JOURNAL DE LA SOCIÉTÉ FRANÇAISE DE STATISTIQUE

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Journal de la société française de statistique, tome 143, nº 1-2 (2002), p. 79-101

http://www.numdam.org/item?id=JSFS_2002__143_1-2_79_0

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MODEL BUILDING USING COVARIATES IN NONLINEAR MIXED-EFFECTS MODELS

José C. PINHEIRO *

ABSTRACT

Nonlinear mixed-effects (NLME) models are useful in describing nonlinear relationships between a response variable and parameters and covariates in data that are grouped according to a classification factor. Examples of such grouped data include longitudinal and repeated measures data, which frequently arise in many areas of application, such as pharmacokinetics, biostatistics, and economics. By associating common random effects to observations sharing the same level of a classification factor, NLME models flexibly represent the covariance structure induced by the grouping of the data. The random effects in an NLME model account for differences in the parameter values among groups. In many applications, additional covariates are collected along with the response and model covariates and can be used to explain, at least partially, the variation among groups. Including covariates in an NLME model to explain inter-group variation generally leads to simplifications in the random effects model and to a better understanding of the model producing the response. This paper describes a model-building strategy for identifying and including covariates in an NLME model, using the capabilities available in the NLME library for S-PLUS and R. The use of the proposed methodology and the capabilities in the NLME software are illustrated with real-life examples from ecophysiology and pharmacokinetics.

RÉSUMÉ

Les modèles non-linéaires à effets mixtes (NLME) sont très utiles pour décrire des relations non-linéaires entre une variable réponse et des paramètres ou covariables lorsque les données observées sont groupées selon un niveau de facteur. De telles données englobent entre autres les données longitudinales, les mesures répétées et concernent beaucoup de domaines applicatifs telles la phamaco-cinétique, la biostatistique, l'économie. En associant aux observations des effets aléatoires communs à chaque niveau au facteur de groupement, la structure de covariance associée est aisément représentée par les modèles NLME : les effets aléatoires caractérisent les différences des valeurs des paramètres entre groupes. Dans beaucoup d'applications, des covariables additionnelles sont mesurées parallèlement à la variable réponse et elles peuvent être utilisées pour expliquer des variations même partielles entre les groupes. Inclure ces covariables dans un modèle NLME pour expliquer des variations inter-groupes amène généralement à une simplification de la modélisation des effets aléatoires et à une meilleure compréhension du modèle explicatif. Ce papier décrit une stratégie de modélisation pour l'identification et la prise en compte de covariables utilisant les capacités de la librairie NLME disponible sous S-PLUS et sous R. La mise en œuvre de sont cette méthodologie est illustrée sur des exemples concrets en écophysiologie et en pharmo-cinétique.

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1. Introduction

Nonlinear mixed-effects (NLME) models are useful in describing nonlinear relationships between a response variable and parameters and covariates in data that are grouped according to a classification factor. Examples of such grouped data include longitudinal and repeated measures data, which frequently arise in many areas of application, such as pharmacokinetics, biostatistics, and economics. NLME models assume that the form of the intragroup model relating the response variable to covariates is common to all groups, but some of the parameters that define the model are allowed to vary with group, through the use of random effects. By associating common random effects to observations in the same group, NLME models flexibly represent the covariance structure induced by the grouping of the data.

The random effects in an NLME model account for individual deviations in the parameters among groups. In many applications, these deviations can be at least partially explained by differences in covariate values among groups. Including covariates in an NLME model to explain inter-group variation often leads to simplifications in the random effects model and generally allows a better understanding of the mechanism producing the response.

During the process of adding covariates to an NLME model, several model building questions need to be addressed, such as :

- Among the candidate covariates, which are potentially useful in explaining the random effects variation?
- Which random effects have their variation best explained by covariates?
- How should the potentially useful covariates be tested for inclusion in the model?
- Should random effects be included in, or eliminated from, the model, after covariates have been included?

This paper describes a model-building strategy for addressing these questions in the context of NLME models, using the capabilities available in the NLME library for S-PLUS and R (Pinheiro and Bates, 2000). The use of the proposed methodology and the capabilities in the NLME software are illustrated with real-life examples from ecophysiology and pharmacokinetics.

The rest of the paper is organized as follows. The nonlinear mixed-effects model is described in Section 2. Section 3 introduces examples from ecophysiology (CO₂ uptake) and pharmacokinetics (clinical study of Quinidine) which are later used to illustrate the model building methodology, described in Section 4. Conclusions and suggestions for further research are presented in Section 5.

2. A nonlinear mixed-effects model

Nonlinear mixed-effects (NLME) models are mixed-effects models in which the response function is nonlinear in at least some of the underlying parameters. Several different nonlinear mixed-effects models have been proposed in the

literature (Sheiner and Beal, 1980; Mallet et al., 1988; Lindstrom and Bates, 1990; Vonesh and Carter, 1992; Davidian and Gallant, 1992; Wakefield et al., 1994). We adopt here the NLME model proposed by Lindstrom and Bates (1990), which can be viewed as a hierarchical model that generalizes both the linear mixed-effects model of Laird and Ware (1982) and the usual nonlinear regression model for independent data (Bates and Watts, 1988). In the first stage, the jth observation on the ith group is described as

$$y_{ij} = f(\phi_{ij}, \mathbf{x}_{ij}) + \epsilon_{ij}, \ i = 1, \dots, M, \ j = 1, \dots, n_i$$
 (1)

where f is a nonlinear function of a group-specific vector of parameters ϕ_{ij} and the vector of covariates x_{ij} , the ϵ_{ij} are normally distributed, independent within-group error terms, M is the total number of groups, and n_i is the number of observations in the ith group. In the second stage the group-specific parameters are modeled as

$$\phi_{ij} = A_{ij}\beta + B_{ij}b_i \tag{2}$$

where β represents the fixed effects; b_i the random effects (varying with i but not with j), which are assumed to be independently distributed as $\mathcal{N}(\mathbf{0}, \Psi)$. A_{ij} and B_{ij} are design matrices for the fixed and random effects respectively, which may depend on the values of some covariates at the jth observation. It is further assumed that the b_i are independent of the ϵ_{ij} .

