping(theta)
```

where `mapping` is an R function that both accepts and returns a numeric vector. The nonzero elements of sparse matrix classes in Matrix are stored in a slot called `x`.

In the current version of lme4 (v. 1.1-7) the mapping from $\theta$ to $\Lambda_\theta$ is represented as an R integer vector, Lind, of indices, so that

```r
> mapping <- function(theta) theta[Lind]
```

The index vector Lind is computed during the model setup and stored in the function’s environment. Continuing the example from Section 2.3.2, consider a new value for `theta`,

```r
> thetanew <- c(1, -0.1, 2, 0.1, -0.2, 3)
```

To put these values in the appropriate elements in `Lambdati`, we use `mapping`,

```r
> Lambdati@x[] <- mapping(thetanew)
> Lambdati
```

6 x 6 sparse Matrix of class "dgTMatrix"

```
[1,]  1 -0.1  2.0 . . .
[2,] .  0.1 -0.2 . . .
[3,] . .  3.0 . . .
[4,] . . .  1 -0.1  2.0
[5,] . . . .  0.1 -0.2
[6,] . . . . .  3.0
```

This Lind approach can be useful for extending the capabilities of lme4 by using the modular approach to fitting mixed models. For example, Appendix A.1 shows how to use Lind to fit
Table 7: Quantities updated during an evaluation of the linear mixed model objective function.

- **Relative covariance factor**: \( \mathbf{\Lambda}_\theta \) (Equation 4), double sparse lower-triangular matrix
- **Random-effects Cholesky factor**: \( \mathbf{L}_\theta \) (Equation 18), double sparse triangular matrix
- **Intermediate vector in the solution of the normal equations**: \( \mathbf{c}_u \) (Equation 49), double vector
- **Block in the full Cholesky factor**: \( \mathbf{R}_{ZX} \) (Equation 18), double dense matrix
- **Cross-product of the fixed-effects Cholesky factor**: \( \mathbf{R}_X^T \mathbf{R}_X \) (Equation 51), double dense matrix
- **Fixed-effects coefficients**: \( \mathbf{\beta} \) (Equation 2), double vector
- **Spherical conditional modes**: \( \mathbf{u} \) (Section 3.1), double vector
- **Non-spherical conditional modes**: \( \mathbf{b} \) (Equation 2), double vector
- **Linear predictor**: \( \mathbf{\mu} = y_{\text{fit}} \) (Equation 13), double vector
- **Weighted residuals**: \( \mathbf{W}^{1/2}(\mathbf{y}_{\text{obs}} - \mathbf{\mu}) \), double vector
- **Penalized weighted residual sum-of-squares**: \( r^2(\theta) \) (Equation 19), double
- **Twice the log determinant random-effects Cholesky factor**: \( \log |\mathbf{L}_\theta|^2 \), double

A model where two random slopes are uncorrelated, but both slopes are correlated with an intercept.

The mapping from the covariance parameters to the relative covariance factor is treated differently in other implementations of the \texttt{lme4} approach to linear mixed models. At the other extreme, the \texttt{flexLambda} branch of \texttt{lme4} and the \texttt{lme4pureR} package provide the capabilities for a completely general mapping. This added flexibility has the advantage of allowing a much wider variety of models (e.g., compound symmetry, auto-regression). However, the disadvantage of this approach is that it becomes possible to fit a much wider variety of ill-posed models. Finally, if one would like such added flexibility with the current stable version of \texttt{lme4}, it is always possible to use the modular approach to wrap the \texttt{Lind} based deviance function in a general mapping function taking a parameter to be optimized, say \( \phi \), into \( \theta \). However, this approach is likely to be inefficient in many cases.

The \texttt{update} method from the \texttt{Matrix} package efficiently updates the random-effects Cholesky factor, \( \mathbf{L}_\theta \), from a new value of \( \theta \) and the updated \( \mathbf{\Lambda}_\theta \).

\[
> \mathbf{L} \leftarrow \text{update}(\mathbf{L}, \text{Lambdat} \%*\% \mathbf{ZtW}, \text{mult} = 1)
\]

The \texttt{mult = 1} argument corresponds to the addition of the identity matrix to the upper-left block on the left-hand-side of Equation 18.
PLS step II: solve normal equations

With the new covariance parameters installed in $\Lambda_\theta$, the next step is to solve the normal equations (Equation 17) for the current estimate of the fixed-effects coefficients, $\hat{\beta}_\theta$, and the conditional mode, $\mu_{U \mid Y=Y_{\text{obs}}}$. We solve these equations using a sparse Cholesky factorization (Equation 18). In a complex model fit to a large data set, the dominant calculation in the evaluation of the profiled deviance (Equation 35) or REML criterion (Equation 42) is this sparse Cholesky factorization (Equation 18). The factorization is performed in two phases; a symbolic phase and a numeric phase. The symbolic phase, in which the fill-reducing permutation $P$ is determined along with the positions of the nonzeros in $L_\theta$, does not depend on the value of $\theta$. It only depends on the positions of the nonzeros in $ZA_\theta$. The numeric phase uses $\theta$ to determine the numeric values of the nonzeros in $L_\theta$. Using this factorization, the solution proceeds by the following steps,

\begin{align}
L_\theta c_u &= PA_\theta^T Z^T W y \\
L_\theta R_{ZX} &= PA_\theta^T Z^T W X \\
R_X^T R_X &= X^T W X - R_{ZX}^T R_{ZX} \\
(R_X^T R_X) \hat{\beta}_\theta &= X^T W y - R_{ZX} c_u \\
L_\theta^T P u &= c_u - R_{ZX} \hat{\beta}_\theta
\end{align}

which can be solved using the Matrix package with,

\begin{verbatim}
> cu[] <- as.vector(solve(L, solve(L, Lambdat %*% ZtWy, system = "P"), system = "L"))
> RZX[] <- as.vector(solve(L, solve(L, Lambdat %*% ZtWX, system = "P"), system = "L"))
> RXtRX <- as(XtWX - crossprod(RZX), "dpoMatrix")
> beta[] <- as.vector(solve(RXtRX, XtWy - crossprod(RZX, cu)))
> u[] <- as.vector(solve(L, solve(L, cu - RZX %*% beta, system = "Lt"), system = "Pt"))
\end{verbatim}

Notice the nested calls to solve. The inner calls of the first two assignments determine and apply the permutation matrix (system = "P"), whereas the outer calls actually solve the equation (system = "L"). In the assignment to u[], the nesting is reversed in order to return to the original permutation.

PLS step III: Update linear predictor and residuals

The next step is to compute the linear predictor, $\mu_{Y\mid U}$ (Equation 13), and the weighted residuals with new values for $\hat{\beta}_\theta$ and $\mu_{B\mid Y=Y_{\text{obs}}}$. In lme4pureR these quantities can be computed as,

\begin{verbatim}
> b[] <- as.vector(crossprod(Lambdat, u))
> mu[] <- as.vector(crossprod(Zt, b) + X %*% beta + offset)
> wtres <- sqrtW * (y - mu)
\end{verbatim}

where b represents the current estimate of $\mu_{B\mid Y=Y_{\text{obs}}}$. 
PLS step IV: Compute profiled deviance

Finally, the updated linear predictor and weighted residuals can be used to compute the profiled deviance (or REML criterion), which in \texttt{lme4pureR} proceeds as,

```r
> pwrss <- sum(wtres^2) + sum(u^2)
> logDet <- 2*determinant(L, logarithm = TRUE)$modulus
> if (REML) logDet <- logDet + determinant(RXtRX, logarithm = TRUE)$modulus
> attributes(logDet) <- NULL
> profDev <- logDet + degFree * (1 + log(2 * pi * pwrss) - log(degFree))
```

The profiled deviance consists of three components: (1) log-determinant(s) of Cholesky factorization (\texttt{logDet}), (2) the degrees of freedom (\texttt{degFree}), and the penalized weighted residual sum-of-squares (\texttt{pwrss}).

3.7. Sparse matrix methods

In fitting linear mixed models, an instance of the PLS problem (17) must be solved at each evaluation of the objective function during the optimization (Section 4) with respect to $\theta$. Because this operation must be performed many times it is worthwhile considering how to provide effective evaluation methods for objects and calculations involving the sparse matrices associated with random-effects terms (Sections 2.3).

The \texttt{CHOLMOD} library of C functions (Chen, Davis, Hager, and Rajamanickam 2008), on which the sparse matrix capabilities of the \texttt{Matrix} package for R and the sparse Cholesky factorization in \texttt{Julia} are based, allows for separation of the symbolic and numeric phases. Thus we perform the symbolic phase as part of establishing the structure representing the model (Section 2). Furthermore, because \texttt{CHOLMOD} functions allow for updating $L_\theta$ directly from the matrix $\Lambda_\theta Z^T$ without actually forming $\Lambda_\theta Z^T Z \Lambda_\theta + I$ we generate and store $Z^T$ instead of $Z$ (note that we have ignored the weights matrix, $W$, for simplicity). We can update $\Lambda_\theta^T Z^T$ directly from $\theta$ without forming $\Lambda_\theta$ and multiplying two sparse matrices. Although such a direct approach is used in the \texttt{MixedModels} package for Julia, in \texttt{lme4} we first update $\Lambda_\theta^T Z^T$ then form the sparse product $\Lambda_\theta Z^T$. A third alternative, which we employ in \texttt{lme4pureR}, is to compute and save the cross-products of the model matrices, $X$ and $Z$, and the response, $y$, before starting the iterations. To allow for case weights, we save the products $X^T W X$, $X^T W y$, $Z^T W X$, $Z^T W y$ and $Z^T W Z$ (see Table 6).

We wish to use structures and algorithms that allow us to take a new value of $\theta$ and evaluate the $L_\theta$ (Equation 18) efficiently. The key to doing so is the special structure of $\Lambda_\theta Z^T W^{1/2}$. To understand why this matrix, and not its transpose, is of interest we describe the sparse matrix structures used in Julia and in the \texttt{Matrix} package for R.

Dense matrices are stored in R and in Julia as a one-dimensional array of contiguous storage locations addressed in \texttt{column-major order}. This means that elements in the same column are in adjacent storage locations whereas elements in the same row can be widely separated in memory. For this reason, algorithms that work column-wise are preferred to those that work row-wise.

Although a sparse matrix can be stored in a \texttt{triplet} format, where the row position, column position and element value of the nonzeros are recorded as triplets, the preferred storage
forms for actual computation with sparse matrices are compressed sparse column (CSC) or compressed sparse row (CSR, Davis 2006, Chapter 2). The CHOLMOD (and, more generally, the SuiteSparse package of C libraries; Davis et al. 2015) uses the CSC storage format. In this format the nonzero elements in a column are in adjacent storage locations and access to all the elements in a column is much easier than access to those in a row (similar to dense matrices stored in column-major order).

The matrices $Z$ and $Z\Lambda_\theta$ have the property that the number of nonzeros in each row, $\sum_{i=1}^k p_i$, is constant. For CSC matrices we want consistency in the columns rather than the rows, which is why we work with $Z^\top$ and $\Lambda_\theta Z^\top$ rather than their transposes.

An arbitrary $m \times n$ sparse matrix in CSC format is expressed as two vectors of indices, the row indices and the column pointers, and a numeric vector of the nonzero values. The elements of the row indices and the nonzeros are aligned and are ordered first by increasing column number then by increasing row number within column. The column pointers are a vector of size $n + 1$ giving the location of the start of each column in the vectors of row indices and nonzeros.

Because the number of nonzeros in each column of $Z^\top$, and in each column of matrices derived from $Z^\top$ (such as $\Lambda_\theta^k Z^\top W^{1/2}$) is constant, the vector of nonzeros in the CSC format can be viewed as a dense matrix, say $N$, of size $(\sum_{i=1}^n p_i) \times n$. We do not need to store the column pointers because the columns of $Z^\top$ correspond to columns of $N$. All we need is $N$, the dense matrix of nonzeros, and the row indices, which are derived from the grouping factor index vectors $i_i$, $i = 1, \ldots, k$ and can also be arranged as a dense matrix of size $(\sum_{i=1}^n p_i) \times n$. Matrices like $Z^\top$, with the property that there are the same number of nonzeros in each column, are sometimes called regular sparse column-oriented (RSC) matrices.

### 4. Nonlinear optimization module

The objective function module produces a function which implements the penalized least squares algorithm for a particular mixed model. The next step is to optimize this function over the covariance parameters, $\theta$, which is a nonlinear optimization problem. The lme4 package separates the efficient computational linear algebra required to compute profiled likelihoods and deviances for a given value of $\theta$ from the nonlinear optimization algorithms, which use general-purpose nonlinear optimizers.

One benefit of this separation is that it allows for experimentation with different nonlinear optimizers. Throughout the development of lme4, the default optimizers and control parameters have changed in response to feedback from users about both efficiency and convergence properties. lme4 incorporates two built-in optimization choices, an implementation of the Nelder-Mead simplex algorithm adapted from Steven Johnson’s NLopt C library (Johnson 2014) and a wrapper for Powell’s BOBYQA algorithm, implemented in the minqa package (Bates, Mullen, Nash, and Varadhan 2014b) as a wrapper around Powell’s original Fortran code (Powell 2009). More generally, lme4 allows any user-specified optimizer that (1) can work with an objective function (i.e., does not require a gradient function to be specified), (2) allows box constraints on the parameters, and (3) conforms to some simple rules about argument names and structure of the output. An internal wrapper allows the use of the optimx package (Nash and Varadhan 2011), although the only optimizers provided via optimx that satisfy the constraints above are the nlminb and L-BFGS-B algorithms that are then-
selves wrappers around the versions provided in base R. Several other algorithms from Steven Johnson’s NLOpt package are also available via the nloptr wrapper package (e.g., alternate implementations of Nelder-Mead and BOBYQA, and Powell’s COBYLA algorithm).

This flexibility assists with diagnosing convergence problems – it is easy to switch among several algorithms to determine whether the problem lies in a failure of the nonlinear optimization stage, as opposed to a case of model misspecification or unidentifiability or a problem with the underlying PLS algorithm. To date we have only observed PLS failures, which arise if $X^T W X - R_{ZX}^T R_{ZX}$ becomes singular during an evaluation of the objective function, with badly scaled problems (i.e., problems with continuous predictors that take a very large or very small numerical range of values).

The requirement for optimizers that can handle box constraints stems from our decision to parameterize the variance-covariance matrix in a constrained space, in order to allow for singular fits. In contrast to the approach taken in the nlme package (Pinheiro et al. 2014), which goes to some lengths to use an unconstrained variance-covariance parameterization (the log-Cholesky parameterization; Pinheiro and Bates 1996), we instead use the Cholesky parameterization but require the elements of $\theta$ corresponding to the diagonal elements of the Cholesky factor to be non-negative. With these constraints, the variance-covariance matrix is singular if and only if any of the diagonal elements is exactly zero. Singular fits are common in practical data-analysis situations, especially with small- to medium-sized data sets and complex variance-covariance models, so being able to fit a singular model is an advantage: when the best fitting model lies on the boundary of a constrained space, approaches that try to remove the constraints by fitting parameters on a transformed scale will often give rise to convergence warnings as the algorithm tries to find a maximum on an asymptotically flat surface (Bolker et al. 2013).

In principle the likelihood surfaces we are trying to optimize over are smooth, but in practice using gradient information in optimization may or may not be worth the effort. In special cases, we have a closed-form solution for the gradients (Equations 46–48), but in general we will have to approximate them by finite differences, which is expensive and has limited accuracy. (We have considered using automatic differentiation approaches to compute the gradients more efficiently, but this strategy requires a great deal of additional machinery, and would have drawbacks in terms of memory requirements for large problems.) This is the primary reason for our switching to derivative-free optimizers such as BOBYQA and Nelder-Mead in the current version of lme4, although as discussed above derivative-based optimizers based on finite differencing are available as an alternative.

There is most likely further room for improvement in the nonlinear optimization module; for example, some speed-up could be gained by using parallel implementations of derivative-free optimizers that evaluated several trial points at once (Klein and Neira 2013). In practice users most often have optimization difficulties with poorly scaled or centered data – we are working to implement appropriate diagnostic tests and warnings to detect these situations.

### 5. Output module

Here we describe some of the methods in lme4 for exploring fitted linear mixed models (Table 8), which are represented as objects of the S4 class ‘lmerMod’. We begin by describing the theory underlying these methods (Section 5.1) and then continue the sleepstudy example
introduced in Section 1.2 to illustrate these ideas in practice.

5.1. Theory underlying the output module

Covariance matrix of the fixed-effect coefficients

In the \texttt{lm} function, the variance-covariance matrix of the coefficients is the inverse of the cross-product of the model matrix, times the residual variance (Chambers 1993). The inverse cross-product matrix is computed by first inverting the upper triangular matrix resulting from the QR decomposition of the model matrix, and then taking its cross-product,

$$\text{Var}_{\theta, \sigma} \left( \begin{bmatrix} \hat{\mu}_{i|y=y_{\text{obs}}} \\ \hat{\beta} \end{bmatrix} \right) = \sigma^2 \begin{bmatrix} L_\theta^T & R_{ZX} \\ 0 & R_X \end{bmatrix}^{-1} \begin{bmatrix} L_\theta & 0 \\ R_{ZX}^T & R_X^T \end{bmatrix}^{-1}. \quad (54)$$

Because of normality, the marginal covariance matrix of $\hat{\beta}$ is just the lower-right $p$-by-$p$ block of $\text{Var}_{\theta, \sigma} \left( \begin{bmatrix} \hat{\mu}_{i|y=y_{\text{obs}}} \\ \hat{\beta} \end{bmatrix} \right)$. This lower-right block is

$$\text{Var}_{\theta, \sigma}(\hat{\beta}) = \sigma^2 R_X^{-1}(R_X^T)^{-1}, \quad (55)$$

which follows from the Schur complement identity (Horn and Zhang 2005, Theorem 1.2). This matrix can be extracted from a fitted \texttt{merMod} object as,