Inclusion of covariates in the NLME model is done primarily through the A_{ij} matrices. As described in Section 4, candidate covariates for inclusion in the model are screened using plots of the estimated random effects against available covariates. The most promising covariates are incorporated in the model by adding corresponding columns to the A_{ij} , with resulting estimated fixed effects being tested for significance. This model building strategy is described and illustrated in detail in Section 4.

Different methods have been proposed to estimate the parameters in the NLME model (Ramos and Pantula, 1995; Davidian and Giltinan, 1995; Vonesh and Chinchilli, 1997); we concentrate here on methods based on the likelihood function. Because the random effects b_i are unobserved quantities, maximum likelihood estimation in NLME models is based on the marginal density of the responses y, which is calculated as

$$\ell(\boldsymbol{\beta}, \boldsymbol{\Psi}, \sigma | \boldsymbol{y}) = \int p(\boldsymbol{y} | \boldsymbol{b}, \boldsymbol{\beta}, \sigma) p(\boldsymbol{b} | \boldsymbol{\Psi}) d\boldsymbol{b}$$
 (3)

Because the model function f in (1) can be nonlinear in the random effects, the integral in (3) generally does not have a closed-form expression. To make the numerical optimization of the likelihood function a tractable problem, different approximations to (3) have been proposed. Some of these methods consist of taking a first-order Taylor expansion of the model function f around the expected value of the random effects (i.e., 0) (Sheiner and Beal, 1980; Vonesh and Carter, 1992), or around the conditional modes of the random effects (\hat{b}_i) (Lindstrom and Bates, 1990). We adopt here the approximation suggested

by Lindstrom and Bates (1990), which is implemented via an alternating algorithm comprising a linear mixed-effects (LME) step and a penalized nonlinear least squares (PNLS) step. This is the algorithm used in the nlme function of the **NLME** library. Inferences on the model parameters, including hypothesis testing, are based on asymptotic results for the linear mixed-effects log-likelihood used in the LME step of the alternating algorithm (Pinheiro and Bates, 2000).

3. Examples

In this section we introduce the two examples that will be used to illustrate the model building methodology for incorporating covariates in an NLME model described in Section 4. We also describe how to fit the associated NLME models using the tools in the NLME library.

3.1. Carbon dioxide uptake

Data from a study of the cold tolerance of a C_4 grass species, *Echinochloa crusgalli* is reported in Potvin *et al.* (1990). A total of 12 four-week-old plants, 6 from Québec and 6 from Mississippi, were divided into two groups – control plants that were kept at 26°C and chilled plants that were subject to 14 h of chilling at 7°C. After 10 h of recovery at 20°C, carbon dioxide (CO_2) uptake rates (in μ mol/m²s) were measured for each plant at seven increasing concentrations of ambient CO_2 (μ L/L). The objective of the experiment was to evaluate the effect of plant type and chilling treatment on the CO_2 uptake. The CO_2 data, displayed in Figure 1, are available in the NLME library as the groupedData object CO_2 (Pinheiro and Bates, 2000).

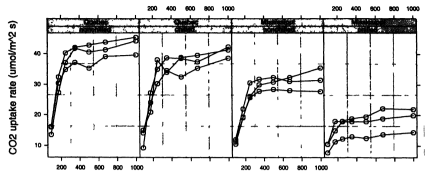
> CO2

Grouped Data : uptake ∼ conc | Plant

	Plant	Туре	Treatment	conc	uptake
1	Qn1	Quebec	nonchilled	95	16.0
2	Qn1	Quebec	nonchilled	175	30.4
	•				
83	Mc3	Mississippi	chilled	675	18.9
84	Mc3	Mississippi	chilled	1000	19.9

It is clear from Figure 1 that the CO_2 uptake rates of Québec plants are greater than those of Mississippi plants and that chilling the plants reduces their CO_2 uptake rates, with this decrease being more pronounced in Mississippi plants than in Québec plants.

An asymptotic regression model with an offset is used in Potvin *et al.* (1990) to represent the expected CO_2 uptake rate U(c) as a function of the ambient CO_2 concentration c



Ambient carbon dioxide concentration (uL/L)

FIG 1 - CO₂ uptake versus ambient CO₂ by chilling treatment and plant type for *Echinochloa crus-qalli* plants

$$U(c) = \phi_1 \left\{ 1 - \exp\left[-\exp\left(\phi_2\right)(c - \phi_3)\right] \right\},\tag{4}$$

where ϕ_1 is the asymptotic CO₂ uptake rate, ϕ_2 is the logarithm of the rate constant, and ϕ_3 is the maximum ambient concentration of CO₂ at which there is no uptake. The logarithm of the rate constant is used to enforce the positivity of the estimated rate constant, while keeping the optimization problem unconstrained.