```r
> RX <- getME(fm1, "RX")
> sigma2 <- sigma(fm1)^2
> sigma2 * chol2inv(RX)
```

which could be computed with \texttt{lme4} as \texttt{vcov(fm1)}.

The square-root diagonal of this covariance matrix contains the estimates of the standard errors of fixed-effects coefficients. These standard errors are used to construct Wald confidence intervals with \texttt{confint(. , method = "Wald")}. Such confidence intervals are approximate, and depend on the assumption that the likelihood profile of the fixed effects is linear on the $\zeta$ scale (Section 5.1.2).

Profiling

The theory of likelihood profiles is straightforward: the deviance (or likelihood) profile, $-2\mathcal{L}_P()$, for a focal model parameter $P$ is the minimum value of the deviance conditioned on a particular value of $P$. For each parameter of interest, our goal is to evaluate the deviance profile for many points – optimizing over all of the non-focal parameters each time – over a wide enough range and with high enough resolution to evaluate the shape of the profile (in particular, whether it is quadratic, which would allow use of Wald confidence intervals and tests) and to find the values of $P$ such that $-2\mathcal{L}_P(P) = -2\mathcal{L}(\hat{P}) + \chi^2(1 - \alpha)$, which represent
the profile confidence intervals. While profile confidence intervals rely on the asymptotic distribution of the minimum deviance, this is a much weaker assumption than the log-quadratic likelihood surface required by Wald tests.

An additional challenge in profiling arises when we want to compute profiles for quantities of interest that are not parameters of our PLS function. We have two problems in using the deviance function defined above to profile linear mixed models. First, a parameterization of the random-effects variance-covariance matrix in terms of standard deviations and correlations, or variances and covariances, is much more familiar to users, and much more relevant as output of a statistical model, than the parameters, \( \theta \), of the relative covariance factor – users are interested in inferences on variances or standard deviations, not on \( \theta \). Second, the fixed-effect coefficients and the residual standard deviation, both of which are also of interest to users, are profiled out (in the sense used above) of the deviance function (Section 3.4), so we have to find a strategy for estimating the deviance for values of \( \beta \) and \( \sigma^2 \) other than the profiled values.

To handle the first problem we create a new version of the deviance function that first takes a vector of standard deviations (and correlations), and a value of the residual standard deviation, maps them to a \( \theta \) vector, and then computes the PLS as before; it uses the specified residual standard deviation rather than the PLS estimate of the standard deviation (Equation 34) in the deviance calculation. We compute a profile likelihood for the fixed-effect parameters, which are profiled out of the deviance calculation, by adding an offset to the linear predictor for the focal element of \( \beta \). The resulting function is not useful for general nonlinear optimization – one can easily wander into parameter regimes corresponding to infeasible (non-positive semidefinite) variance-covariance matrices – but it serves for likelihood profiling, where one focal parameter is varied at a time and the optimization over the other parameters is likely to start close to an optimum.

In practice, the profile method systematically varies the parameters in a model, assessing the best possible fit that can be obtained with one parameter fixed at a specific value and comparing this fit to the globally optimal fit, which is the original model fit that allowed all the parameters to vary. The models are compared according to the change in the deviance, which is the likelihood ratio test statistic. We apply a signed square root transformation to this statistic and plot the resulting function, which we call the profile zeta function or \( \zeta \), versus the parameter value. The signed aspect of this transformation means that \( \zeta \) is positive where the deviation from the parameter estimate is positive and negative otherwise, leading to a monotonically increasing function which is zero at the global optimum. A \( \zeta \) value can be compared to the quantiles of the standard normal distribution. For example, a 95% profile deviance confidence interval on the parameter consists of the values for which \(-1.96 < \zeta < 1.96\). Because the process of profiling a fitted model can be computationally intensive – it involves refitting the model many times – one should exercise caution with complex models fit to large data sets.

The standard approach to this computational challenge is to compute \( \zeta \) at a limited number of parameter values, and to fill in the gaps by fitting an interpolation spline. Often one is able to invert the spline to obtain a function from \( \zeta \) to the focal parameter, which is necessary in order to construct profile confidence intervals. However, even if the points themselves are monotonic, it is possible to obtain non-monotonic interpolation curves. In such a case, \texttt{lme4} falls back to linear interpolation, with a warning.
The last part of the technical specification for computing profiles is deciding on a strategy for choosing points to sample. In one way or another one wants to start at the estimated value for each parameter and work outward either until a constraint is reached (i.e., a value of 0 for a standard deviation parameter or a value of ±1 for a correlation parameter), or until a sufficiently large deviation from the minimum deviance is attained. lme4’s profiler chooses a cutoff \( \phi \) based on the \( 1 - \alpha_{\text{max}} \) critical value of the \( \chi^2 \) distribution with a number of degrees of freedom equal to the total number of parameters in the model, where \( \alpha_{\text{max}} \) is set to 0.05 by default. The profile computation initially adjusts the focal parameter \( p_i \) by an amount \( \epsilon = 1.01 \hat{p}_i \) from its ML or REML estimate \( \hat{p}_i \) (or by \( \epsilon = 0.001 \) if \( \hat{p}_i \) is zero, as in the case of a singular variance-covariance model). The local slope of the likelihood profile \( (\zeta(\hat{p}_i + \epsilon) - \zeta(\hat{p}_i))/\epsilon \) is used to pick the next point to evaluate, extrapolating the local slope to find a new \( \epsilon \) that would be expected to correspond to the desired step size \( \Delta \zeta \) (equal to \( \phi/8 \) by default, so that 16 points would be used to cover the profile in both directions if the log-likelihood surface were exactly quadratic). Some fail-safe testing is done to ensure that the step chosen is always positive, and less than a maximum; if a deviance is ever detected that is lower than that of the ML deviance, which can occur if the initial fit was wrong due to numerical problems, the profiler issues an error and stops.

**Parametric bootstrapping**

To avoid the asymptotic assumptions of the likelihood ratio test, at the cost of greatly increased computation time, one can estimate confidence intervals by parametric bootstrapping – that is, by simulating data from the fitted model, refitting the model, and extracting the new estimated parameters (or any other quantities of interest). This task is quite straightforward, since there is already a \texttt{simulate} method, and a \texttt{refit} function which re-estimates the (RE)ML parameters for new data, starting from the previous (RE)ML estimates and re-using the previously computed model structures (including the fill-reducing permutation) for efficiency. The \texttt{bootMer} function is thus a fairly thin wrapper around a \texttt{simulate/refit} loop, with a small amount of additional logic for parallel computation and error-catching. (Some of the ideas of \texttt{bootMer} are adapted from merBoot (Sánchez-Espigares and Ocaña 2009), a more ambitious framework for bootstrapping lme4 model fits which unfortunately seems to be unavailable at present.)

**Conditional variances of random effects**

It is useful to clarify that the conditional covariance concept in lme4 is based on a simplification of the linear mixed model. In particular, we simplify the model by assuming that the quantities, \( \beta \), \( \Lambda \), and \( \sigma \), are known (i.e., set at their estimated values). In fact, the only way to define the conditional covariance is at fixed parameter values. Our approximation here is using the estimated parameter values for the unknown “true” parameter values. In this simplified case, \( U \) is the only quantity in the statistical model that is both random and unknown.

Given this simplified linear mixed model, a standard result in Bayesian linear regression modeling (Gelman et al. 2013) implies that the conditional distribution of the spherical random effects given the observed response vector is Gaussian,

\[
(U | Y = y_{\text{obs}}) \sim \mathcal{N}(\mu_U | Y = y_{\text{obs}}, \sigma^2 V),
\]  

(56)
where

\[ V = \left( \Lambda_\theta^T Z^T W Z \Lambda_\theta + I_z \right)^{-1} = \left( L_{\theta}^{-1} \right)^T \left( L_{\theta}^{-1} \right) \]  

(57)

is the unscaled conditional variance and

\[ \mu_{(Y | \theta = y_{\text{obs}})} = V \Lambda_\theta^T Z^T W \left( y_{\text{obs}} - o - X \hat{\beta} \right) \]  

(58)

is the conditional mean/mode. Note that in practice the inverse in Equation 57 does not get computed directly, but rather an efficient method is used that exploits the sparse structures.

The random-effects coefficient vector, \( b \), is often of greater interest. The conditional covariance matrix of \( B \) may be expressed as

\[ \hat{\sigma}^2 \Lambda_\theta^T V \Lambda_\theta^T. \]  

(59)

**The hat matrix**

The hat matrix, \( H \), is a useful object in linear model diagnostics (Cook and Weisberg 1982). In general, \( H \) relates the observed and fitted response vectors, but we specifically define it as,

\[ \left( \mu_{Y | \theta = o} - o \right) = H \left( y_{\text{obs}} - o \right). \]  

(60)

To find \( H \) we note that

\[ \left( \mu_{Y | \theta = o} - o \right) = \left[ Z \Lambda X \right] \left[ \begin{array}{c} \mu_{(Y | \theta = y_{\text{obs}})} \\ \hat{\beta}_\theta \end{array} \right]. \]  

(61)

Next we get an expression for \( \left[ \mu_{(Y | \theta = y_{\text{obs}})} \right] \hat{\beta}_\theta \) by solving the normal equations (Equation 17),

\[ \begin{bmatrix} \mu_{(Y | \theta = y_{\text{obs}})} \\ \hat{\beta}_\theta \end{bmatrix} = \begin{bmatrix} L_{\theta} & R_{ZX} \\ 0 & R_X \end{bmatrix}^{-1} \begin{bmatrix} L_{\theta} & 0 \\ R_{ZX} & R_X \end{bmatrix}^{-1} \begin{bmatrix} \Lambda_\theta^T Z^T \\ X^T \end{bmatrix} W \left( y_{\text{obs}} - o \right). \]  

(62)

By the Schur complement identity (Horn and Zhang 2005),

\[ \begin{bmatrix} L_{\theta} & R_{ZX} \\ 0 & R_X \end{bmatrix}^{-1} = \begin{bmatrix} \left( L_{\theta} \right)^{-1} & - \left( L_{\theta} \right)^{-1} R_{ZX} R_X^{-1} \\ 0 & R_X^{-1} \end{bmatrix}. \]  

(63)

Therefore, we may write the hat matrix as

\[ H = \left( C_L^T C_L + C_R^T C_R \right), \]  

(64)

where,

\[ L_{\theta} C_L = \Lambda_\theta^T Z^T W^{1/2} \]  

(65)

and,

\[ R_X^T C_R = X^T W^{1/2} - R_{ZX}^T C_L. \]  

(66)

The trace of the hat matrix is often used as a measure of the effective degrees of freedom (e.g., Vaida and Blanchard 2005). Using a relationship between the trace and vec operators, the trace of \( H \) may be expressed as,

\[ \text{tr}(H) = \| \text{vec}(C_L) \|^2 + \| \text{vec}(C_R) \|^2. \]  

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<tr>
<td>df.residual</td>
<td>Residual degrees of freedom.</td>
</tr>
<tr>
<td>drop1</td>
<td>Drop allowable single terms from the model.</td>
</tr>
<tr>
<td>extractAIC</td>
<td>Generalized Akaike information criterion</td>
</tr>
<tr>
<td>fitted</td>
<td>Fitted values given conditional modes (Equation 13).</td>
</tr>
<tr>
<td>fixef (5.2.2)</td>
<td>Estimates of the fixed-effects coefficients, $\hat{\beta}$</td>
</tr>
<tr>
<td>formula (2.2.1)</td>
<td>Mixed-model formula of fitted model.</td>
</tr>
<tr>
<td>logLik</td>
<td>Maximum log-likelihood.</td>
</tr>
<tr>
<td>model.frame</td>
<td>Data required to fit the model.</td>
</tr>
<tr>
<td>model.matrix</td>
<td>Fixed-effects model matrix, $X$.</td>
</tr>
<tr>
<td>ngrps (5.2.2)</td>
<td>Number of levels in each grouping factor.</td>
</tr>
<tr>
<td>nobs (5.2.2)</td>
<td>Number of observations.</td>
</tr>
<tr>
<td>plot</td>
<td>Diagnostic plots for mixed-model fits.</td>
</tr>
<tr>
<td>predict (5.2.8)</td>
<td>Various types of predicted values.</td>
</tr>
<tr>
<td>print</td>
<td>Basic printout of mixed-model objects.</td>
</tr>
<tr>
<td>profile (5.1.2)</td>
<td>Profiled likelihood over various model parameters.</td>
</tr>
<tr>
<td>ranef (5.2.2)</td>
<td>Conditional modes of the random effects.</td>
</tr>
<tr>
<td>refit (5.1.3)</td>
<td>A model (re)fitted to a new set of observations of the response variable.</td>
</tr>
<tr>
<td>refitML (5.2.4)</td>
<td>A model (re)fitted by maximum likelihood.</td>
</tr>
<tr>
<td>residuals (5.2.2)</td>
<td>Various types of residual values.</td>
</tr>
<tr>
<td>sigma (5.2.2)</td>
<td>Residual standard deviation.</td>
</tr>
<tr>
<td>simulate (5.2.8)</td>
<td>Simulated data from a fitted mixed model.</td>
</tr>
<tr>
<td>summary (5.2.2)</td>
<td>Summary of a mixed model.</td>
</tr>
<tr>
<td>terms</td>
<td>Terms representation of a mixed model.</td>
</tr>
<tr>
<td>update (5.2.1)</td>
<td>An updated model using a revised formula or other arguments.</td>
</tr>
<tr>
<td>VarCorr (5.2.2)</td>
<td>Estimated random-effects variances, standard deviations, and correlations.</td>
</tr>
<tr>
<td>vcov (5.2.2)</td>
<td>Covariance matrix of the fixed-effect estimates.</td>
</tr>
<tr>
<td>weights</td>
<td>Prior weights used in model fitting.</td>
</tr>
</tbody>
</table>

Table 8: List of currently available methods for objects of the class ‘merMod’.
5.2. Using the output module

The user interface for the output module largely consists of a set of methods (Table 8) for objects of class `merMod`, which is the class of objects returned by `lmer`. In addition to these methods, the `getME` function can be used to extract various objects from a fitted mixed model in `lme4`. Here we illustrate the use of several of these methods.

Updating fitted mixed models

To illustrate the `update` method for ‘merMod’ objects we construct a random intercept only model. This task could be done in several ways, but we choose to first remove the random-effects term `(Days | Subject)` and add a new term with a random intercept,