We initially consider an NLME version of the CO_2 uptake model (4) with all parameters as mixed effects and no treatment covariates. The corresponding model for the CO_2 uptake u_{ij} of plant i at ambient CO_2 concentration c_{ij} is

$$u_{ij} = \phi_{1i} \left\{ 1 - \exp\left[- \exp\left(\phi_{2i}\right) \left(c_{ij} - \phi_{3i} \right) \right] \right\} + \epsilon_{ij},$$

$$\phi_{i} = \begin{bmatrix} \phi_{1i} \\ \phi_{2i} \\ \phi_{3i} \end{bmatrix} = \begin{bmatrix} \beta_{1} \\ \beta_{2} \\ \beta_{3} \end{bmatrix} + \begin{bmatrix} b_{1i} \\ b_{2i} \\ b_{3i} \end{bmatrix} = \boldsymbol{\beta} + \boldsymbol{b}_{i},$$

$$\boldsymbol{b}_{i} \sim \mathcal{N} \left(\mathbf{0}, \boldsymbol{\Psi} \right), \quad \epsilon_{ij} \sim \mathcal{N} \left(\mathbf{0}, \sigma^{2} \right),$$

$$(5)$$

where ϕ_{1i} , ϕ_{2i} , and ϕ_{3i} have the same interpretation as in model (4), but are now allowed to vary with plant. The fixed effects, β , represent the population average of the individual parameters, ϕ_{i} , and the random effects, b_{i} , represent the deviations of the ϕ_{i} from their population average. In the notation of the general NLME model (2), $A_{ij} = B_{ij} = I$.

The **NLME** library includes an SSasympOff function with a self-starting implementation of model (4), which is used to automatically generate starting estimates for the parameters in the model. We use it to fit the NLME model (5).

```
> fm1CO2 <- nlme(uptake ~ SSasympOff(conc, Asym, 1rc, c0),</pre>
    data = CO2, fixed = Asym + lrc + c0 \sim 1)
> fm1CO2
  Fixed: list(Asym \sim 1, lrc \sim 1, c0 \sim 1)
             1rc
 32.474
        -4.6362
                  43.543
Random effects :
 Formula : list(Asym \sim 1, lrc \sim 1, c0 \sim 1)
 Level: Plant
 Structure : General positive-definite
             StdDev
                        Corr
     Asym
            9.50999
                      Asym
                                  1rc
      1rc
            0.12828
                     -0.160
       c0
           10.40519
                       0.999
                              -0.139
 Residual
            1.76641
```

with $\phi_1 = Asym$, $\phi_2 = 1rc$, and $\phi_3 = c0$.

The very high correlation between Asym and c0 gives indication that the fit corresponds to a numerically unstable solution. In general, from practical experience, this occurs when the random effects model is over-parameterized and the optimization algorithm attempts to converge to a lower dimension $\widehat{\Psi}$ but, because of the parameterization used in the NLME software, ends up converging to a boundary solution. In this particular case, the estimated $\widehat{\Psi}$ suggests that a single random effect could be used to represent both the Asym and the c0 random effects. The scatter plot matrix of the estimated random effects (not shown) confirms that Asym and c0 are in almost perfect linear alignment. A possible model for the random effects in this case would express the c0 random effect as a multiple δ of the Asym random effect. This model, however, leads to numerical difficulties in the estimation of δ using the alternating algorithm implemented in the NLME software, resulting in singular gradient matrices.

An alternative approach is to consider simpler models with one of the highly correlated random effects removed. Dropping the Asym random effect results in a highly significant decrease in the log-likelihood of the fitted model (p-value < 0.0001), while dropping the c0 random effect gives an equivalent fit of the data (p-value of 0.41 for the likelihood ratio test comparing the two nested models). Therefore, we chose the model with Asym and 1rc random effects only, which can be fit in NLME with

This is the model that will be used as a starting point for covariate model building in Section 4.

Formally, the hypotheses being tested when deciding whether a random effect could be dropped from model (5) are

$$H_0: \mathbf{\Psi} = \begin{pmatrix} \mathbf{\Psi}_0 & \mathbf{0}' \\ \mathbf{0} & 0 \end{pmatrix}, \ \mathbf{\Psi}_0 \in M_2^+ \quad vs. \quad H_1: \mathbf{\Psi} \in M_3^+,$$
 (6)

where M_q^+ represents the cone of symmetric positive-definite matrices of order q. This corresponds to a non-classical likelihood ratio test scenario, as the null hypothesis H_0 lies in the boundary of the parameter space. As discussed in Stram and Lee (1994) (see also the correction to the original paper in Stram and Lee (1995)), using the results of Self and Liang (1987), classical likelihood ratio tests under this scenario tend to be conservative. That is, the p-value calculated from the χ_1^2 distribution is greater than it should be, under the correct asymptotic distribution. As described in Stram and Lee (1994), the asymptotic distribution of the likelihood ratio test under the boundary conditions (6) is given by a mixture of χ^2 distributions with degrees-of-freedom ≤ 1 . The conclusions regarding the choice of which random effects to keep in the NLME model for the CO₂ data are unchanged when the corrected likelihood ratio test distribution is used (the corresponding p-value for eliminating the c0 random effect is 0.20).

3.2. Clinical study of quinidine

Routine clinical data on patients receiving the drug quinidine as a treatment for cardiac arrythmia (atrial fibrillation of ventricular arrythmias) were reported in Verme et al. (1992). All patients were receiving oral quinidine doses. At irregular intervals blood samples were drawn and serum concentrations of quinidine were determined. These data, shown in Figure 2, are available as the groupedData object Quinidine in the NLME library.

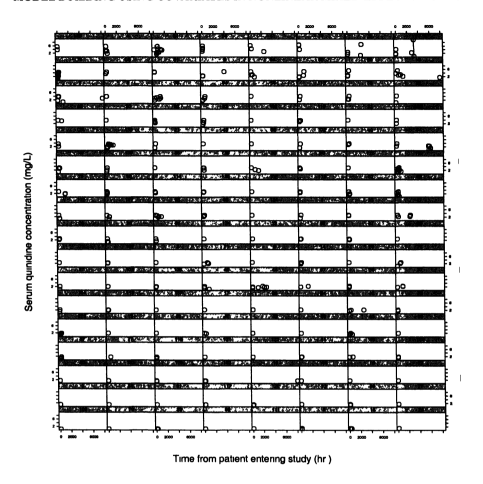


Fig 2 – Serum concentrations of quinidine in 136 hospitalized patients under varying dosage regimens versus time since entering the study.