```r
> fm3 <- update(fm1, . ~ . - (Days | Subject) + (1 | Subject))
> formula(fm3)

Reaction ~ Days + (1 | Subject)
```

Model summary and associated accessors

The `summary` method for ‘merMod’ objects provides information about the model fit. Here we consider the output of `summary(fm1)` in four steps. The first few lines of output indicate that the model was fitted by REML as well as the value of the REML criterion (Equation 40) at convergence (which may also be extracted using `REMLcrit(fm1)`). The beginning of the summary also reproduces the model formula and the scaled Pearson residuals,

```
Linear mixed model fit by REML ['lmerMod']
Formula: Reaction ~ Days + (Days | Subject)
Data: sleepstudy

REML criterion at convergence: 1743.6

Scaled residuals:
    Min  1Q Median  3Q     Max
-3.954 -0.463  0.023  0.463  5.179
```

This information may also be obtained using standard accessor functions,

```r
> formula(fm1)
> REMLcrit(fm1)
> quantile(residuals(fm1, "pearson", scaled = TRUE))
```

The second piece of the `summary` output provides information regarding the random-effects and residual variation,

```
Random effects:
Groups   Name     Variance Std.Dev. Corr
Subject  (Intercept) 612.1   24.74
```

Days 35.1  5.92  0.07
Residual 654.9  25.59
Number of obs: 180, groups: Subject, 18

which can also be accessed using,

```r
> print(vc <- VarCorr(fm1), comp = c("Variance", "Std.Dev."))
> nobs(fm1)
> ngrps(fm1)
> sigma(fm1)
```

The print method for objects returned by `VarCorr` hides the internal structure of these `VarCorr.merMod` objects. The internal structure of an object of this class is a list of matrices, one for each random-effects term. The standard deviations and correlation matrices for each term are stored as attributes, `stddev` and `correlation`, respectively, of the variance-covariance matrix, and the residual standard deviation is stored as attribute `sc`. For programming use, these objects can be summarized differently using their `as.data.frame` method,

```r
> as.data.frame(VarCorr(fm1))
```

which contains one row for each variance or covariance parameter. The `grp` column gives the grouping factor associated with this parameter. The `var1` and `var2` columns give the names of the variables associated with the parameter (`var2` is `<NA>` unless it is a covariance parameter). The `vcov` column gives the variances and covariances, and the `sdcor` column gives these numbers on the standard deviation and correlation scales.

The next chunk of output gives the fixed-effect estimates,

```r
Fixed effects:
  Estimate Std. Error t value
(Intercept) 251.41   6.82   36.84
  Days      10.47   1.55    6.77
```

Note that there are no p values (see Section 5.2.5). The fixed-effect point estimates may be obtained separately via `fixef(fm1)`. Conditional modes of the random-effects coefficients can be obtained with `ranef` (see Section 5.1.4 for information on the theory). Finally, we have the correlations between the fixed-effect estimates