A total of 361 quinidine concentration measurements were made on 136 hospitalized patients under varying dosage regimens. The times since hospitalization at which the quinidine concentrations were measured varied between 0.13 and 8095.5 hours. Most patients have only a few concentration measurements – 34% have only one and 80% have three or fewer. Only 5% of the patients have seven or more observations. Additional demographic and physiological data were collected for each subject. The additional available covariates are described in Table 1. Some of these covariates, such as age, body weight, and creatinine clearance, were "time-varying." That is, their value for a particular patient could change during the course of the study. Others, such as race, remained constant. One of the main objectives of the study was to investigate relationships between the individual pharmacokinetic parameters and the covariates. Statistical analyses of these data using different modeling approaches

are given in Davidian and Gallant (1992), Davidian and Giltinan (1995) and Wakefield (1996).

Table 1. - Demographic and physiological covariates in the quinidine data.

Age (yr)	42–92
Glycoprotein concentration (mg/100 dL)	0.39-3.16
Body weight (kg)	41-119
Congestive heart failure	no/mild, moderate, severe
Creatinine clearance (ml/min)	< 50, ≥ 50
Ethanol abuse	none, current, former
Height (in.)	60-79
Race	Caucasian, Latin, Black
Smoking status	no, ves

The model that has been suggested for the quinidine data is the one-compartment open model with first-order absorption. This model can be defined recursively as follows. Suppose that, at time t, the patient receives a dose d_t and prior to that time the last dose was given at time t'. The expected concentration in the serum compartment, C_t , and in the absorption compartment, C_{a_t} , are given by

$$C_{t} = C_{t'} \exp\left[-K(t - t')\right] + \frac{Ca_{t'}k_{a}}{k_{a} - K} \left\{\exp\left[-K(t - t')\right] - \exp\left[-k_{a}(t - t')\right]\right\}$$

$$Ca_{t} = Ca_{t'} \exp\left[-k_{a}(t - t')\right] + \frac{d_{t}}{V}$$
(7)

where V is the apparent volume of distribution, k_a is the absorption rate constant, and K is the elimination rate constant.

When a patient receives the same dose d at regular time intervals Δ , model (7) converges to the steady state model

$$C_{t} = \frac{d k_{a}}{V (k_{a} - K)} \left[\frac{1}{1 - \exp(-K\Delta)} - \frac{1}{1 - \exp(-k_{a}\Delta)} \right]$$

$$Ca_{t} = \frac{d}{V \left[1 - \exp(-k_{a}\Delta) \right]}$$
(8)

Patients considered to be in steady state conditions have concentrations modeled as above.

Finally, for a between-dosages time t, the model for the expected concentration C_t , given that the last dose was received at time t', is identical to (7).

Using the fact that the elimination rate constant K is equal to the ratio between the clearance (Cl) and the volume in distribution (V), we can reparametrize models (7) and (8) in terms of V, k_a , and Cl. To ensure that the estimates of V, k_a , and Cl are positive, we can rewrite models (7) and (8) in terms of $lV = \log(V)$, $lk_a = \log(k_a)$ and $lCl = \log(Cl)$.

The initial conditions for the recursive model (7) and (8) are $C_0 = 0$ and $Ca_0 = d_0/V$, with d_0 denoting the first dose received by the patient. It has been assumed throughout the model's definition that the bioavailability of the drug, *i.e.*, the percentage of the administered dose that reaches the measurement compartment, is equal to one.

The function quinModel in the **NLME** library implements the recursive models (7) and (8), parameterized in terms of lV, lKa and lCl. This is not a self-starting model, so initial values for the fixed effects need to be provided when calling nlme. We used values reported in the literature as starting estimates for the fixed effects.

Preliminary analyses of the data, without using any covariates to explain intersubject variation, indicate that only lCl and lV need random effects to account for their variability in the patient population, and that the corresponding random effects can be assumed to be independent. The corresponding models for the fixed and random effects are

$$lCl_{i} = \beta_{1} + b_{1i}, \quad lV_{i} = \beta_{2} + b_{2i}, \quad lKa_{i} = \beta_{3},$$

$$\boldsymbol{b}_{i} = \begin{bmatrix} b_{1i} \\ b_{2i} \end{bmatrix} \sim \mathcal{N} \left(\boldsymbol{0}, \begin{bmatrix} \psi_{1} & 0 \\ 0 & \psi_{2} \end{bmatrix} \right), \tag{9}$$

which is fitted in S-PLUS or R with

```
> fm1Ouin <-
  nlme(conc ~ quinModel(Subject, time, conc, dose, interval,
                           1V, 1Ka, 1C1),
       data = Quinidine, fixed = 1V + 1Ka + 1C1 \sim 1,
       random = pdDiag(lV + lCl \sim 1), groups = \sim Subject,
       start = list(fixed = c(5, -0.3, 2)),
       na.action = na.include, naPattern = \sim !is.na(conc))
> fm1Quin
  Fixed : 1V + 1Ka + 1C1 \sim 1
     1V
              1Ka
                      1C1
5.3796 -0.20535 2.4687
Random effects :
 Formula : list(lV \sim 1, lCl \sim 1)
 Level: Subject
 Structure : Diagonal
                1V
                        1Cl Residual
StdDev: 0.31173 0.32276
                              0.73871 ...
```

This will be the starting model used for covariate model building of the quinidine data in the next section. Note that the 1V and 1C1 are assumed to be independent, so only estimates for the corresponding standard deviations are presented in the S output.

4. Incorporating covariates in the NLME model

The general model building approach to be used for incorporating covariates in the model consists in starting with an NLME model with no covariates to explain inter-group variation, and using plots of the estimated random effects \hat{b}_i versus the candidate covariates to identify interesting patterns. Because the random effects accommodate individual departures from the population mean, plotting the estimated random effects against the candidate covariates provides useful information for the model-building process. A systematic pattern in a given random effect with respect to a covariate would indicate that the covariate should be included in the model.

If no interesting patterns are observed, the current model is kept, else, the covariate-coefficient pair with the most promising pattern is tested for inclusion in the model. The procedure is then applied sequentially, until no further interesting patterns are found.

The number of additional parameters to be estimated tends to grow considerably with the inclusion of covariates and, possibly, their associated random effects in the model. If the number of covariate-coefficient combinations is large, we suggest using a *forward stepwise* approach in which covariate-coefficient pairs are included in the model one at a time and the potential importance of the remaining covariates is graphically assessed at each step. The significance of the fixed-effects associated with a covariate included in the model is assessed using the Wald-type tests, based on asymptotic results for LME models (Pinheiro and Bates, 2000).

The inclusion of new random effects in the model when a covariate is added is rare, but should be investigated. The more common situation is that random effects can be eliminated from the model after covariates are included to account for inter-group variation. In both cases we proceed by comparing nested models using either likelihood ratio tests, or information criterion statistics (AIC and BIC). We illustrate the use of the proposed model-building strategy with the CO₂ uptake and Quinidine examples described in Section 3.

4.1. CO₂ uptake

The primary question of interest for the CO_2 data is the effect of plant type and chilling treatment on the individual model parameters ϕ_i . To plot the estimated random effects against the covariates, we first need to extract the \hat{b}_i from the fitted model and combine them with the covariates. The ranef function in NLME accomplishes that.

- > fm2CO2.RE <- ranef(fm2CO2, augFrame = T)</pre>
- > fm2CO2.RE

	Asym	lrc	Туре	Treatment	conc	uptake				
Qn1	6.17160	0.0483563	Quebec	nonchilled	435	33.229				
Qn2	10.53264	-0.1728531	Quebec	nonchilled	435	35.157				
Qn3	12.21810	-0.0579930	Quebec	nonchilled	435	37.614				

The augFrame argument, when TRUE, indicates that summary values for all the variables in the data frame should be returned along with the estimated random effects. When a covariate is constant within a group, such as Treatment and Type in the CO2 data, its unique values per group are returned. Otherwise, if the covariate varies within the group and is numeric, such as conc and uptake in CO2, the group means are returned; if it is a categorical variable, the most frequent values (modes) within each group are used.

The plot method for objects produced by the ranef function is the most useful tool for identifying relationships between individual parameters and covariates. The simple call below produces the plot in Figure 3.

> plot(fm2CO2.RE, form = ~ Type * Treatment)

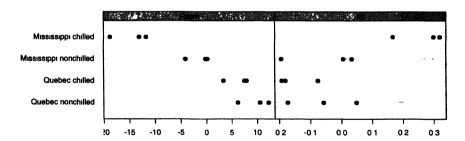


FIG 3. – Dotplots of estimated random effects corresponding to fm2CO2 versus all combinations of plant type and chilling treatment.

Figure 3 shows a strong relationship between the estimated random effects and the covariates – Asym decreases when the plants are chilled and is higher among Québec plants than Mississippi plants, with the increase in Asym from chilled to nonchilled plants being larger among Mississippi plants than among Québec plants, suggesting an interaction between Type and Treatment. There is also evidence of a Type by Treatment interaction on the log-rate – 1rc increases with chilling for Mississippi plants and decreases with chilling for Québec plants. We include both covariates in the model to explain the Asym and 1rc plot-to-plot variation. This leads to the following changes in ϕ_{1i} and ϕ_{2i} in model (5).

$$\phi_{1i} = \beta_1 + \gamma_{11}x_{1i} + \gamma_{21}x_{2i} + \gamma_{31}x_{1i}x_{2i} + b_{1i},$$

$$\phi_{2i} = \beta_2 + \gamma_{12}x_{1i} + \gamma_{22}x_{2i} + \gamma_{32}x_{1i}x_{2i} + b_{2i},$$

$$x_{1i} = \begin{cases} -1, & \text{Type}_i = \text{Qu\'ebec}, \\ 1, & \text{Type}_i = \text{Mississippi}, \end{cases}$$

$$x_{2i} = \begin{cases} -1, & \text{Treatment}_i = \text{nonchilled}, \\ 1, & \text{Treatment}_i = \text{chilled}, \end{cases}$$

where β_1 represents the average asymptotic uptake rate, γ_{11} and γ_{12} represent the plant type main effects, γ_{21} and γ_{22} represent the chilling treatment main effects, and γ_{31} and γ_{32} represent the plant type–chilling treatment interactions.

We update the fitted model, incorporating the covariates as in (10) through the fixed argument.

```
> fm3CO2 <- update(fm2CO2,
+ fixed = list(Asym + lrc \sim Type * Treatment, c0 \sim 1),
+ start = c(32.4, 0, 0, 0, -4.6, 0, 0, 0, 49.3))
```

Because the fixed-effects model has changed, new starting values must be provided. We use the previous estimates for β_1 , β_2 and β_3 and set the initial values for the new fixed effects to zero. Wald-type tests are then used to assess the significance of the individual fixed effects.