```r
Correlation of Fixed Effects:
  (Intr)
Days -0.138
```
The full variance-covariance matrix of the fixed-effects estimates can be obtained in the usual R way with the \texttt{vcov} method (Section 5.1.1). Alternatively, \texttt{coef(summary(fm1))} will return the full fixed-effects parameter table as shown in the summary.

**Diagonal plots**

\texttt{lme4} provides tools for generating most of the standard graphical diagnostic plots (with the exception of those incorporating influence measures, e.g., Cook’s distance and leverage plots), in a way modeled on the diagnostic graphics of \texttt{nlme} (Pinheiro and Bates 2000). In particular, the familiar \texttt{plot} method in base R for linear models (objects of class ‘\texttt{lm}’) can be used to create the standard fitted vs. residual plots,

\begin{verbatim}
> plot(fm1, type = c("p", "smooth"))
\end{verbatim}

scale-location plots,

\begin{verbatim}
> plot(fm1, sqrt(abs(resid(.))) ~ fitted(.),
  type = c("p", "smooth"))
\end{verbatim}

and quantile-quantile plots (from \texttt{lattice}),

\begin{verbatim}
> qqmath(fm1, id = 0.05)
\end{verbatim}

In contrast to \texttt{plot.lm}, these scale-location and Q-Q plots are based on raw rather than standardized residuals.

In addition to these standard diagnostic plots, which examine the validity of various assumptions (linearity, homoscedasticity, normality) at the level of the residuals, one can also use the \texttt{dotplot} and \texttt{qqmath} methods for the conditional modes (i.e., ‘\texttt{ranef.mer}’ objects generated by \texttt{ranef(fit)}) to check for interesting patterns and normality in the conditional modes. \texttt{lme4} does not provide influence diagnostics, but these (and other useful diagnostic procedures) are available in the dependent packages \texttt{HLMdiag} and \texttt{influence.ME} (Loy and Hofmann 2014; Nieuwenhuis, Te Grotenhuis, and Pelzer 2012).

Finally, posterior predictive simulation (Gelman and Hill 2006) is a generally useful diagnostic tool, adapted from Bayesian methods, for exploring model fit. Users pick some summary statistic of interest. They then compute the summary statistic for an ensemble of simulations (Section 5.2.8), and see where the observed data falls within the simulated distribution; if the observed data is extreme, we might conclude that the model is a poor representation of reality. For example, using the sleep study fit and choosing the interquartile range of the reaction times as the summary statistic:

\begin{verbatim}
> iqrvec <- sapply(simulate(fm1, 1000), IQR)
> obsval <- IQR(sleepstudy$Reaction)
> post.pred.p <- mean(obsval >= c(obsval, iqrvec))
\end{verbatim}

The (one-tailed) posterior predictive \( p \) value of 0.78 indicates that the model represents the data adequately, at least for this summary statistic. In contrast to the full Bayesian case, the procedure described here does not allow for the uncertainty in the estimated parameters. However, it should be a reasonable approximation when the residual variation is large.
Sequential decomposition and model comparison

Objects of class ‘merMod’ have an anova method which returns F statistics corresponding to the sequential decomposition of the contributions of fixed-effects terms. In order to illustrate this sequential ANOVA decomposition, we fit a model with orthogonal linear and quadratic Days terms,

\[
> \text{fm4} \leftarrow \text{lmer}(\text{Reaction} \sim \text{polyDays[, 1]} + \text{polyDays[, 2]} + \\
(\text{polyDays[, 1]} + \text{polyDays[, 2]} | \text{Subject}), \\
\text{within(sleepstudy, polyDays} <\text{ poly(Days, 2))})
\]

> anova(fm4)

Analysis of Variance Table

<table>
<thead>
<tr>
<th></th>
<th>npar</th>
<th>Sum Sq</th>
<th>Mean Sq</th>
<th>F value</th>
</tr>
</thead>
<tbody>
<tr>
<td>polyDays[, 1]</td>
<td>1</td>
<td>23875</td>
<td>23875</td>
<td>46.08</td>
</tr>
<tr>
<td>polyDays[, 2]</td>
<td>1</td>
<td>340</td>
<td>340</td>
<td>0.66</td>
</tr>
</tbody>
</table>

The relative magnitudes of the two sums of squares indicate that the quadratic term explains much less variation than the linear term. Furthermore, the magnitudes of the two F statistics strongly suggest significance of the linear term, but not the quadratic term. Notice that this is only an informal assessment of significance as there are no p values associated with these F statistics, an issue discussed in more detail in the next subsection (“Computing p-values”, p. 35).

To understand how these quantities are computed, let \( \mathbf{R}_i \) contain the rows of \( \mathbf{R}_X \) (Equation 18) associated with the ith fixed-effects term. Then the sum of squares for term \( i \) is,

\[
SS_i = \hat{\beta}^\top \mathbf{R}_i^\top \mathbf{R}_i \hat{\beta}
\]

If \( DF_i \) is the number of columns in \( \mathbf{R}_i \), then the F statistic for term \( i \) is,

\[
F_i = \frac{SS_i}{\hat{\sigma}^2 DF_i}
\]

For multiple arguments, the anova method returns model comparison statistics.

> anova(fm1, fm2, fm3)

refitting model(s) with ML (instead of REML)

Data: sleepstudy
Models:
fm3: Reaction ~ Days + (1 | Subject)
fm2: Reaction ~ Days + ((1 | Subject) + (0 + Days | Subject))
fm1: Reaction ~ Days + (Days | Subject)

<table>
<thead>
<tr>
<th></th>
<th>npar</th>
<th>AIC</th>
<th>BIC</th>
<th>logLik</th>
<th>deviance</th>
<th>Chisq</th>
<th>Df</th>
<th>Pr(&gt;Chisq)</th>
</tr>
</thead>
<tbody>
<tr>
<td>fm3</td>
<td>4</td>
<td>1802</td>
<td>1815</td>
<td>-897</td>
<td>1794</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>fm2</td>
<td>5</td>
<td>1762</td>
<td>1778</td>
<td>-876</td>
<td>1752</td>
<td>42.08</td>
<td>1</td>
<td>8.8e-11</td>
</tr>
<tr>
<td>fm1</td>
<td>6</td>
<td>1764</td>
<td>1783</td>
<td>-876</td>
<td>1752</td>
<td>0.06</td>
<td>1</td>
<td>0.8</td>
</tr>
</tbody>
</table>
The output shows $\chi^2$ statistics representing the difference in deviance between successive models, as well as $p$ values based on likelihood ratio test comparisons. In this case, the sequential comparison shows that adding a linear effect of time uncorrelated with the intercept leads to an enormous and significant drop in deviance ($\Delta$deviance $\approx 42$, $p \approx 10^{-10}$), while the further addition of correlation between the slope and intercept leads to a trivial and non-significant change in deviance ($\Delta$deviance $\approx 0.06$). For objects of class ‘lmerMod’ the default behavior is to refit the models with ML if fitted with REML = TRUE, which is necessary in order to get sensible answers when comparing models that differ in their fixed effects; this can be controlled via the refit argument.

**Computing $p$ values**

One of the more controversial design decisions of lme4 has been to omit the output of $p$ values associated with sequential ANOVA decompositions of fixed effects. The absence of analytical results for null distributions of parameter estimates in complex situations (e.g., unbalanced or partially crossed designs) is a long-standing problem in mixed-model inference. While the null distributions (and the sampling distributions of non-null estimates) are asymptotically normal, these distributions are not $t$ distributed for finite size samples – nor are the corresponding null distributions of differences in scaled deviances $F$ distributed. Thus approximate methods for computing the approximate degrees of freedom for $t$ distributions, or the denominator degrees of freedom for $F$ statistics (Satterthwaite 1946; Kenward and Roger 1997), are at best ad hoc solutions.

However, computing finite-size-corrected $p$ values is sometimes necessary. Therefore, although the package does not provide them (except via parametric bootstrapping, Section 5.1.3), we have provided a help page to guide users in finding appropriate methods:

```r
> help("pvalues")
```

This help page provides pointers to other packages that provide machinery for calculating $p$ values associated with ‘merMod’ objects. It also suggests framing the inference problem as a likelihood ratio test (achieved by supplying multiple ‘merMod’ objects to the anova method (Section 5.2.4), as well as alternatives to $p$ values such as confidence intervals (Section 5.2.6). Previous versions of lme4 provided the mcmcsamp function, which generated a Markov chain Monte Carlo sample from the posterior distribution of the parameters (assuming flat priors). Due to difficulty in constructing a version of mcmcsamp that was reliable even in cases where the estimated random-effect variances were near zero, mcmcsamp has been withdrawn.

**Computing confidence intervals**

As described above (Section 5.1.1), lme4 provides confidence intervals (using confint) via Wald approximations (for fixed-effect parameters only), likelihood profiling, or parametric bootstrapping (the boot.type argument selects the bootstrap confidence interval type).

As is typical for computationally intensive profile confidence intervals in R, the intervals can be computed either directly from fitted ‘merMod’ objects (in which case profiling is done as an interim step), or from a previously computed likelihood profile (of class ‘thpr’, for “theta profile”). Parametric bootstrapping confidence intervals use a thin wrapper around the bootMer function that passes the results to boot.ci from the boot package (Canty and Ripley 2015; Davison and Hinkley 1997) for confidence interval calculation.
In the running sleep study example, the 95% confidence intervals estimated by all three methods are quite similar. The largest change is a 26% difference in confidence interval widths between profile and parametric bootstrap methods for the correlation between the intercept and slope random effects \((-0.54,0.98)\) vs. \((-0.48,0.68)\).

**General profile zeta and related plots**

**lme4** provides several functions (built on lattice graphics, Sarkar 2008) for plotting the profile zeta functions (Section 5.1.2) and other related quantities.

- **The profile zeta plot** (Figure 2; `xyplot`) is simply a plot of the profile zeta function for each model parameter; linearity of this plot for a given parameter implies that the likelihood profile is quadratic (and thus that Wald approximations would be reasonably accurate).

- **The profile density plot** (Figure 3; `densityplot`) displays an approximation of the probability density function of the sampling distribution for each parameter. These densities are derived by setting the cumulative distribution function (c.d.f) to be \(\Phi(\zeta)\) where \(\Phi\) is the c.d.f of the standard normal distribution. This is not quite the same as evaluating the distribution of the estimator of the parameter, which for mixed models can be very difficult, but it gives a reasonable approximation. If the profile zeta plot is linear, then the profile density plot will be Gaussian.

- **The profile pairs plot** (Figure 4; `splom`) gives an approximation of the two-dimensional profiles of pairs of parameters, interpolated from the univariate profiles as described in Bates and Watts (1988, Chapter 6). The profile pairs plot shows two-dimensional 50%, 80%, 90%, 95% and 99% marginal confidence regions based on the likelihood ratio, as well as the profile traces, which indicate the conditional estimates of each parameter for fixed values of the other parameters. While panels above the diagonal show profiles with respect to the original parameters (with random-effects parameters on the standard deviation/correlation scale, as for all profile plots), the panels below the diagonal show plots on the \((\zeta_i, \zeta_j)\) scale. The below-diagonal panels allow us to see distortions from an elliptical shape due to nonlinearity of the traces, separately from the one-dimensional distortions caused by a poor choice of scale for the parameters. The \(\zeta\) scales provide, in some sense, the best possible set of single-parameter transformations for assessing the contours. On the \(\zeta\) scales the extent of a contour on the horizontal axis is exactly the same as the extent on the vertical axis and both are centered about zero.

For users who want to build their own graphical displays of the profile, there is a method for `as.data.frame` that converts profile (``thpr`’) objects to a more convenient format.

**Computing fitted and predicted values; simulating**

Because mixed models involve random coefficients, one must always clarify whether predictions should be based on the marginal distribution of the response variable or on the distribution that is conditional on the modes of the random effects (Equation 12). The **fitted** method retrieves fitted values that are conditional on all of the modes of the random effects; the **predict** method returns the same values by default, but also allows for predictions to be
made conditional on different sets of random effects. For example, if the `re.form` argument is set to `NULL` (the default), then the predictions are conditional on all random effects in the model; if `re.form` is `-0` or `NA`, then the predictions are made at the population level (all random-effect values set to zero). In models with multiple random effects, the user can give `re.form` as a formula that specifies which random effects are conditioned on. `lme4` also provides a `simulate` method, which allows similar flexibility in conditioning on random effects; in addition it allows the user to choose (via the `use.u` argument) between conditioning on the fitted conditional modes or choosing a new set of simulated conditional modes (zero-mean Normal deviates chosen according to the estimated random-effects variance-covariance matrices). Finally, the `simulate` method has a method for `formula` objects, which allows for `de novo` simulation in the absence of a fitted model. In this case, the user must specify the random-effects ($\theta$), fixed-effects ($\beta$), and residual standard deviation ($\sigma$) parameters via the `newparams` argument. The standard simulation method (based on a `merMod` object) is the basis of parametric bootstrapping (Section 5.1.3) and posterior predictive simulation (Section 5.2.3); `de novo` simulation based on a formula provides a flexible framework for power analysis.

6. Conclusion

Mixed modeling is an extremely useful but computationally intensive technique. Computational limitations are especially important because mixed models are commonly applied to moderately large data sets ($10^4$–$10^6$ observations). By developing an efficient, general, and (now) well-documented platform for fitted mixed models in R, we hope to provide both a practical tool for end users interested in analyzing data and a reusable, modular framework...
for downstream developers interested in extending the class of models that can be easily and efficiently analyzed in \textit{R}.

We have learned much about linear mixed models in the process of developing \textit{lme4}, both from our own attempts to develop robust and reliable procedures and from the broad community of \textit{lme4} users; we have attempted to describe many of these lessons here. In moving forward, our main priorities are (1) to maintain the reference implementation of \textit{lme4} on the Comprehensive \textit{R} Archive Network (CRAN), developing relatively few new features; (2) to improve the flexibility, efficiency and scalability of mixed-model analysis across multiple compatible implementations, including both the \texttt{MixedModels} package for \textit{Julia} and the experimental \texttt{flexLambda} branch of \textit{lme4}.

\section*{Acknowledgments}

Rune Haubo Christensen, Henrik Singmann, Fabian Scheipl, Vincent Dorie, and Bin Dai contributed ideas and code to the current version of \textit{lme4}; the large community of \textit{lme4} users has exercised the software, discovered bugs, and made many useful suggestions. Søren Hojsgaard participated in useful discussions and Xi Xia and Christian Zingg exercised the code and reported problems. We would like to thank the Banff International Research Station for hosting a working group on \textit{lme4}, and an NSERC Discovery grant and NSERC postdoctoral fellowship for funding.
Figure 4: Profile pairs plot: `splom(prof.obj)`
References


A. Modularization examples

The modularized functions in \texttt{lme4} allow for finer control of the various steps in fitting mixed models (Table 1). The main advantage of these functions is the ability to extend the class of models that \texttt{lmer} is capable of fitting. In general there are three steps to this process.

- Construct a mixed-model formula that is similar to the intended model.
- Using the modular functions, hook into and modify the fitting process so that the desired model is estimated.
- Use the estimated parameter values to make inferences.

Note that the many tools of the output module become unavailable for sufficiently extensive departures from the class of models fitted by \texttt{lmer}. Therefore, this last inference step may be more involved in the modular approach than with inference from models fitted by \texttt{lmer}. Here we provide two examples that illustrate this process.

A.1. Zero slope-slope covariance models

Consider the following mixed-model formula,

\begin{verbatim}
f \texttt{orm <- respVar ~ 1 + (explVar1 + explVar2 | groupFac)}
\end{verbatim}

in which we have a fixed intercept and two random-effects terms, each with a random slope and intercept. The \texttt{lmer} function would fit this formula to data by estimating an unstructured \(3 \times 3\) covariance matrix giving the variances and covariances among the intercept and the two slopes. It is not possible to use the current version of \texttt{lmer} to fit a model with the following two properties: (1) the slopes are uncorrelated and (2) the intercept is correlated with both of the slopes. We illustrate the use of the modular functions in \texttt{lme4} to fit such a model.

\textit{Simulate from a zero slope-slope covariance model}

We simulate data from such a model using a balanced design with a single grouping factor. We use \texttt{lme4}'s utilities for constructing a template from the model formula, with which to conduct these simulations. We simulate a balanced design with 500 levels to the grouping factor and 20 observations per group,

\begin{verbatim}
> set.seed(1)
> dat <- mkDataTemplate(form, 
> nGrps = 500, 
> nPerGrp = 20, 
> rfunc = rnorm)
> head(dat)
\end{verbatim}

\begin{verbatim}
   groupFac   respVar  explVar1  explVar2
1       1   0.62645 -0.80433   0.23535
2       1   0.18364 -1.05653   0.24483
3       1  -0.83563 -1.03540  -0.64219
\end{verbatim}
The initial parameters for this model may be obtained using `mkParsTemplate`.

```r
> (pars <- mkParsTemplate(form, dat))

beta
(Intercept) 0

theta
groupFac.(Intercept) groupFac.explVar1.(Intercept)
1 0
groupFac.explVar2.(Intercept) groupFac.explVar1
0 1
groupFac.explVar2.explVar1 groupFac.explVar2
0 1

sigma
[1] 1
```

We wish to simulate random effects from the following covariance matrix,

```r
> vc <- matrix(c(1.0, 0.5, 0.5,
                  0.5, 1.0, 0.0,
                  0.5, 0.0, 1.0), 3, 3)
```

Here the intercept is correlated with each of the two slopes, but the slopes are uncorrelated. The `theta` vector that corresponds to this covariance matrix can be computed using `lme4`'s facilities for converting between various representations of (co-)variance structures (see `?vcconv`).

```r
> pars$theta[] <- Vv_to_Cv(mlist2vec(vc))
```

With these new parameters installed, we simulate a response variable.

```r
> dat$respVar <- simulate(form,
                           newdata = dat,
                           newparams = pars,
                           family = "gaussian")[[1]]
```

Indeed this data set appears to contain intercept-slope correlations, but not slope-slope correlations, as we see using the `lmList` function.