Value Std.Error DF t-value p-value Asym. (Intercept) 32.342 0.7849 64 41.208 <.0001 Asym.Type -7.990 0.7785 64 -10.264<.0001 Asym. Treatment -4.210 0.7781 64 -5.410 <.0001 Asym.Type: Treatment -2.725 0.7781 64 -3.502 0.0008 -4.509 0.0809 64 -55.743 lrc.(Intercept) <.0001 lrc.Type 0.133 0.0552 64 2.417 0.0185 1rc.Treatment 0.100 0.0551 64 1.812 0.0747 lrc.Type :Treatment 0.185 0.0554 64 3.345 0.0014

Fixed effects : list(Asym + lrc \sim Type * Treatment, c0 \sim 1)

. .

4.3646 64

11.573

<.0001

c0 50.511

All fixed effects introduced in the model to explain the variability in Asym are highly significant and the Type by Treatment interaction for 1rc is also highly significant, confirming the previous conclusions from Figure 3.

The inclusion of the experimental factors in the model resulted in a reduction in the estimated standard deviation for the Asym random effects from 9.66 to 2.35 and in the estimated standard deviation for the 1rc random effects from 0.20 to 0.08, indicating that a substantial part of the plot-to-plot variation in these two coefficients is explained by differences in plant type and chilling treatment.

Figure 4 displays the plots of the estimated random effects corresponding to fm3CO2 versus the experimental factors. As expected, no systematic patterns can be found.

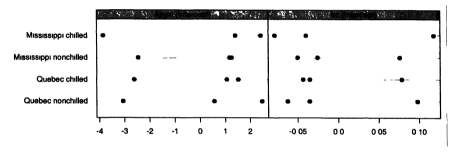


FIG. 4. – Dotplots of estimated random effects corresponding to fm3CO2 versus all combinations of plant type and chilling treatment.

After covariates have been introduced in the model to account for intergroup variation, a natural question to ask is which random effects, if any, are still needed. The ratio between a random-effect's standard deviation and the absolute value of the corresponding fixed effect gives an idea of the relative inter-group variability for the coefficient and is useful in deciding which random effects should be tested for deletion from the model. For the fm3CO2 fit these ratios are 7.3% for Asym. (Intercept) and 1.8% for 1rc. (Intercept), suggesting that the latter should be tested for exclusion first. The likelihood ratio test comparing fm3CO2 to a model with a single Asym random effect gives a non-significant p-value of 0.27, indicating that the 1rc random effect could be dropped after the inclusion of the covariates. A subsequent test comparing the simpler NLME model to a model without any random effects resulted in a highly significant p-value, indicating that the Asym random effect needs to kept in the model to account for unexplained plot-to-plot variation. The same remarks about the conservativeness of the classical likelihood ratio test for evaluating the removal of random effects from the model, included at the end of Section 3.1, also apply here. As before, the conclusions are unchanged when the corrected distribution of the test is used.

4.2. Quinidine

To investigate which covariates may account for patient-to-patient variation in the pharmacokinetic parameters, we first extract the estimated random effects, augmented with summary values for the available covariates (the modal value is used for time-varying categorical variables and the mean for time-varying numeric variables).

```
> fm1Quin.RE <- ranef( fm1Quin, aug = T )</pre>
> fm1Ouin.nlmeRE[1 :3,]
              1v
                         1C1
                               time
                                                    interval Age
                                                                   Height
                                        conc
                                              dose
109
      0.0005212 -0.0028369
                              61.58 0.50000
                                                           NA
                                                                70
                                                                        67
                                                NA
  70
                   0.3227614
       0.0362214
                               1.50
                                     0.60000
                                                           NA
                                                                68
                                                                        69
                                                NA
     -0.0254211
                   0.4402551 91.14 0.56667
                                                NA
                                                           NA
                                                                75
                                                                        72
     Weight
                   Race Smoke Ethanol
                                           Heart Creatinine
                                                                 glyco
 109
          58
                                         No/Mild
                                                        >= 50
                                                               0.46000
             Caucasian
                            no
                                   none
  70
          75
             Caucasian
                            no
                                 former No/Mild
                                                        >= 50
                                                               1.15000
  23
         108
                                   none No/Mild
                                                        >= 50 0.83667
             Caucasian
                           yes
```

The dotplot displays used to visualize the relationships between the estimated random effects and the covariates in the CO₂ example do not scale up well when there are a large number of groups, or a large number of covariates in the data, as in the quinidine study. Also, they cannot be used with numeric covariates, like Weight and Age. The plot method for objects returned by the ranef function actually allows a more flexible type of display for these situations. Relationships between estimated random effects and categorical variables are displayed using boxplots, while scatter plots are used for displaying the relationships between the estimated random effects and numeric covariates. Specifying a two-sided formula in the form argument, with the random effect on left-hand side and the desired covariates, separated by the + operator, on the right-hand side, indicates to the plot method that the more general display should be used. For example, to plot the estimated 1C1 random effects against the available covariates we use

```
> plot( fm1Quin.RE, form = 1Cl ~ Age + Smoke + Ethanol +
+ Weight + Race + Height + glyco + Creatinine + Heart)
```

The resulting plot, shown in Figure 5, indicates that clearance decreases with glycoprotein concentration and age, and increases with creatinine clearance and weight. There is also evidence that clearance decreases with severity of congestive heart failure and is smaller in Blacks than in both Caucasians and Latins. The glycoprotein concentration is clearly the most important covariate for explaining the 1C1 inter-individual variation. A straight line seems adequate to model the observed relationship.