```r
> formLm <- form
> formLm[[3]] <- findbars(form[[1]])
> print(formLm)
```
respVar ~ explVar1 + explVar2 | groupFac

> cor(t(sapply(lmList(formLm, dat), coef)))

<p>| | | | | | |</p>
<table>
<thead>
<tr>
<th></th>
<th></th>
<th></th>
<th></th>
<th></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>(Intercept)</td>
<td>1.00</td>
<td>0.53</td>
<td>0.43</td>
<td></td>
<td></td>
</tr>
<tr>
<td>explVar1</td>
<td>0.53</td>
<td>1.00</td>
<td>0.06</td>
<td></td>
<td></td>
</tr>
<tr>
<td>explVar2</td>
<td>0.43</td>
<td>0.06</td>
<td>1.00</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

This `lmList` function computes ordinary linear models for each level of the grouping factor. Next we show how to fit a proper mixed model to these data.

**Fitting a zero slope-slope covariance model**

The main challenge with fitting a zero slope-slope model is the need to figure out which Cholesky decompositions correspond to variance-covariance matrices with structural zeros between the slopes. There are two general approaches to solving this problem. The first is to reorder the rows and columns of the covariance matrix so that the intercept column comes last. In this case, setting the second component of the covariance parameter vector, \( \theta \), to zero will achieve the desired covariance structure. However, we consider a second approach, which retains the ordering produced by the `lFormula` function: (1) intercept, (2) first slope, and (3) second slope. The covariance structure for the random effects in this model is given by the following covariance matrix and its associated Cholesky decomposition.

\[
\begin{pmatrix}
\sigma_1^2 & \sigma_{12} & \sigma_{13} \\
\sigma_{21} & \sigma_2^2 & 0 \\
\sigma_{31} & 0 & \sigma_3^2
\end{pmatrix} =
\begin{pmatrix}
\theta_1 & 0 & 0 \\
\theta_2 & \theta_4 & 0 \\
\theta_3 & \theta_5 & \theta_6
\end{pmatrix}
\begin{pmatrix}
\theta_1 & \theta_2 & \theta_3 \\
0 & \theta_4 & \theta_5 \\
0 & 0 & \theta_6
\end{pmatrix}
\]

(70)

where the \( \sigma \) parameters give the standard deviations and covariances, and the \( \theta \) parameters are the elements of the Cholesky decomposition. The key feature of this covariance matrix is the structural zero indicating no covariance between the two slopes. Therefore, in order to ensure the structural zero, we must have,

\[
\theta_5 = -\frac{\theta_2 \theta_3}{\theta_4}.
\]

(71)

To represent this restriction in R we introduce a length five vector, `phi`, which we convert into an appropriate `theta` vector using the following function.

> phiToTheta <- function(phi) {
    theta5 <- -(phi[2]*phi[3])/phi[4]
    c(phi[1:4], theta5, phi[5])
}

We may therefore optimize the structured model by optimizing over `phi` instead of `theta`. We begin to construct an appropriate objective function for such an optimization by taking the first two modular steps, resulting in a deviance function of `theta`.

> lf <- lFormula(formula = form, data = dat, REML = TRUE)
> devf <- do.call(mkLmerDevfun, lf)
This deviance function is a function of the unconstrained \texttt{theta} vector, but we require a deviance function of the constrained \texttt{phi} vector. We therefore construct a wrapper around the deviance function as follows.

\begin{verbatim}
> devfWrap <- function(phi) devf(phiToTheta(phi))
\end{verbatim}

Finally, we optimize the deviance function,

\begin{verbatim}
> opt <- bobyqa(par = lf$reTrms$theta[-5],
             fn = devfWrap,
             lower = lf$reTrms$lower[-5])
\end{verbatim}

To extract and label the estimated covariance matrix of the random effects, we may use the conversion functions (?vcconv) and the column names of the random-effects model matrix (\texttt{cnms}). We construct this estimated matrix and compare it with the true matrix used to simulate the data.

\begin{verbatim}
> vcEst <- vec2mlist(Cv_to_Vv(phiToTheta(opt$par)))[[1]]
> dimnames(vcEst) <- rep(lf$reTrms$cnms, 2)
> round(vcEst, 2)
\end{verbatim}

\begin{verbatim}
groupFac
  groupFac (Intercept) explVar1 explVar2
  (Intercept)  1.00  0.57  0.42
  explVar1    0.57  1.11  0.00
  explVar2    0.42  0.00  0.99
\end{verbatim}

\begin{verbatim}
> vc
\end{verbatim}

\begin{verbatim}
[,1] [,2] [,3]
[1,] 1.0  0.5  0.5
[2,] 0.5  1.0  0.0
[3,] 0.5  0.0  1.0
\end{verbatim}

We see that the estimated covariance matrix provides a good estimate of the true matrix.

\section*{A.2. Additive models}

It is possible to interpret additive models as a particular class of mixed models (Wood and Scheipl 2014). The main benefit of this approach is that it bypasses the need to select a smoothness parameter using cross-validation or generalized cross-validation. However, it is inconvenient to specify additive models using the \texttt{lme4} formula interface. The \texttt{gamm4} package wraps the modularized functions of \texttt{lme4} within a more convenient interface (Wood and Scheipl 2014). In particular, the strategy involves the following steps,

1. Convert a \texttt{gamm4} formula into an \texttt{lme4} formula that approximates the intended model.
2. Parse this formula using \texttt{lFormula}.
3. Modify the resulting transposed random-effects model matrix, $Z^\top$, so that the intended additive model results.

4. Fit the resulting model using the remaining modularized functions (Table 1).

Here we illustrate this general strategy using a simple simulated data set,

```r
> library("gamm4")
> library("splines")
> set.seed(1)
> n <- 100
> pSimulation <- 3
> pStatistical <- 8
> x <- rnorm(n)
> Bsimulation <- ns(x, pSimulation)
> Bstatistical <- ns(x, pStatistical)
> beta <- rnorm(pSimulation)
> y <- as.numeric(Bsimulation %*% beta + rnorm(n, sd = 0.3))
```

where `pSimulation` is the number of columns in the simulation model matrix and `pStatistical` is the number of columns in the statistical model matrix.

We plot the resulting data along with the predictions from the generating model,

```r
> par(mar = c(4, 4, 1, 1), las = 1, bty = "l")
> plot(x, y, las = 1)
> lines(x[order(x)], (Bsimulation %*% beta)[order(x)])
```

We now set up an approximate `lme4` model formula, and parse it using `lFormula`,

\[ y \sim \beta_0 + \beta_1 x + \beta_2 x^2 + \epsilon \]

\[ \epsilon \sim N(0, \sigma^2) \]
We now insert a spline basis into the `parsedFormula` object,

```r
> parsedFormula$reTrms <- within(parsedFormula$reTrms, {
  Bt <- t(as.matrix(Bstatistical))[]
  cnms$pseudoGroups <- "spline"
  Zt <- as(Bt, class(Zt))
})
```

Finally we continue with the remaining modular steps,

```r
> devianceFunction <- do.call(mkLmerDevfun, parsedFormula)
> optimizerOutput <- optimizeLmer(devianceFunction)
> mSpline <- mkMerMod( rho = environment(devianceFunction),
  opt = optimizerOutput,
  reTrms = parsedFormula$reTrms,
  fr = parsedFormula$fr)
> mSpline

Linear mixed model fit by REML ['lmerMod']
REML criterion at convergence: 60.734
Random effects:
  Groups   Name   Std.Dev.
  pseudoGroups  spline 0.293
  Residual     0.300
Number of obs: 100, groups: pseudoGroups, 8
Fixed Effects:
  (Intercept)    x
  -0.0371   -0.1714
```

Computing the fitted values of this additive model requires some custom code, and illustrates the general principle that methods for `merMod` objects constructed from modular fits should only be used if the user knows what she is doing,

```r
> xNew <- seq(min(x), max(x), length = 100)
> newBstatistical <- predict(Bstatistical, xNew)
> yHat <- cbind(1, xNew) %*% getME(mSpline, "fixef") +
  newBstatistical %*% getME(mSpline, "u")
> par(mar = c(4, 4, 1, 1), las = 1, bty = "l")
> plot(x, y)
> lines(xNew, yHat)
> lines(x[order(x)], (Bsimulation %*% beta)[order(x)], lty = 2)
> legend("topright", bty = "n", c("fitted", "generating"), lty = 1:2, col = 1)
```
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