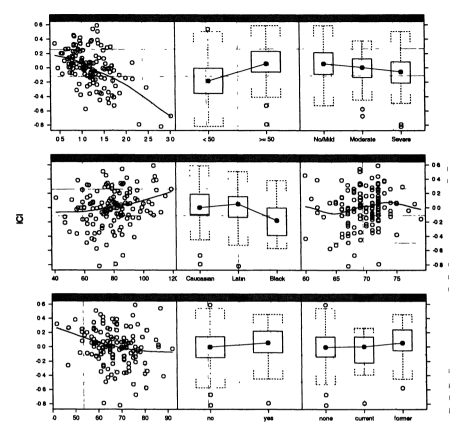


FIG. 5. – Estimated log-clearance random effects from model fm1Quin versus demographic and physiological covariates in the quinidine data. A loess smoother is included in the scatter plots of the continuous covariates to aid in visualizing possible trends.

Because of the number of observations per individual varies considerably in the quinidine study, the random effects are, as a result, estimated with different precisions. An alternative plot, taking the different precisions into account, would use standardized estimates $\hat{b}_{i}^{*} = \hat{b}_{i}/sd(\hat{b}_{i})$, where $sd(\hat{b}_{i})$ represents the estimated standard deviation of \hat{b}_{i} . This type of plot is not currently implemented in NLME, so we will stick here to plots based on the unnormalized \hat{b}_{i} .

Figure 6 presents the plots of the estimated 1V random effects versus the available covariates. None of the covariates seems helpful in explaining the variability of this random effect and we do not pursue the modeling of its variability any further.

Following the forward stepwise approach mentioned earlier, initially only the glycoprotein concentration is included in the model to explain the 1C1

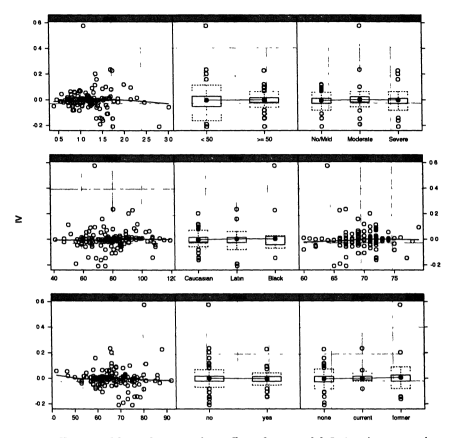


FIG. 6. – Estimated log-volume random effects from model fm1Quin versus demographic and physiological covariates in the quinidine data.

inter-subject variation according to a linear model. The modified version of model (9) is

$$lCl_{ij} = (\beta_1 + b_{1i}) + \beta_4 \text{glyco}_{ij}.$$

Because the glycoprotein concentration may change with time on the same patient, the random effects for 1C1 need to be indexed by both patient i and time j. The corresponding model is fitted with

```
Fixed effects : list(1V + 1Ka \sim 1, 1C1 \sim glyco)
                    Value Std.Error
                                        DF
                                           t-value
                                                     p-value
              ıν
                    5.3085
                              0.10244
                                       222
                                             51.818
                                                      <.0001
                  -0.6662
             lKa
                              0.30251
                                       222
                                             -2.202
                                                      0.0287
 1Cl.(Intercept)
                    3.1067
                              0.06473
                                       222
                                             47.997
                                                      <.0001
       1Cl.glyco
                  -0.4914
                              0.04263
                                       222
                                            -11.527
                                                      <.0001
```

As expected, the estimated lCl.glyco fixed effect is very significant, indicating that the glycoprotein concentration should be kept in the model.

To search for further covariates to include in the model, we investigate the plots of the estimated 1C1. (Intercept) random effects from the fm2Quin fit versus the covariates, presented in Figure 7.

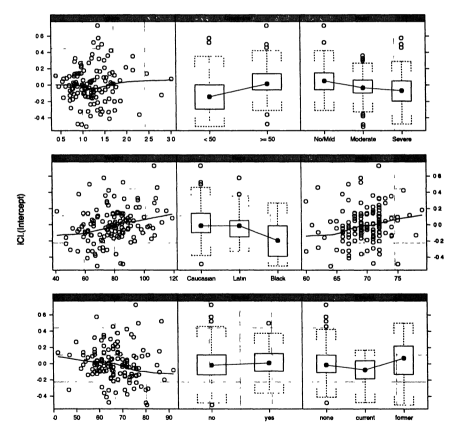


FIG. 7. – Estimated log-clearance random effects from model fm2Quin versus demographic and physiological covariates in the quinidine data.

As expected, there is no relation between the estimated random effects and glyco. The plots indicate that the estimated IC1. (Intercept) random

effects increase with creatinine clearance, weight, and height, decrease with age and severity of congestive heart failure, and are smaller in Blacks than in Caucasians and Latins. The most relevant variable appears to be the creatinine clearance, which is included in the model as a binary variable taking value 0 when creatinine is < 50 and 1 when creatinine is ≥ 50 .

```
> options(contrasts = c("contr.treatment", "contr.poly")
> fm3Ouin <- update( fm2Ouin,</pre>
   fixed = list(lV + lKa \sim 1, lCl \sim glyco + Creatinine),
   start = c(5.31, -0.67, 3.11, -0.49, 0)
> summary( fm3Ouin.nlme )
Fixed effects : list(1V + 1Ka \sim 1, 1C1 \sim glyco + Creatinine)
                    Value Std.Error
                                       DF t-value p-value
              JV.
                   5.2900
                             0.10631 221
                                           49.761
                                                    <.0001
             1Ka -0.7462
                             0.29635 221 -2.518
                                                     0.0125
 1C1.(Intercept)
                  2.9229
                             0.07221 221
                                            40.477
                                                     < .0001
       1Cl.glyco -0.4632
                             0.04117 221 -11.251
                                                    < .0001
  1Cl.Creatinine 0.2125
                             0.04491 221
                                             4.733
                                                     <.0001
```

The fixed effect corresponding to 1C1.Creatinine is very significant, as expected.

The final model produced by this stepwise model-building approach includes extra terms for indicator functions of the events race = Black, heart = No/Mild, and ethanol = former to explain the clearance variation. The corresponding model for the log-clearance is expressed as

```
lCl_{ij} = (\beta_1 + b_{1i}) + \beta_4 \text{glyco}_{ij} + \beta_5 \text{Creatinine}_i +
\beta_6 \text{I}(\text{Race} = \text{Black})_i + \beta_7 \text{I}(\text{Heart} = \text{No/Mild})_i + \beta_8 \text{I}(\text{Ethanol} = \text{former})_i
and the resulting fit gives
 > \text{summary}(\text{fm4Quin.nlme}) 
 . . .
 \text{Random effects} :
 . . .
 \text{IV 1C1.(Intercept) Residual}
```

0.62565

0.222

StdDev : 0.29855

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Fixed effects : list($IV+IKa \sim 1$, $IC1 \sim glyco+Creatinine+I(Race == "Black")+I(Heart == "No/Mild")+I(Ethanol == "former"))$

```
Value Std.Error
                                                 DF
                                                     t-value p-value
                        lV
                             5.3036
                                       0.10494 218
                                                      50.540
                                                               <.0001
                       1Ka
                            -0.7288
                                       0.29299 218
                                                      -2.487
                                                               0.0136
           1C1.(Intercept)
                             2.8585
                                       0.07459 218
                                                      38.322
                                                               <.0001
                 lCl.glyco
                            -0.4617
                                       0.03987 218
                                                     -11.582
                                                               <.0001
            1Cl.Creatinine
                             0.2086
                                       0.04366 218
                                                       4.779
                                                               <.0001
    lCl.I(Race == "Black")
                            -0.2697
                                       0.09409 218
                                                      -2.867
                                                               0.0046
1C1.I(Heart == "No/Mild")
                             0.1473
                                       0.05397 218
                                                       2.729
                                                               0.0069
1Cl.I(Ethanol == "former")
                                       0.06531 218
                             0.1528
                                                       2.340
                                                               0.0202
```

All 1C1 coefficients are significant at the usual levels.

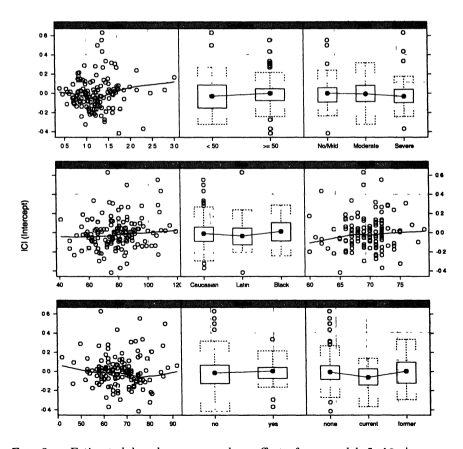


FIG. 8. – Estimated log-clearance random effects from model fm4Quin versus demographic and physiological covariates in the quinidine data.



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The plots of the estimated random effects for model fm4Quin versus covariates, displayed in Figure 8, do not suggest any further covariates to be included in the model. Even though there was a reduction in the estimated standard deviations for the 1C1 and 1V random effects, neither could be removed from the model.

As reported in previous analyses of the quinidine data (Davidian and Giltinan,1995), there is evidence that the variability in the concentration measurements increases with the quinidine concentration. This can be investigated using the tools available in the **NLME** library, but we will not pursue such analysis here.

5. Conclusions

Incorporating covariates in an NLME model allows inter-group variation to be explained, at least partially, via fixed effects. Besides enhancing the understanding of the model and reducing the dependency on random effects, this allows better predictions to be made from the model, especially for groups not previously observed. Often times, the inclusion of covariates in the model leads to the removal of random effects, or a simplification of the random effects model.

The NLME library in S-PLUS and R provides a flexible and rich environment for model building with covariates in NLME models, including powerful graphical tools for identifying interesting covariates and useful functions for summarizing covariates and combining them with estimated random effects.

The general model building strategy proposed in this paper is to, when feasible, start with a model with all coefficients having both fixed and random effects. This may lead to over-parameterization in the random effects model and the resulting $\widehat{\Psi}$ should be investigated for possible indications of numerical instability (the most common problem being very high correlations, in absolute value, between some of the random effects). When this happens, a more parsimonious random effects model should be identified via testing of nested models (information criteria such as the AIC and the BIC could also be used at this stage), in order to achieve greater numerical stability. This model should then be used as a starting point to identify potentially useful covariates to explain inter-group variation. Plots of the estimated random effects versus available covariates are particularly useful at this step. A forward stepwise approach is proposed, with the most promising covariate identified in the plots being included in the model at each step, then tested for significance, and plots of the new estimated random effects versus covariates investigated. The procedure continues until no further covariates are available, or promising covariates can be identified in the plots. Finally, after all useful covariates have been identified and included in the model, the remaining random effects should be re-evaluated for their need in the model. Comparisons of coefficients of variation and confidence intervals for standard deviations corresponding to the random effects are useful at this step.

With genetic data being increasingly collected in a variety of areas where NLME models are used, such as pharmacogenetic data in clinic trials, thousands of potentially useful covariates are becoming available for model building. This creates challenging statistical and computational problems, such as the need for automatic methods of selection that account for multiple testing, algorithms that scale-up with the number of fixed effects, etc. Further research is needed before the methods and software described here can be used in this type of applications.

Acknowledgments

I would like to thank Robert Faivre, Didier Concordet, and an anonymous referee for many helpful comments and suggestions that improved the quality and readability of this article.

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